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# Theory, practice, and design criteria for utilizing artificial reefs to increase production of marine fishes

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Increasing the production of marine fishes is a tractable goal with wide socioeconomic and ecological appeal. Ecosystem restoration projects that increase the amount of suitable habitat in an area and/or habitat quality enhance both fishery production and ecosystem services. Fortunately, there are a wealth of studies documenting the specific examples of restoring these services from successful artificial reef deployments. Considering the need to create future structures in a variety of scenarios and locations, it is salient to summarize the mechanisms through which increased secondary production occurs and the design considerations. To achieve this objective maximizing ecological processes including the provisioning of planktonic and epibenthic food resources and related trophic pathways, and those associated with life-stage specific habitat use (e.g., recruitment, juvenile survival, reproductive output) are critical mechanisms of productive reefs. We synthesized this information by addressing the structural and ecological theory of artificial reef design based upon physical attributes such as complexity, vertical relief, habitat heterogeneity, and spatial scale. Within this framework we summarized the mechanisms that may be used to increase secondary fish production and propose a general theory for optimization of these variables.

## KEYWORDS

secondary fish production, habitat restoration, shelter, habitat complexity, habitat heterogeneity, trophic pathways, ecosystem services

## Introduction

There is no debate that marine resources are crucial for current and future socioeconomic processes. Unfortunately, these resources have declined over time due to a variety of stressors including the depletion of species at higher trophic levels, habitat loss, pollution, and climate change, dramatically impacting marine ecosystems worldwide

(Jackson et al., 2001; Worm et al., 2006). Reduced fishery stocks and deterioration or loss of habitat, compounded by the increasing socioeconomic need to access marine resources highlights the theory that current secondary (fish) production and services were below the potential, and certainly historical potential, for marine ecosystems. While this theory is generally not tested directly (i.e., manipulation and/or calculation of a maximum versus current production potential), it has been indirectly demonstrated and is at the core of a variety of fishery management actions to improve stock sizes and sustainability (Knowlton, 2021). Well-designed ecosystem restoration projects increase the amount of suitable habitat area and/or quality, enhancing both fishery production and ecological services. Similarly, marine protected areas increase fishery stock biomass within their boundaries (Claudet et al., 2008; Edgar and Stuart-Smith, 2009), supporting this fundamental concept: current production is less than potential production in marine ecosystems.

Ongoing macroscale alterations of marine habitats and the implementation of infrastructure projects challenge us from a variety of perspectives. For thousands of years ‘human-made’, or as the literature primarily refers to them ‘artificial’ reefs, were installed to enhance marine habitats and there are continuous proposals for future implementation (Whitmarsh et al., 2008; Seaman, 2019; Tickell et al., 2019; Bugnot et al., 2020). Optimal artificial reef design is salient for deploying these structures to adequately restore lost ecosystem services. Herein, we delineate the theory and applicability of reversing or offsetting these losses, synthesizing the current empirical evidence for contributions of artificial reef habitats to local and regional ecosystems worldwide and the mechanisms supporting this goal. These theoretical constructs are essential for establishing design and evaluation criteria for utilizing reefing technology to improve ecosystem health and associated services.

Artificial reefs deployments have resulted in substantial increases of local and regional fish and invertebrate production. Johnson et al. (1994) found the rate of secondary production for a reef fish assemblage on a quarry rock artificial reef in southern California, USA was nine times greater than the sand-bottom fish assemblage the artificial reef replaced (i.e., was constructed on top of). Similarly, artificial reef habitat in Delaware Bay, USA supported up to two orders of magnitude higher secondary production (benthic macrofauna) per unit area than the comparable soft bottom habitat it replaced (Steimle et al., 2002). Large-scale artificial reef deployment assessments provide evidence for regional increases of production in fishes and invertebrates. In Japan, analyses of catch data revealed regional increases in octopus production in response to approximately 50,000 m<sup>3</sup> of artificial reef habitat construction (Polovina and Sakai, 1989). Following the deployment of an artificial reef complex in Algarve, southern Portugal, Roa-Ureta et al. (2019) analyzed 27 years of landings data to estimate the regional carrying capacity (K) of two-banded seabream, *Diplodus vulgaris*, increased by 35% (an additional 895 tons).

This large-scale reefing effort comprised over 21,500 concrete modules (small 3 m<sup>3</sup> or large 174 m<sup>3</sup>) covering an area of 43 km<sup>2</sup> (Santos et al., 2011). The Algarve artificial reef design focused on adding habitat for high-value (€12–18/kg in 2011) juvenile and adult fishes, which were targets of an artisanal fishery, to a soft bottom area where this habitat was previously rare (Santos et al., 2011). An economic analysis found revenues from fishing the areas around this artificial reef complex was greater than at control sites and catch rates and associated revenues had risen consistently over a 15 year period following deployment of the reef (Whitmarsh et al., 2008). Additionally, the artificial reef in Algarve relieved pressure from natural reefs in the region by redistributing fishing effort across a larger area (Leitao et al., 2009). These researchers also emphasized the need for additional fishery management regulations to prevent overfishing (Santos et al., 2011; Roa-Ureta et al., 2019). Thus, increased regional secondary production is achievable, but necessitates holistic design strategies that incorporate local and regional ecological processes, proper spatial scale and location, and socioeconomic factors such as management.

Impacts of a given artificial reef project were typically species- or taxa-specific (or even specific to a life-stage within a species), because responses to artificial reef habitats depend on species-specific functional attributes (e.g., diet, habitat use, life history) (Leitão, 2013; Smith et al., 2015; Cresson et al., 2019). Impacts were so diverse across taxa that predicting potential effects for entire assemblages was difficult and therefore a ‘case-by-case’ approach was generally more accurate (Brickhill et al., 2005). In paired comparisons between natural rocky or coral reefs and nearby artificial habitats (e.g., artificial reefs, shipwrecks, energy infrastructure), fish density, biomass and richness were often similar, demonstrating that artificial reefs have the potential to mimic natural reefs, if that is the objective (Paxton et al., 2020b). Most assessments of artificial reef designs examining specific objectives concentrated on important focal species (e.g., fishery or keystone species) and/or functional groups (e.g., those with specific trophic roles or habitat use patterns). Similarly, comparing reef success across ecosystems was highly context-specific and necessitates an understanding of regional goals as well as fine-scale biological and physical processes. The goal of increasing local and/or regional production is feasible, but a “one size fits all” approach is not advised (Komyakova et al., 2019; Paxton et al., 2020b; Blount et al., 2021).

Thus, a substantial amount of literature documented specific effects of individual artificial reef deployments, the mechanisms through which increased production occurred, and design considerations for achieving these objectives. Recently there has been more focus in the literature on the specific mechanisms, sometimes referred to as “enhanced ecological functions” (Glarou et al., 2020), that lead to increased secondary production of fish on artificial reefs, even above rates observed on natural reefs (e.g., Cresson et al., 2014a;

Glarou et al., 2020; Rouse et al., 2020; Puckeridge et al., 2021). These mechanisms provide benefits to individual species, typically through provision of shelter and/or food supporting survival or growth. While in some instances data were not available to directly quantify regional or local secondary production of fishes, evidence of these mechanisms, often in the context of a focal species' life history and behavioral ecology, support an assumption that reefs were contributing to secondary production. Yet, across ecosystems, there were consistent elements in artificial reef design that support increased secondary production and these criteria are useful for planning, evaluation, and design of current and future projects and have the potential to increase production on a regional scale. We posit that the most tractable metric to evaluate ecological services across habitats is secondary production.

## Ecological process considerations of reef design

### Planktonic and epibenthic food resources

A sizable proportion of the secondary production of fishes and invertebrates in artificial reef systems is fueled through planktonic resources, with a lesser proportion being supported by macroalgae or detritus. In a stable isotope and diet study of the largest artificial reef complex in the Mediterranean Sea (RECIFS PRADO program), organic matter of pelagic origin (i.e., phytoplankton and zooplankton) was the primary source supporting the system through artificial reef resident fishes feeding on sessile filter feeding invertebrates or directly on zooplankton (Cresson et al., 2014a; Cresson et al., 2014b). Similarly, a study of trophic pathways of fishes on a natural reef system in Australia observed that planktonic resources supported 60% of fishes, compared to approximately 30% by macroalgae and 10% by detritus (Truong et al., 2017). Current flow that delivers the associated flux of plankton and nutrients are critical factors supporting secondary production and thus are essential for overall performance. Therefore, understanding trophic bottlenecks and maximizing planktonic flux is necessary for increasing secondary production.

Zooplanktivorous fishes on artificial reefs are often the key trophic link between zooplankton and the larger reef-associated community (Bray et al., 1981; Cresson et al., 2014a; Holland et al., 2021; Puckeridge et al., 2021). Schooling water column and zooplanktivorous fishes were in greater abundance above higher relief purpose-built artificial reefs in temperate Australia (Puckeridge et al., 2021) and wrecks off of North Carolina, USA (Lemoine et al., 2019), when compared to lower relief natural reefs within each region. An Australian study directly documented the trophic link between Bluespotted Flathead

(*Platycephalus caeruleopunctatus*), a resident benthic ambush predator, feeding on pelagic forage fish above the high relief artificial reef modules (Holland et al., 2021). This mechanism has also involved nutrient transfer, such as the case with diurnal planktivorous pomacentrids in California that increase overall reef production with their metabolic byproducts being used by algae (Bray et al., 1981). Many artificial reef systems are therefore more productive than natural reefs because they provide sufficient habitat for zooplanktivorous fishes (Champion et al., 2015; Pondella et al., 2018; Holland et al., 2021).

With this in mind, flux of planktonic resources and even inducing upwelling to increase primary productivity (e.g., Itosu et al., 1995; Otake et al., 1995) are important considerations. Further, for reefs that receive higher current flow, such as offshore oil and gas platforms in California, this translated to increased growth rates for mussels (*Mytilus* spp.), a dominant filter feeding invertebrate, and the planktivorous Blue Rockfish (*Sebastes mystinus*) than individuals in nearby natural nearshore habitats (Page and Hubbard, 1987; Blanchette et al., 2007; Love et al., 2007). Similarly, a study of an experimental artificial reef off of Scotland demonstrated the importance of water flow for the secondary production of a suspension feeding bryozoan both on the outer-facing reef surfaces and into internal reef spaces, with higher secondary invertebrate production found on reefs with more complex designs and larger internal reef spaces (Rouse et al., 2020). Clearly in order to maximize secondary production, optimizing both benthic and pelagic food resources is essential.

### Shelter

Proper shelter allows fishes to maximize the acquisition of planktonic and epibenthic food resources while minimizing the amount of energy needed to maintain themselves. In a system with underutilized food resources, the increase of shelter and associated use by various life history stages of fishes theoretically results in greater production. Beyond increasing appropriate habitat where it is limited or lost, properly designed habitats provide a direct survival and metabolic benefit to the fish sheltering on them. In Atlantic Cod, metabolic rates associated with reef habitat (as compared to sand) were lower, suggesting that the energy saved by sheltering contributed to increased somatic growth and thus production rates (Schwartzbach et al., 2020). Reef holes and cavities (i.e., void spaces) provide refuges, and hole size was often positively associated with the body sizes of fishes found sheltering in them (Hixon and Beets, 1993; Beets and Hixon, 1994; Friedlander and Parrish, 1998). For example, smaller reef fishes had higher survival rates on reefs with smaller hole sizes compared to similar sized fish on reefs with larger holes (Hixon and Beets, 1993). Some reef designs utilized models to predict hole/crevice size and number based on the rock size (Barry and Wickins, 1992), suggesting the possibility of

designing reefs with appropriate hole sizes for a specific taxa or life history stages (e.g., [Glarou et al., 2020](#)). Survival and growth of fishes is a function of both predator avoidance and the reduced energy associated with sheltering from current flow in appropriately sized holes. Productive reefs have shelter characteristics enabling fishes to minimize their energy expenditures from flow regimes and predator avoidance while allowing them to maximize the flux of energy from the plankton and epibenthic prey. Theoretically increasing shelter increases the secondary production a reef supports. This is ultimately limited by the carrying capacity of the system (e.g., food availability, species and life-stage specific survival rates) in a density dependent manner resulting in an asymptotic relationship between shelter and production ([Figure 1](#)).

## Recruitment and juvenile survivorship

The function of artificial reefs as shelter for recruitment, survivorship, and growth of young-of-year and juvenile reef-associated fishes is another essential component for increasing local and regional production. Recruitment and juvenile success,

the hallmarks of fishery models ([Ricker, 1975](#)), are a critical factor for stock management. There are multiple examples of artificial reefs providing habitat for newly settled and/or juvenile fish, which increased recruitment rates and cohort success, which in turn increased ecosystem-wide services. High prevalence of newly recruited and/or juvenile fishes on artificial reefs was common (e.g., [Santos et al., 2011](#); [Reubens et al., 2014](#); [Granneman and Steele, 2015](#); [Krone et al., 2017](#)). Fish assemblages on more complex, larger-scale artificial reefs supported a higher proportion of smaller juvenile fishes (e.g., [Santos et al., 2011](#); [Granneman and Steele, 2015](#); [Love et al., 2019](#)) suggesting these patterns documented on small experimental reefs (cm<sup>2</sup> to m<sup>2</sup>) are scalable to larger systems (10's to 100's of m<sup>2</sup>). Additionally, [Komyakova et al. \(2019\)](#) found in temperate southern Australia that the location relative to larger reefs, either natural rocky reefs or large breakwaters, had a greater influence on fish densities at smaller artificial reef modules. This was attributed to increased recruitment of young-of-the-year fishes to these larger reefs and subsequent movement of those fishes to the artificial reef modules. Undoubtedly, incorporation of a nursery design is a key component for successful reefing programs.

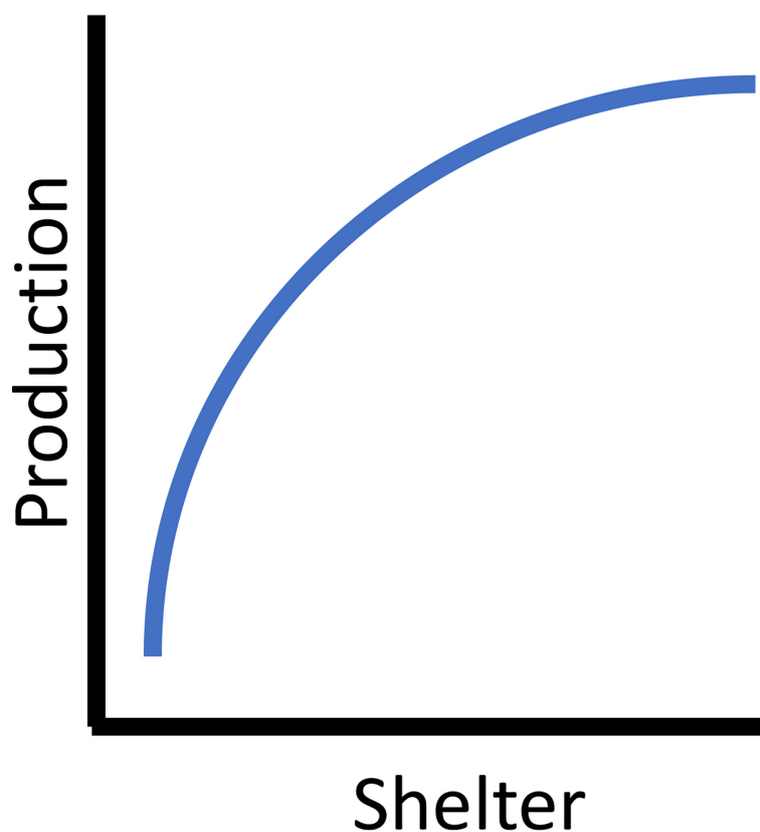


FIGURE 1

The asymptotic theoretical relationship between available shelter and secondary (fish) production.

In a variety of ecosystems, increasing habitat for juvenile fish resulted in corresponding production increases. Windmill artificial reefs (WARs), consisting of monopile offshore wind turbines surrounded by quarry rock scour protection and deployed in largely soft bottom regions of the North Sea, supported high abundances of juvenile fishes. Specifically, they were suitable feeding grounds for juvenile Atlantic Cod (*Gadus morhua*) and Pouting (*Trisopterus luscus*), which fed upon epifaunal invertebrate prey and showed high seasonal site fidelity (De Troch et al., 2013; Reubens et al., 2014). Juvenile Atlantic Cod sheltered on WARs between dawn and dusk (crepuscular) feeding periods, suggesting sheltering from predators provides a survival advantage (Reubens et al., 2014), in addition to metabolic energy savings (Schwartzbach et al., 2020). In passive acoustic tracking of Atlantic Cod, these patterns were consistent over 2–5 month periods, with fish sheltering on the same WAR reef daily (Reubens et al., 2014). While these increases were in local production and regional-scale increases had not yet been observed, these juvenile fish likely emigrated as they matured, benefitting commercial fisheries. Similarly, in a mark-recapture study, juvenile Gag Grouper (*Mycteroperca microlepis*) had high site fidelity (average 300 days) on an experimental concrete artificial reef off Florida. Adults then moved further offshore where they became part of the broader fished stock, with two fish being recaptured on other side of the Gulf of Mexico in Texas and Mexico (Lindberg et al., 2006). In the Algarve artificial reef complex, which was designed to specifically provide shelter for juvenile fishes of fishery importance (Santos et al., 2011), this juvenile sheltering effect made an important contribution to the increases in regional carrying capacity (K) and production of Two-Banded Seabream (Roa-Ureta et al., 2019). Up to 88% of the fish assemblage associated with the Algarve reef modules were juveniles, with consistent recruitment of young-of-the-year fishes observed for some species. Juveniles were also documented as sheltering in the structure and feeding directly on the benthic macrofauna or in the water column near the structures (Leitao et al., 2009; Santos et al., 2011). These examples illustrate the progression necessary for increasing production: creating habitat where appropriate, documenting the density of critical life stages and their associated habitat use, then linking this to regional production. Clearly, reducing habitat and recruitment limitations by increasing nursery areas has significant benefits.

## Reproductive output

In addition to providing a nursery function and sheltering juvenile fishes with high rates of somatic production, a frequently overlooked component in artificial reef studies is reproductive output, or gonadal production. Reproductive output is a function of the number of larger adult fishes residing on a reef, and is a

“major component of energy flow through the fishes in [a] reef system” (Demartini et al., 1994). In this study, fish gonadal production on artificial reefs was equivalent to, or a large multiple of, the somatic production estimates. For territorial species, reefs with more suitable habitat have a corresponding greater number of potential feeding and mating territories (Hixon, 1980), which can lead to increased secondary production compared to less complex natural reefs (Pondella et al., 2002). These concepts are frequently incorporated in fisheries management and conservation network designs, such as MPA connectivity, with the overall goal of increasing localized spawning stock biomass and exporting it throughout the ecosystem (Russ et al., 2004; Pérez-Ruzafa et al., 2008; Almany et al., 2009; Botsford et al., 2009; Cudney-Bueno et al., 2009; Foley et al., 2010; Berumen et al., 2012; Schmiing et al., 2017; Baetscher et al., 2019; Marshall et al., 2019). Further, since most fishes have a pelagic larval stage, understanding gonadal production was a critical feature in modeling connectivity and the link between biomass density and recruitment (Pondella et al., 2015). Particularly well-studied for fishes, gonadal production and egg quality is a function of female size and age. Larger females have higher fecundity and allocate more energy for egg production, and in some instances, produce more fit larvae than smaller females (Berkeley et al., 2004). Importantly, this relationship was not linear, instead being hyperallometric with size (Barneche et al., 2018), and supported the idea that the protection and production of larger females disproportionately and positively affects stock structure. Larger adults have lower annual somatic production but theoretically are a major link to regional increases, especially if habitats are not limiting recruitment. Thus, habitat design and/or management actions taking this into account (e.g., the implementation of MPAs or fishery management practices such as slot limits and protection of spawning aggregations) have a greater impact on recovery due to increased gonadal production and potentially more fit larvae (Claudet et al., 2008; Edgar and Stuart-Smith, 2009; Marshall et al., 2019). Habitat characteristics that optimize spawning aggregations, generating larval sources throughout an ecosystem were also important considerations (Schmiing et al., 2017). Some studies identified larval production as a key management action for fishery replenishment (Kough et al., 2019). Higher quality habitats also support greater densities of large individuals, magnifying this effect.

## Physical considerations of reef design

### Structural complexity

Increasing reef complexity, both externally (rugosity) and internally (void space), improves secondary production (Blount et al., 2021). Simply increasing the complexity of reef modules in the

French Mediterranean doubled overall fish species richness, increased density by ten times, and increased biomass by forty times (Charbonnel et al., 2002). Experimental artificial reefs across a broad geographic range consistently found increased fish density, biomass, and/or species diversity associated with more complex structures (North Carolina, USA : Potts and Hulbert, 1994, South Carolina, USA: Kellison and Sedberry, 1998, Gulf of Mexico, Florida, USA: Sherman et al., 2002, Mediterranean, Italy: Relini et al., 2007, Scotland, UK: Hunter and Sayer, 2009, temperate southern Australia: Komyakova et al., 2019). Often fish density and size structure have a greater impact on rates of secondary production than relative differences in individual growth rates. For example, Granneman and Steele (2014) compared somatic growth rates, estimated from otolith back-calculation, on paired artificial and adjacent natural reefs in southern California, USA. When combined with observed density and size structure of the fish assemblage, they found secondary production rates on quarry rock artificial reefs were similar to or greater than natural reefs, and positively correlated with habitat characteristics (i.e., the abundance of larger boulders on artificial reefs). The authors also found that size-specific differences in fish density made a large contribution to the differences in production, while growth rates among sites were similar. The general mechanism discussed for this phenomenon relates direct sheltering opportunities for a variety of sizes of fishes

provided by the increased number and diversity of sizes of holes and crevices in the reefs. Further, more rugose reefs are more suitable for attachment of both algae and sessile invertebrates (Harlin and Lindbergh, 1977; Walters and Wethey, 1996). As such, there is a positive relationship between complexity and production. In addition, three-dimensional complexity adds increased surface area, elevating the overall abundance of benthic forage resources, which results in increased fish density, an integral factor in secondary production models. Theoretically, this relationship asymptotes with increasing complexity (Figure 2). While increasing complexity adds structural elements that correspond to species-specific functional attributes (e.g., diet, habitat use, life history) of the fish assemblage, similar to within reef habitat heterogeneity, this is ultimately limited by the surface area to volume structural relationship and the availability of internal spaces of a reef of a given size (Kim et al., 1994; Lemoine et al., 2019; Rouse et al., 2020). Thus, increasing complexity to maximize production is a theoretical and practical design consideration.

## Vertical relief

Vertical relief (i.e., height above the seafloor) is another commonly investigated factor in reef design and artificial reefs

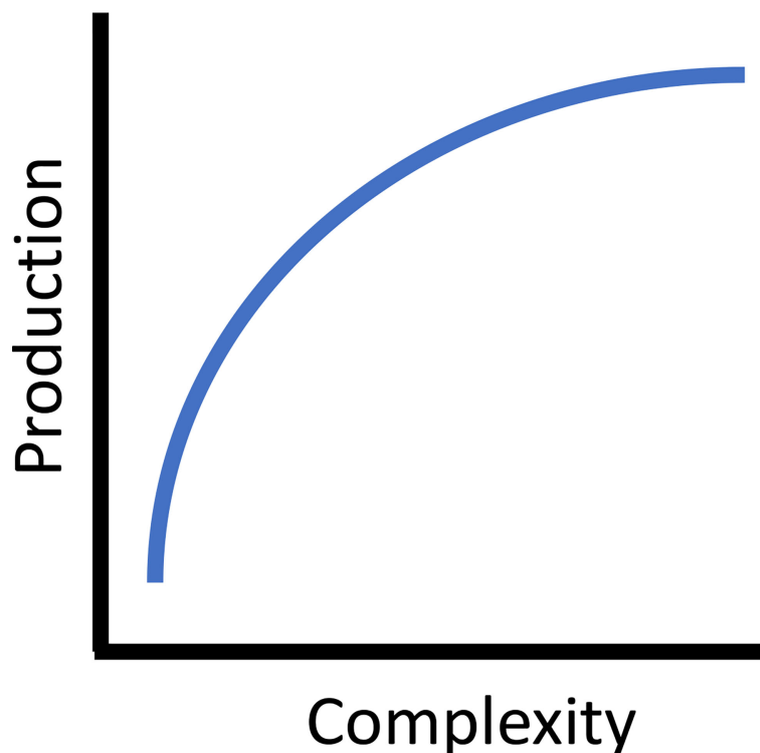


FIGURE 2

The asymptotic theoretical relationship between reef complexity and secondary (fish) production.

that provide sufficient vertical relief support greater taxonomic diversity (Ogawa, 1967; Molles and Manuel, 1978; Bohnsack et al., 1994). Positive relationships between vertical relief and fish density and/or biomass were documented at quarry rock artificial reefs in California, USA (e.g., Ambrose and Swarbrick, 1989; Pondella et al., 2006; Granneman and Steele, 2015). Studies comparing fish assemblages on shipwrecks (i.e., high-relief artificial reefs) with natural reefs and/or concrete artificial reefs found higher-relief wrecks have higher mean densities of fishes (northeast Pacific: Bulger et al., 2019, North Carolina, USA: Lemoine et al., 2019; Paxton et al., 2020a), as well as differences in trophic guilds. Similarly, Paxton et al. (2020a) observed wrecks had higher densities of transient predators (e.g., jacks, mackerel, barracuda, sharks) than concrete and natural reefs, and their review of the artificial reef literature suggested this pattern of more high trophic level (i.e., predatory, typically transient) fishes on artificial reefs than natural reefs was a global pattern. Yet, shipwrecks were not necessarily adequate surrogates for natural reefs (Medeiros et al., 2021). This is representative of a general pattern of predators being more common on artificial reefs globally, with transient predators being observed in higher densities on taller structures (Paxton et al., 2020a). While there were numerous objective and

anecdotal observations of this phenomenon, salient are the mechanisms and their potential utility in reef design.

Reefs with greater relief perform better than low relief reefs. At the spatial scale of typical reefs (meters), theoretically this will asymptote (Figure 3). While many species of fish orient to high relief structures for feeding purposes (as described previously) they also utilize them for spawning sites. Such spawning aggregations were often associated with the highest relief components within a reef and in some taxa optimal spawning sites were favored. Additionally, adults return to traditional spawning sites based upon a learned response, yet in the absence of this learning, they chose new sites based upon reef features implying there was an optimization of assessed reef features to maximize spawning success (Warner, 1988). Interestingly, on newly deployed reefs adult fishes sometimes within hours, but certainly within days orient to these structures, which, since they were just placed in the water, do not yet have epibenthic resources (Turner et al., 1969). Over the long term, large predatory fishes migrate and orient to these vertical features (Barilotti et al., 2020; Burns et al., 2020; Paxton et al., 2020a). However, very high-relief reefs, particularly those with an intertidal component (i.e., breakwaters, some shipwrecks, and oil platforms), had increased ecosystem heterogeneity and

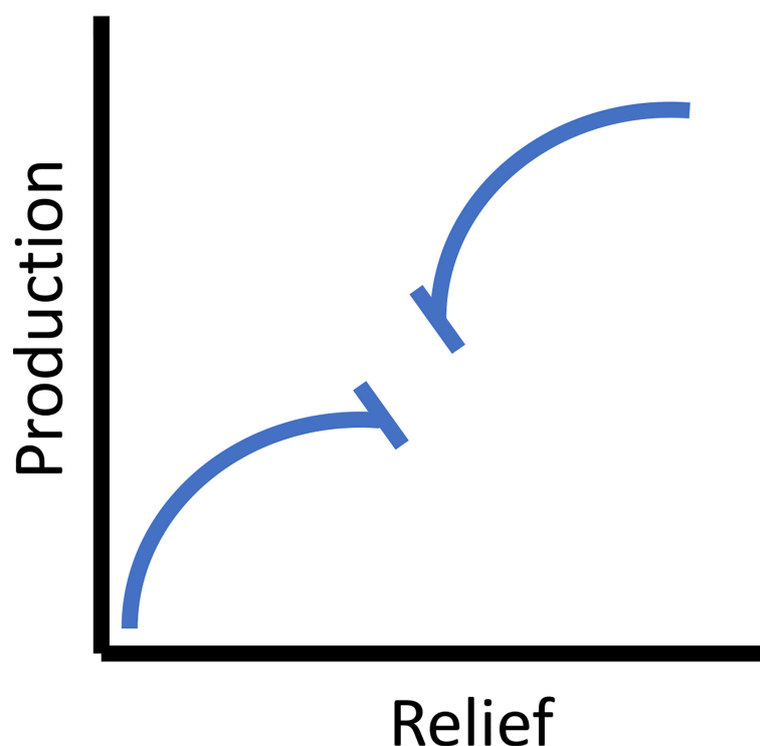


FIGURE 3

The theoretical relationship between vertical relief and secondary (fish) production. For the scale of typical reefs (meters) the relationship asymptotes (left side on x-axis), and then for very high relief reefs (10's of meters) and those that include an intertidal component (e.g., breakwaters, some shipwrecks, and oil platforms) the relationship is extended until it asymptotes again (right side on x-axis).

associated production (e.g., Stephens et al., 1994; Claisse et al., 2014). For example, breakwaters incorporate the intertidal zone, increasing habitat heterogeneity and have high rates of secondary production (Pondella et al., 2002). These processes are consistent with the inclusion of kelp, whereby a three-dimensional relief component is extended to the surface by a canopy providing habitat heterogeneity and trophic resources that support an increased suite of life stages and species (Graham, 2004). Structures with these high-relief components also act as fish aggregating devices (FADs) for more transient pelagic fishes. This attraction can occur on a non-random consistent basis, resulting in increased overall production through the inclusion of planktivores into the system (Smith et al., 2016). Relief is an attraction mechanism of adults and larval fishes, potential spawning sites, and food resources. On a typical reef module without an intertidal component, production as a function of relief asymptotes (especially if resource dependent), then adding an additional three-dimensional FAD/kelp/intertidal component increases production to a theoretical maximum (Figure 3).

## Habitat heterogeneity, reef size, and spacing

In many systems, nursery areas are often decoupled from adult habitats. Thus, increasing habitat heterogeneity (i.e., variability of habitat types across a given landscape area) (Wedding and Yoklavich, 2015), is a critical step towards optimizing habitat for multiple life stages. Fishes, either through ontogenetic changes or during specific life history stages, often use various habitats within an ecosystem. For many reef associated species this includes foraging in surrounding soft-bottom areas. A “halo effect” documented around artificial and natural reef structures that were surrounded by soft-bottom habitat largely refers to direct and secondary effects of the organic enrichment of these sediments. These halo effects were due to complex processes. Hydrodynamic changes from the increased vertical relief of an artificial reef directly affected the composition and size of sediments, and created a plume of detritus and waste generated by reef inhabitants (e.g., filter feeding invertebrates, planktivorous fishes, macroalgae) organically enriching sediments at scales of meters to tens of meters around an artificial reef (Heery et al., 2017; Bugnot et al., 2020). Changes in the composition and densities of sediment infauna (e.g., polychaetes and amphipods) at these scales [e.g., out to 30 m in Reeds et al. (2018) and out to 80 m in Bortone et al. (1998)], were a result of being fed by the additional organic material and increased predation from reef-associated fishes and crabs. These soft-bottom foraging grounds were important in supporting the diets (i.e., production) of many species that shelter in reefs. Many taxa utilize these reef/soft bottom ecotones, emphasizing the importance of the halo effects. Multiple studies have focused

on assessing the foraging distance of fishes from artificial reefs (Bortone et al., 1998; Brandt and Jackson, 2013) to minimize the overlap of foraging areas in reef design (Champion et al., 2015; Rosemond et al., 2018), which has the potential to limit growth and reef productivity (e.g., Lindberg et al., 2006). Johnson et al. (1994) found 20-25% of reef associated fish diets on a California artificial quarry rock reef derived from sand-rock ecotone habitats. These ecotones in southern California support increased densities of a commercial serranid, Barred Sandbass (*Paralabrax nebulifer*) (Anderson et al., 1989). Pondella et al. (2018) increased the amount of ecotone habitat by designing a reef with offset modules to increase the reef perimeter. Rosemond et al. (2018) found that reef-associated fish species were mostly foraging in the soft-bottom habitat up to 30 m away from a variety of artificial reef types (e.g., concrete pipes, metal ships) off North Carolina, USA. Reeds et al. (2018) documented reef fish foraging at elevated densities only out to 15 m from a steel fishery enhancement, but because the reef had a relatively small footprint (12 m x 16 m), they calculated that reef associated fishes were using an area 15 times greater than the reef footprint itself, demonstrating that for some species, a relatively large area of these surrounding sediments play an important ecological role. In the Adriatic Sea, Tyrrhenian Sea, and southern California, physical factors associated with artificial reef proximity outweighed the influence of predators (Ambrose and Anderson, 1990; Danovaro et al., 2002; Fabi et al., 2002) illustrating the importance of these ‘halos’.

Extending the ‘halo effect’ by incorporating the utilization of planktonic resources is conceptually a ‘dome effect’, as reefs extend three-dimensionally up into the water column. Multiple studies have taken a mechanistic approach in examining how the amount of shelter a reef provides interacts with the trophic processes of artificial reef residents who feed in adjacent habitats (e.g., water column, soft bottom, or a mix of soft bottom and natural reef). These studies suggested there is a trade-off (and potentially thresholds) between providing shelter for more individuals within the artificial reef and competition among those individuals for planktonic (Champion et al., 2015) or benthic food resources in the surrounding habitats (Lindberg et al., 2006; Brandt and Jackson, 2013), or both (Lindquist et al., 1994). Champion et al. (2015) empirically derived the amount of zooplankton that flowed across an artificial reef in Australia and the amount consumed per day by individuals of a resident fish species (*Atypichthys strigatus*, a smaller Kyphosidae). They modeled thresholds at which reefs were able to shelter more fish than the zooplankton flow could support, suggesting an optimal reef size exists to balance this tradeoff. They also noted that reef shape was important. Lindberg et al. (2006) found that growth was density-dependent in larger juvenile and young adult Gag Grouper (*Mycteroperca microlepis*) sheltering in concrete artificial reefs off Florida, USA and attributed it to competition for food in the surrounding feeding grounds. When also factoring in the potential for increased fishing mortality, if

fish were in higher densities, the authors argued that most positive effect on stocks was achieved through widely scattered patch reefs with appropriately sized cavities, as that should increase growth rates of individual fish while they were residing in the reefs. It is also important to note that this species used these reefs during an older juvenile/younger adult stage prior to moving further offshore, again emphasizing that habitat use patterns associated with artificial reefs are species and life stage dependent. For example, adult wrasses (Labridae) shelter nocturnally in sand proximate to reefs. Some wrasses on coral reefs have a post-settlement stage where they shelter in the sand prior to recruitment to the reef (Hamilton et al., 2008). For overall fish productivity, there are tradeoffs between distance between reef modules, size, spacing, and maximization of the halo and ecotone effects, as these effects are synergistic to overall performance (Figure 4). But adding more structure without optimizing spacing criteria may have diminishing returns. As such, in a particular reef system, the heterogeneity of subhabitats has a maximum (Figure 4). Theoretically, each increase in subhabitat type theoretically increases the production to a maximum.

Construction logistics, available funding, engineering, regulation restrictions or boundaries, and costs constrain reef size and associated local and regional effects (Lan and Hsui,

2006). Often it is necessary to weigh the benefits of one or a few larger reefs versus many smaller reefs optimizing the use of limited resources (e.g., Pondella et al., 2018). Some studies that have directly addressed reef size suggested intermediate sizes were most productive (reef volumes from 400–4000 cubic meters, (Ogawa et al., 1977), as cited and discussed in Glarou et al. 2020). For example, on reefs constructed with concrete cubes with holes in Taiwan, the biomass of resident fish species increased sigmoidally when compared to reef size, indicating that intermediate sizes produced the best ratio of standing stock biomass to construction cost (Jan et al., 2003). Further, reefs that are too large will reduce habitat heterogeneity and ultimately reduce production for the suite of species that depend on these varied resources (Figure 5) The trade-offs between costs and design criteria are important considerations for maximizing fish production with available resources.

## Additional considerations for realizing benefits

There are species-specific timescales for expected fishery benefits, based on life history and behavioral ecological traits. These benefits follow periods of good larval survival and

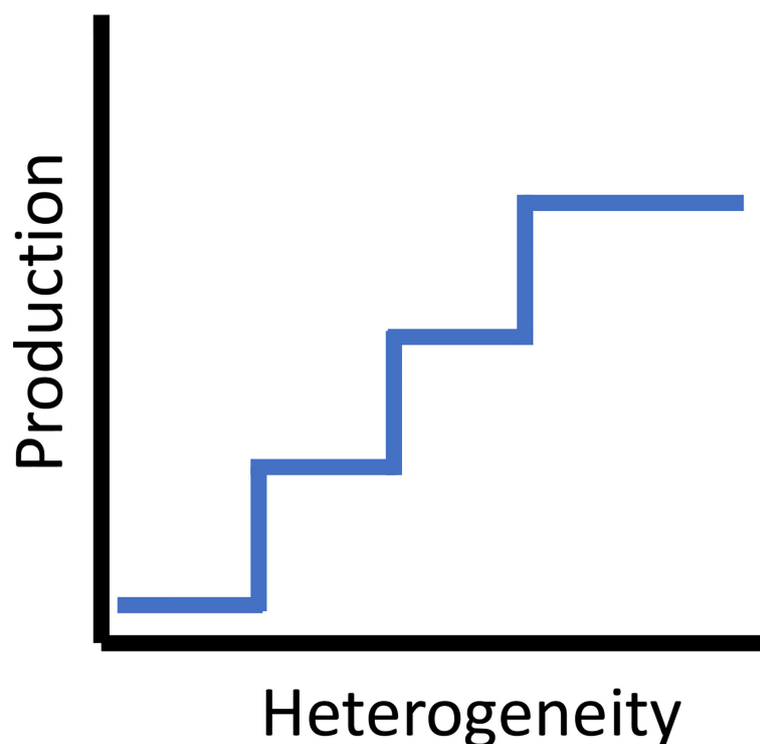


FIGURE 4  
The theoretical step relationship between habitat heterogeneity and secondary (fish) production.

recruitment. In Algarve, the Seabream fishery regional carrying capacity increased by 35% eight years post construction (construction took five years) as new production attributed to the artificial reef spilled over to the surrounding continental shelf. It followed multiple strong recruitment years, and then a four year lag for the species to enter the fishery (Roa-Ureta et al., 2019). An earlier economic analysis for the initial pilot-scale deployments of artificial reefs in this region also demonstrated fishery value (i.e., revenue) per unit effort slowly increased over a 15 year period following reef deployment and was higher near artificial reef modules when compared to fishing near control areas (Whitmarsh et al., 2008). These are consistent with timelines reported for maturity in Marine Protected Areas (MPA). Typically, post-closure fish densities and biomass asymptote at an interval of at least 15-20 years with older reserves outperforming younger reserves (Abesamis and Russ, 2005; Claudet et al., 2006). These time frames for maximizing reef contribution on both local and regional scales parallel typical age at maturity, the significantly greater gonadal output for larger females, and overall longevity for many focal species.

The installation of an artificial reef complex took pressure off natural reefs in the region by redistributing fishing effort across a larger area (Leitao et al., 2009). As such, spacing and connectivity to other reefs are important design criteria. In some instances, reef development and spacing protect soft-

bottom species by acting as barriers to bottom trawling (Guillen et al., 1994). Shallow anthropogenic seafloor structures influence the foraging of marine mammals (Arnould et al., 2015; Todd et al., 2020). The interplay among the potential to redistribute and/or enhance mammal populations as well as attracting these predators to artificial reefs (Fernandez-Betelu et al., 2022) is an open question necessitating further investigation as these predators could increase the natural mortality of fish and invertebrate populations. Determining mechanisms for (or direct evidence of) increased survival or growth of fishes also provided evidence that artificial reef systems are not functioning as ecological traps (Reubens et al., 2014). In such traps, an organism is attracted to and settles preferably in a habitat with suboptimal conditions relative to other available and more suitable habitats (Robertson and Hutto, 2006). If an ecological trap occurs, growth, recruitment and survival rates are lower than at reference habitats, as was recently described for Reef Ball reefs in Australia (Komyakova et al., 2021). Although better alternative habitats are available, the suboptimal habitat is unfortunately chosen, resulting in reduced production. While there are few examples of this in the literature, demonstrating a net benefit to a species with certain characteristics (e.g., life history, habitat use or movement patterns, trophic traits/groups) is critical. There is also the potential for artificial reefs to support or expand the

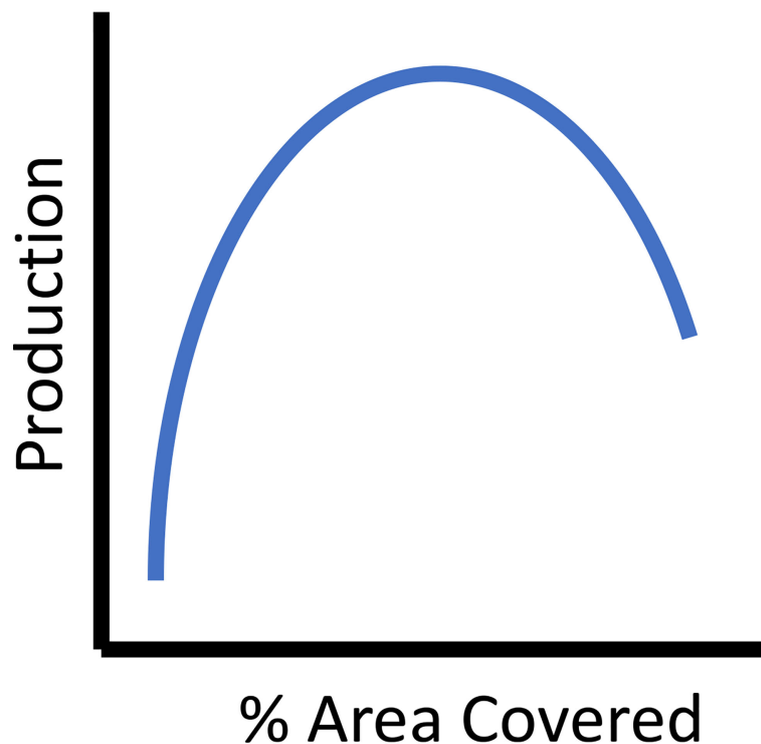


FIGURE 5

The theoretical relationship between percent of area covered, as a metric of reef spacing criteria and secondary (fish) production.

range of invasive species (Dahl and Patterson, 2014; Dafforn, 2017; Schulze et al., 2020). In heavily overfished populations (i.e., the current standing stock is not producing enough reproductive output to begin with), adding additional, even higher quality, habitat will not realize increased production if recruitment levels are low (Powers et al., 2003). In these cases, considering larger location-specific integrative coastal management approaches (Komyakova et al., 2019; Paxton et al., 2020b) and additional fishery management is necessary to achieve or sustain positive outcomes (Powers et al., 2003; Whitmarsh et al., 2008; Roa-Ureta et al., 2019).

Artificial reefs have a long history of use in fishery management (Bortone et al., 2011). An emerging concern in the literature was the consideration of artificial reefs as part of larger location-specific integrative coastal management approaches (Komyakova et al., 2019; Paxton et al., 2020b) and the potential necessity of additional fishery management for achieving or sustaining positive outcomes (Powers et al., 2003; Whitmarsh et al., 2008; Santos et al., 2011; Roa-Ureta et al., 2019). While not a concern when artificial reefs are primarily providing a nursery function to life stages with sizes below those targeted by fisheries (e.g., oyster reef restoration in estuaries in southeastern USA: Peterson et al., 2003, WARs in the North Sea: Reubens et al., 2014), this was a consideration when artificial reefs are being used by adult stages of species targeted heavily in local fisheries. On one hand, if an artificial reef or complex is relatively large (with respect to natural reefs in the area), then it may take pressure off natural reefs in the region by redistributing fishing effort across a larger total area reef habitat (Leitao et al., 2009). There are complex interactions within habitats, for instance reviews cite many studies where artificial reefs essentially stop bottom trawling in the area and may have positive impacts for soft bottom species (Relini et al., 2007; Brandini, 2014; Heery et al., 2017; Glarou et al., 2020). However, if the artificial reefs instead aggregated existing biomass in the region making target species easier to exploit (the goal of Fish Aggregating Devices [FADs]), and if this fishing mortality was not counter-balanced with a proportional increase in localized production (Lindberg et al., 2006; Smith et al., 2015), essentially the crux of the 'attraction vs production' debate (Osenberg et al., 2002), then consideration of additional fisheries management actions is prudent. In the case where an artificial reef serves as a spawning aggregation site (Schmiing et al., 2017), this necessitates potential permanent or reproductive season fishing closures. Balancing this against the potential secondary production increases due to increasing gonadal production are the challenges managers face. More broadly though, if objectives include amplifying regional impacts of deploying an artificial habitat (e.g., population recovery or increasing regional carrying capacity), then considerations including additional management actions focused on spawning stock biomass recovery, and particularly the protection of larger/older individuals, like in the implementation of MPAs and size limits, were practical

(Claudet et al., 2008; Edgar and Stuart-Smith, 2009; Barneche et al., 2018; Marshall et al., 2019). Thus, incorporating an ecosystem-wide approach in reef design is necessary.

Overall successful reefing projects optimize all these discussed criteria (shelter, complexity, heterogeneity, relief and spacing) to maximize reproductive output and recruitment success. These parameters are ecosystem- and taxa- specific; thus, wherever possible a thorough and understanding of the geographical, ecological, and socioeconomic context is necessary.

## Author contributions

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

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## Conflict of interest

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