Modified habitats change ecological processes affecting a non-indigenous epibiont

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ABSTRACT: Urbanisation of coastal habitats, particularly the increased numbers of pier-pilings, jetties and seawalls along shorelines, affects natural systems. Epibiota on secondary substrata (kelps, Ecklonia radiata) in man-made structures differ from those in natural habitats, but they have received considerably less attention. To understand the consequences of changes in the structure of these assemblages, it is therefore necessary to determine which ecological processes are being affected and the factors influencing them. In Sydney Harbour, kelps on pier-pilings supported greater covers of bryozoans, particularly of the non-indigenous species Membranipora membranacea, than found on natural reefs. Experimental transplants of kelps without epibiota from reefs to pilings showed that recruitment and growth of colonies of M. membranacea were much greater on pilings than on reefs. Patterns of distribution and abundance of this epibiotic bryozoan are determined by a combination of these processes and probably influenced by differences in abiotic and biotic characteristics between habitats. Understanding how these components of habitats affect ecological processes is necessary to allow sensible prediction of the effects of modifying habitats on the ecology of organisms.

KEY WORDS: Urbanisation · Modification of habitat · Kelp · Epibiota · Bryozoans · Recruitment · Growth

INTRODUCTION

Habitats are being modified by human activities. Besides direct effects of modified habitats, which change abiotic factors, there are potential indirect effects through interactions with other stressors (Sala et al. 2000, Didham et al. 2007). In particular, alterations of environmental factors in modified habitats can facilitate the introduction of species, which, in turn, may act synergistically affecting ecological processes and, consequently, the function of natural systems (Didham et al. 2007). Concerns about increasing rates of anthropogenic disturbances therefore make it necessary to go beyond the study of ecological patterns to investigate the processes that determine them (Pressey et al. 2007). Understanding such processes is crucial to develop successful strategies for management and conservation.

In coastal waters surrounding cities, natural habitats are often replaced by man-made structures, such as marinas, breakwaters and seawalls. These structures provide novel substrata for hard-bottom organisms but have characteristics that make them intrinsically different from natural habitats. For instance, these structures affect hydrodynamic processes, reducing water flow and wave energy (Floerl & Inglis 2003), usually causing greater sedimentation rates. The presence of piers and wharves also reduces light (Glasby 1999). As a result, the structure of assemblages in these modified habitats differs from those
on natural substrata (e.g. Connell & Glasby 1999, Bulleri 2003). For example, many non-indigenous species become established on artificial structures that act as entry points facilitating their spread across other habitats (e.g. Glasby et al. 2007, Dafforn et al. 2009). Recent studies have shown that the addition of these structures also affects epibiota on habitat-forming species occurring in these habitats (e.g. macroalgae; Marzinelli et al. 2009).

Stands of kelps provide habitat and other resources (e.g. food) to many species. In Sydney Harbour, the kelp Ecklonia radiata occurs on pier pilings in marinas and on natural rocky reefs. Covers of epibiota on laminae of kelps on pilings are, however, much greater than on kelps on adjacent reefs (Marzinelli et al. 2009). Greater covers of epibiota can have detrimental effects on kelps and, in turn, on the organisms that use kelps as a resource (Wahl & Hay 1995, Levin et al. 2002, Wahl 2008). For example, greater covers of epibiota can have negative effects on kelps by reducing the area available for photosynthesis (Cancino et al. 1987), which can be enhanced by the reduced levels of light under piers (Hepburn et al. 2006), thereby altering primary productivity. Epibiota can also make kelps more susceptible to drag (D’Antonio 1985); fronds with greater covers of epibiota are more likely to be torn apart by currents and waves (Lambert et al. 1992). This can contribute to the loss of stands of kelps (Steneck et al. 2002, Schiel et al. 2004, Coleman et al. 2008) and subsequent changes in the structure and function of benthic assemblages (Levin et al. 2002). In Sydney, most of the observed difference in covers of epibiota is due to the non-indigenous bryozoan Membranipora membranacea (Hewitt et al. 2004), which can cover 30 to 50% of the laminae of kelps on pilings, but only covers ~5% of laminae of kelps on reefs (Marzinelli 2009). Greater covers of bryozoans on pilings appear to be influenced by properties of the primary habitat (pier pilings) and not due to indirect modifications of kelps by the habitat (Marzinelli et al. 2009). The piers supported by the pilings reduce light significantly, and pilings support much smaller abundances of the sea urchin Holopneustes purpurascens, which lives on the canopy of kelps and feeds on its laminae. Manipulative experiments showed that greater shade and smaller numbers of urchins on pilings are the main factors influencing covers of bryozoans (Marzinelli et al. 2011). These factors are likely to affect several ecological processes, such as recruitment and growth of the bryozoans.

The population structure of many species is often determined by recruitment processes (Underwood & Fairweather 1989), which encompass the production and fertilization of gametes, dispersal and survival of larvae, and settlement and survival of individuals until census (Underwood 1979, Keough & Downes 1982, Todd 1998, Keough & Swearer 2007). Recruitment of marine invertebrates is affected by several factors. For example, shade increases larval settlement because larvae of many marine invertebrates exhibit negative phototaxis (Thorson 1964, Rodriguez et al. 1993). Also, differential mortality of early settlers due to biological disturbance or predation can affect recruitment (Hunt & Scheibling 1997). Thus, greater shade and smaller post-settlement mortality in pilings (due to low abundances of urchins) could, therefore, increase recruitment of bryozoans to kelps.

Another ecological process that can determine the structure of populations is growth. In assemblages of sessile invertebrates, for instance, growth rates of species often determine the outcome of competitive interactions (Nandakumar & Tanaka 1994). Also, the feeding performance of many filter-feeders is affected by the size of the colonies (Pratt 2005). Artificial structures affect waves and currents, which might increase availability of food through passive hydrodynamic processes. This, in turn, may lead to greater growth of bryozoans on pilings (Okamura 1992), resulting in greater covers on laminae of kelps.

The aim of this study was therefore to determine the extent to which these 2 ecological processes are being affected by the modified habitat formed by pilings. We examined the models that differences in covers of Membranipora membranacea between kelps on pier pilings and rocky reefs are caused by: (1) greater recruitment to kelps on pilings, (2) faster growth of colonies on kelps on pilings, or (3) a combination of both processes. Because covers of bryozoans are much greater on kelps on pilings (Marzinelli et al. 2009), the space available for recruitment is significantly smaller than on kelps on reefs. Therefore, to provide similar space for recruitment in the 2 habitats, we transplanted kelps from reefs to pilings. By doing this, we avoided possible confounding due to the space available to recruit and other potential sources of confounding, such as presence of greater numbers of conspecifics. This methodology could be employed because we first showed that the pattern of differences in covers of bryozoans was not due to differences in the type of kelps that grow in each habitat (Marzinelli et al. 2009). Model 1 leads to the prediction that numbers of colonies and percentage covers of bryozoans recruiting to laminae of kelps transplanted from reefs to pilings will be greater than...
on those on reefs. Model 2 leads to the prediction that growth of colonies of *M. membranacea* will be faster on pilings than on reefs. Finally, model 3 leads to the prediction that recruitment to kelps and growth rates of colonies will be greater on pilings than on reefs.

**MATERIALS AND METHODS**

**Recruitment**

*Ecklonia radiata* without epibiota on their laminae were experimentally transplanted from rocky reefs to pilings at 3 places in Sydney Harbour, Australia (Fig. 1): Chowder Bay (CB; 33° 50' S, 151° 15' E) and 2 sites (~100 m apart) at Balmoral Beach (BB; 33° 49' S, 151° 15' E; see Marzinelli et al. 2009). Each location has natural rocky reefs (sandstone platforms) and wharves with wooden pilings and decking built over soft sediments (~2 to 6 m depth). The distance between reefs and pilings was ~300 m at each place.

Fifteen kelps were collected haphazardly (~3 m apart) from reefs at the same depth (1 to 2 m) by carefully detaching the holdfast from the substratum. Five individuals from each place were then randomly assigned to each of 3 treatments: (1) individuals Transplanted (TP) to pilings; (2) Disturbed (D) individuals, which were disturbed in the same manner as required for transplantation, but returned to their original site on reefs; (3) Translocated (TL) individuals, which were similarly disturbed, but taken to a different site on reefs. In addition, 5 Undisturbed (U) individuals were marked *in situ* at each place on reefs.

Disturbed and Translocated treatments allowed distinguishing among the possible influences of disturbances due to the procedure involved in the transplantation of kelps or moving the kelps to an unfamiliar place in the same habitat, respectively (Chapman 1986). Transplanted, Disturbed and Translocated kelps were held in place on the reefs or pilings by a transplanting device (Marzinelli et al. 2009). A cable tie was placed around the stipe and 3 cable ties were threaded through the former and cable tied onto a square piece of plastic mesh (361 cm²) placed underneath the holdfast, keeping the kelp in an upright position. On rocky reefs, the mesh was attached to the holdfast of other individuals in the kelp-stand using 3 cable ties. A rope was placed around pilings and the devices were cable-tied to this rope. On pilings, 5 kelps from each place were marked *in situ* and assigned to a fifth treatment, Undisturbed pilings (P), to estimate recruitment to kelps with great covers of epibiota and to control for possible differences in recruitment between kelps transplanted to pilings and those already on pilings. Most of the kelps on pilings had bryozoans on their laminae. On each kelp, 1 secondary lamina without epibiota was marked using cable ties (around the lamina) and assigned to the P treatment. Comparing recruitment to secondary lamina of undisturbed kelps on pilings with recruitment to the whole thallus of undisturbed kelps on reefs would be confounded because the space available for recruitment (and the numbers and/or covers of conspecifics) differs between the 2 habitats. Thus, to determine differences in recruitment between habitats, recruitment to undisturbed kelps on reefs was compared with recruitment to kelps transplanted from reefs to pilings (and appropriate procedural controls, see above), which have similar covers of bryozoans and space availability (see 'Introduction').

This experiment was performed 3 times to test for consistency over time: 16 October to 22 November 2007, 1 March to 14 April 2008 and 11 December 2008 to 7 January 2009. Experiments lasted for 4 to 6 wk to allow recruitment and growth of epibiota. Longer experiments were not necessary because...
when kelps were transplanted from reefs to pilings and vice versa in previous experiments, percentage covers became similar to that on the opposite habitat after 1 mo (Marzinelli et al. 2009). Each kelp was sampled by taking 5 random photographs of the laminae (covers do not seem to differ among sections or surfaces of laminae at these locations; Marzinelli 2009) with an Olympus 7 megapixel underwater digital camera, which allowed sampling colonies larger than ~0.05 cm². A frame was mounted to the camera to ensure that each photo was always the same distance from the substratum (6 cm) and covered the same area (4 × 5 cm), which provided the greatest possible resolution and precision. Photographs were analysed on a computer screen; the colonies of bryozoans were counted (including those partially within the quadrat) and the percentage cover estimated, using 30 regularly-spaced points over the entire photograph. Taxa that were in the quadrat but not under these points were assigned an arbitrary cover of 0.5%. Whenever possible, animals were identified to species.

**Growth**

A pilot study was done in May 2007 at each of 2 sites on pilings or reefs at BB to test the methodology. *Ecklonia radiata* with colonies of *Membranipora membranacea* occupying areas <16 cm² were sampled haphazardly every ~3 m by snorkelling. Ten colonies, each from different individuals of *E. radiata*, were marked *in situ* by attaching a circular plastic earring (0.75 to 1 cm diameter) to the lamina ~2 cm from each colony. Tags were also attached to the stipe of each kelp with cable ties. At the start of the experiment and after 1 and 7 d, each colony was photographed as above (‘Materials and methods: Recruitment’). Photographs were analysed using the Image-J computer software and the area of each colony was calculated for each time of sampling. A scale bar drawn on the frame was used to calibrate measurements. Size-specific growth rate per unit time (1 wk) of *M. membranacea* colonies were calculated as (area_end − area_start)/(area_start × time).

Results of the pilot study indicated that the methodology used produced reliable data on growth, so that hypotheses could be tested. In reefs, ~50% of the tags were lost. Thus, many colonies sampled at the start could not be identified with reliability after 7 d, so these were not included in the analyses. More colonies were therefore tagged on the later experiments. Because there was no significant variability in growth of bryozoans between sites in each habitat (see Results: Growth), subsequent experiments were conducted at a greater spatial scale (Location; see below).

This experiment was then repeated 3 times: 1 to 8 May, 15 to 22 May, and 19 to 27 December 2008. Colonies of *Membranipora membranacea* were sampled on kelps on pilings or rocky reefs at CB and BB. At each location, kelps with *M. membranacea* colonies with an area <16 cm² were tagged and sampled as described above. Fifty colonies on kelps on reefs (n = 1 to 2 colonies per kelp) and 25 colonies on kelps on pilings (n = 1 to 3 colonies per kelp) were marked *in situ* and sampled at the start of the experiments and after 7 d as described above.

**Analyses of data**

Analyses of variance (ANOVA) were used to examine differences among means. When Cochran’s test for heterogeneity of variances was significant and no transformation was possible, the analysis of variance was still done because it is robust to departures from the assumptions when sample size is large (Underwood 1981, 1997). Non-significant interactions with p > 0.25 were pooled (Underwood 1997). Where significant interaction terms were detected, Student-Newman-Keuls (SNK) comparisons of means were used to determine which treatments differed (Underwood 1981, 1997). All tests were done using GMAV 5 statistical software for Windows (Underwood & Chapman 1997). Analyses are explained in detail in each Table.

**RESULTS**

**Recruitment**

Few species recruited to kelps during the experiments. *Membranipora membranacea* recruited to kelps on pilings and reefs, representing >90% of the colonies in each habitat. *Bugula stolonifera* recruited to kelps in each habitat, whereas *Tubulipora* sp. and *Disporella novaehollandiae* recruited only on reefs.

In 2007, several kelps were lost during the experiment, so only 3 were used in analyses. Recruitment of *Membranipora membranacea* did not differ among treatments (Fig. 2, Table 1a). In the second site at BB, however, recruitment only occurred on kelps transplanted to pilings (Fig. 2). Total recruitment was greater at CB (Fig. 2, Table 1). Covers of *M. membranacea* differed among treatments across places.
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(Table 1b). In CB, covers on kelps transplanted to pilings (TP) were significantly greater than on kelps assigned to any of the other treatments (U, D, TL, P; Fig. 2). In the first site at BB, covers on undisturbed kelps on pilings (P) were significantly greater than in all the other treatments, despite a trend of greater covers on kelps transplanted to pilings (TP) than on undisturbed or controls on reefs (U, D, TL; Fig. 2). In the second site at BB, there were no significant differences among treatments. Recruitment, however, only occurred to kelps transplanted to pilings (TP; Fig. 2).

In 2008 and 2009, only 2 colonies of bryozoans recruited to undisturbed kelps on pilings (P), so this treatment was not included in the analyses. In 2008, there was significantly greater recruitment of *Membranipora membranacea* to kelps transplanted to pilings (TP) than to undisturbed kelps on reefs (U) or controls (D, TL) at the 2 sites in BB (Fig. 3, Table 2). Covers of *M. membranacea* were significantly greater on kelps transplanted to pilings (TP) than on undisturbed kelps on reefs (U) or controls (D, TL) at CB and the first site in BB (Fig. 3).

In 2009, recruitment of *Membranipora membranacea* was significantly greater on kelps transplanted to pilings (TP) than on undisturbed kelps on reefs (U) or controls (D, TL) at CB and the first site in BB (Fig. 3).

In 2009, recruitment of *Membranipora membranacea* was significantly greater to kelps transplanted to pilings (TP) than to undisturbed kelps on reefs (U) or controls (D, TL) at the 3 places the experiment was done (Fig. 4; Table 3). The area of colonies that recruited to kelp on pilings or reefs was small (<4 cm²), so percentage cover was not estimated.

### Growth

Growth rates were estimated by calculating the difference between final and initial colony areas and dividing this difference by initial area of each colony. Data were ln-transformed prior to analyses.
In the pilot study at 2 sites in BB in May 2007, growth of *Membranipora membranacea* after 1 d was in most cases <10% of the growth after 7 d (Table 4). Because the measurements used to estimate measurement error were small (<10%), this measurement error was not considered in the other analyses. Growth rates (means ± SE) of *M. membranacea* after 7 d were greater on kelps on pilings (1.3 ± 0.2 wk⁻¹) than on reefs (0.6 ± 0.1 wk⁻¹). This was consistent across sites (Table 5).

During the experiments in 2008, ~50% tags were lost. Because there was no significant variability among kelps (ANOVA, $F_{16,20} = 1.05$, $p > 0.45$), this factor was not considered in further analyses (i.e. the replicates were the colonies). In each experiment in May 2008, mean growth rate of *Membranipora membranacea* colonies was significantly greater on kelps on pilings than on those on reefs at BB (Fig. 5, Table 6). On the contrary, at CB, growth rates were significantly greater on reefs than on pilings in the first experiment. No significant differences were found in the second experiment (Fig. 5; Table 6). In December 2008, growth rates were significantly greater on pilings than on reefs at BB and CB (Fig. 5, Table 7).

### DISCUSSION

In this system, pier-pilings affected recruitment and growth (Model 3) of the non-indigenous bryozoan *Membranipora membranacea* on kelps.

Recruitment, as for many other ecological processes, is very variable in space and time. This variability can influence the structure of populations. For instance, if recruitment is consistently sparse, it can limit populations (see Underwood & Fairweather 1989). It is also more difficult to make predictions about the effect of disturbances (here, human modification of habitat) when variability is great (Keough & Swearer 2007). Recruitment to kelps
varied among places and times, although, when different, it was always greater on pilings than on reefs. Recruitment of *Membranipora membranacea* to kelps appears, therefore, to be affected by the pier-pilings.

One of the earliest components of recruitment is the production of larvae. On developed shorelines, the reproductive output of adults can be affected by the structure of the habitat. For example, seawalls sustained smaller limpets than natural rocky shores; limpets on seawalls produced fewer and smaller egg masses than do those on rocky shores (Moreira et al. 2006). Differences in reproductive outputs of bryozoans between pilings and reefs could explain differences in recruitment between these habitats. If, instead, the production of larvae is similar on pilings and reefs, differences in recruitment could be explained by the arrival of propagules (Sousa 1984, Keough 1986, Underwood & Fairweather 1989). Arrival of propagules to a patch can be influenced by the distance between the source of propagules (i.e. adult individuals producing larvae) and the patch (e.g. Sousa 1984). Distances from potential suppliers of larvae may be smaller for pilings than for reefs, possibly because of greater covers of bryozoans and smaller variability in covers among kelps (Marzinelli et al. 2009). These 2 models seem, however, unlikely to explain differences in recruitment between pilings and reefs because these habitats are in close proximity (~300 m apart).

Larvae disperse through the water column where they can be affected by many factors. Predation, for example, can significantly reduce the numbers of larvae (Thorson 1950). Also, waves and currents affect dispersal of larvae. These physical factors not only influence the numbers of propagules arriving to a patch, but also their survival after settlement. The longer larvae spend swimming, the less selective they become in their responses to settle-
larvae, thereby affecting their survival as adults. Also, greater swimming time can reduce the quality of the settlement cues (e.g. Marshall & Keough 2003). This, in turn, may lead to settlement in unsuitable places, e.g. where there is less food or greater predation. Also, greater swimming time can reduce the quality of the larvae, thereby affecting their survival as adults (Marshall et al. 2003).

After larvae arrive at a patch, their settlement can be triggered by physical and/or chemical cues. One of these cues is intensity of light (Thorson 1964). Larvae may also respond to other physical factors, e.g. water-flow (Rodriguez et al. 1993). Chemical cues produced by conspecifics may also trigger settlement (Keough 1984, Raimondi 1988). Cues provided by the pilings or by greater numbers of adult colonies of bryozoans on kelps on pilings could, therefore, influence recruitment to this habitat. The effect of these cues might, however, vary among species (Todd & Keough 1994) and even among individuals of the same species (Raimondi & Keough 1990).

Because initial settlement and recruitment were not distinguished, there could have been equal settlement in the 2 habitats, but greater survival of the settlers on kelps on pilings, perhaps due to increased food. Alternatively, post-settlement mortality could have been greater in reefs due to predation (Hunt & Scheibling 1997). For example, predation of new settlers by fish affected the distribution and abundances of bryozoans (Keough & Downes 1982). Other studies have also shown the effect of post-settlement predation or disturbance of other organisms (e.g. Osman & Whitlatch 2004, but see Sams & Keough 2007 for example of weak effects). Urchins may affect settlers of bryozoans on kelp on reefs through these processes. Recruitment to undisturbed kelps on pilings was often lower than to transplanted kelps from reefs. This could be explained by differences in the availability of space to settle on kelps in these 2 treatments. The area of a secondary lamina is on average 1/10 of the area of the whole frond (Wernberg et al. 2003). Only recruitment to a secondary lamina was recorded for undisturbed kelps on pilings because they were already fouled with bryozoans, whereas recruitment was recorded for the whole frond of transplanted kelps.

Similar differences in recruitment between pilings and reefs were found when recruitment was estimated as percentage covers, although the difference between habitats became greater. This indicates that, after settlement, the growth rates of colonies may have been greater on pilings than on reefs. Several models may explain the observed differences in growth-rates between pilings and reefs. For example, increased food on pilings because of differences in water flow in comparison to reefs may cause differences in growth (Okamura 1992, but see Okamura & Partridge 1999). Also, the presence of neighbour bryozoans could have caused reduced growth on reefs. For example, bryozoans growing close to neighbours grew more slowly than did corresponding isolated colonies (Nandakumar & Tanaka 1994, Okamura & Partridge 1999). This model is, however, unlikely to explain slower growth on reefs because very small numbers of bryozoans occur on kelp in this habitat (Marzinelli et al. 2009) and because none of the colonies measured came in contact with other colonies.

At CB, differences in growth between habitats were variable. Growth was faster on pilings at one time, faster on reefs at another time, or did not differ. A possible explanation for this could be great variability in food supply between habitats (Okamura 1992). Alternatively, growth may have been slower due to the presence of neighbours, although this is unlikely because none of the colonies measured came in contact with other colonies. Several studies have, however, shown that growth is very variable in time and space (e.g. Keough 1986, O’Dea & Okamura 2000), pointing out the necessity of replication (Keough 1986). Although this experiment was done at 2 locations and 3 times, this may be insufficient to test for consistency in the observed patterns.

Further research should focus on studying the link between growth and recruitment of Membranipora membranacea. Faster growth of bryozoans leads to larger colonies, which may survive better than small ones. This, in turn, may affect the production of larvae because larger colonies have more zooids, each

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one of which is capable of producing larvae (Yoshiioka 1982). If the reproductive output of individual zooids in larger colonies is similar to those in smaller ones, larger colonies can release greater numbers of larvae into the water column because they have more zooids (Yoshioka 1982). This could facilitate the dispersal and establishment of this non-indigenous species elsewhere.

This information can be used to minimise the impacts of the replacement of natural habitats by artificial structures. For instance, trying to eliminate non-indigenous epibiota such as Membranipora membranacea to reduce negative effects on kelps and other organisms that inhabit them (Cancino et al. 1987, Wahl & Hay 1995) will prove to be costly and ineffective because their recruitment and growth are enhanced by the artificial structures, and these processes alone can produce rapid and significant increases in epibiont cover. Also, these non-indigenous species are already established on natural reefs. So, designing artificial structures that mimic natural reefs may be the most effective strategy for management. This is particularly relevant for Sydney Harbour, where natural rocky shores are being replaced by artificial structures, such as seawalls. Altering the design of seawalls to resemble natural shores had proved to successfully enhance diversity (Chapman & Blockley 2009). Several physical and biological properties of pilings may explain differences between habitats in recruitment of bryozoans to kelps (see Introduction). Of the factors shown to affect bryozaos on kelps in this system (Marzinelli et al. 2011), shade is likely to affect settlement of larvae because they often respond to reduced amounts of light (Thorson 1964, Rodriguez et al. 1993). For example, experimental shading of primary substra-

Table 7. Membranipora membranacea. Analysis of growth rate on kelp on pilings or rocky reefs during the experiment in December 2008. Habitat (pilings vs rocky reefs) was a fixed factor with 2 levels; Location was a random factor with 2 levels. The replicates were the colonies (n = 15). Cochran’s test (C) was used to test assumptions of homogeneity. Data were ln (x + 1) transformed. SNK tests of means were done where there were interactions: *p < 0.05; ns: p > 0.05

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tum increased recruitment of bryozoans to pilings (Glasby 1999) and other substrata (Ryland 1960). Conversely, greater densities of sea urchins may reduce settlement of larvae in reefs by enfoldong the fronds for attachment or increase post-settlement mortality by grazing on the kelps (Wahl & Hay 1995). Reducing the shade caused by pilings and transplanting urchins to pilings may thus minimize the changes in biodiversity caused by pilings (Marzinelli et al. 2011).

Patterns of biodiversity are generated and maintained by dynamic ecological processes. Human activities are rapidly altering natural habitats and disrupting natural processes, having important consequences on biodiversity (Vitousek et al. 1997). Conservation planning that ignores anthropogenic influences on ecological processes cannot successfully promote the persistence of biodiversity (Pressey et al. 2007). It is therefore necessary to increase our understanding of the effects of modified habitats on ecological patterns and processes to provide information and practical advice for conservation and management. Artificial structures are replacing natural habitats along urbanized shorelines globally. Altering the design of built structures to be better mimics of natural habitats will contribute to the conservation of local biodiversity by preserving natural patterns of abundances and distribution of organisms and the processes that determine them. Further, this can mitigate other potential adverse effects of anthropogenic modification of habitats, e.g. by reducing the ‘invisibility’ of natural systems and increasing their resilience and stability.

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