Variation in the structure of subtidal landscapes in the NW Mediterranean Sea

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ABSTRACT: Compounded effects of climate change and local human activities are threatening marine biodiversity worldwide. At a regional scale (10s to 100s km), comparisons among areas characterized by the prevalence of different human activities provide an insight into the effects of anthropogenic disturbances at multiple levels of ecological organization (i.e. from landscapes to assemblages). At the landscape scale (1000s m), we hypothesized that patchiness in habitat distribution and proportion of degraded assemblages would increase with increasing levels of disturbance, as a result of the decline of habitat-forming species. In addition, we hypothesized that prevailing human influences would affect the structure and variability of rocky benthic assemblages at smaller spatial scales (10s cm to 10s m). An extensive survey encompassing areas subjected to different human influences (i.e. from urbanized to protected areas) was carried out along the coasts of Tuscany (NW Mediterranean Sea). Seagrass beds and macroalgal canopy stands were the dominant habitats in relatively pristine areas, while macroalgal turfs and dead rhizomes of Posidonia oceanica were the most extended habitats in urbanized areas. In general, habitat fragmentation did not vary among areas subjected to different human influences. At a smaller scale (10s cm to 10s m), urbanization favored dominance by opportunistic species and promoted biotic homogenization. Our study shows that regional variations in the composition of landscapes and assemblages can be predicted on the basis of prevailing human activities. Our results also suggest that variations in landscape composition could be an effective descriptor of the effects of multiple human stressors in marine environments.

KEY WORDS: Habitat degradation · Urbanization · Benthic assemblage · Biotic homogenization · Landscape · Fragmentation · Marine protected area · MPA · Mediterranean

INTRODUCTION

Compounded effects of global climate change and local human activities, such as coastal development, organic and inorganic pollution, and overfishing, pose severe threats to marine biodiversity (Gray 1997). Recent efforts to map the spatial distribution of human activities have suggested that the number and severity of overlapping threats can vary locally in coastal areas (Halpern et al. 2008). Thus, prevailing environmental conditions (i.e. climate, geology) and human activities (either negative or positive, such as the establishment of marine protected areas, MPAs) set the regional scenario within which factors operating at finer scales, from the seascapes to the habitat, down to the community, operate to regulate patterns of abundance and distribution of marine organisms. Nonetheless, little effort has been made, over regional scales, to assess variation at different levels of ecological organization among areas characterized by the prevalence of different human activities.

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Shallow rocky reefs support highly diverse and productive assemblages that are among the most susceptible to human influences (Halpern et al. 2007). According to the regional geomorphology, rocky reefs can be found in proximity to the mainland or in off-shore waters (i.e. shoals or islands). Thus, they can be exposed to different environmental conditions (e.g. currents, salinity, riverine inputs) and human pressures that vary in both nature and intensity. For instance, coastal reefs are more likely to be exposed to degraded environmental conditions (i.e. inputs of sediments, nutrients and pollutants, development of coastal infrastructures) than off-shore or island reefs. On the other hand, activities such as trawling, recreational boating, diving, and angling can be widespread or more common on reefs relatively far from major human conglomerates.

At the community scale, human influences can alter the richness and relative abundance of species in benthic assemblages (Claudet & Fraschetti 2010), increase the susceptibility to invasion by introduced species (Bulleri et al. 2010), and reduce food-web complexity (Guidetti 2006). In addition, anthropogenic stressors can either increase (Warwick & Clarke 1993) or reduce (Chapman et al. 1995) the spatial variability of natural communities.

At the landscape scale, human influences can cause the loss or fragmentation of habitats (Airoldi & Beck 2007). For instance, the regression of canopy-forming macroalgal stands or seagrass beds and the consequent expansion of alternative habitats (e.g. algal turfs or dead rhizomes of *Posidonia oceanica* [hereafter also referred to as matte], respectively) have been documented in urban areas worldwide (Benedetti-Cecchi et al. 2001, Balestri et al. 2004, Mangialajo et al. 2008, Gorman & Connell 2009). Thus, the ultimate effect of the regression of habitat-forming species highly susceptible to altered environmental conditions is often the formation of mosaics of habitats. Several studies have attempted to elucidate the causal mechanisms which promote switches between alternative habitat pairs (i.e. canopy-forming macroalgae versus algal turfs: Bulleri et al. 2002a, Mangialajo et al. 2008; erect macroalgae versus encrusting coralline barrens: Bulleri et al. 2002b, Guidetti 2006; seagrass beds versus dead rhizome matte: Balestri et al. 2004).

Fragmentation and switches to alternative habitats can be driven by physical (e.g. sedimentation, hydrologic alterations, dredging, or anchoring), chemical (eutrophication, inorganic pollution), or biological disturbances (overfishing, biological invasions) that trigger a complex process of habitat degradation or loss and the consequent switch in habitat structure and patch quality (for a comprehensive list of causal mechanisms see Boström et al. 2011). However, despite the growing interest in coastal biogenic habitats like seagrass meadows, salt marshes, and coral reefs, alterations in the structure of landscapes have been rarely investigated in the marine realm (Boström et al. 2011). In particular, no study has, to the best of our knowledge, examined how the relative proportion of different habitats and their spatial configuration (i.e. size and patchiness) vary among areas subjected to different levels of human influence.

Here we explored the effects of multiple human activities at a regional scale, encompassing both an along-shore and an off-shore gradient of disturbance. First, we evaluated how the relative availability and spatial configuration of habitats vary among areas subjected to decreasing levels of human pressure, by selecting study locations in urban and extra-urban settings, off-shore shoals, and in MPAs established off islands. In particular, we hypothesized that areas exposed to higher intensity of human disturbance would show a greater patchiness in habitat mosaics and a greater proportion of degraded types of habitat (e.g. algal turfs or dead *Posidonia oceanica* matte). Second, focusing on rocky bottoms, we assessed the variability in the structure and composition of benthic communities along the human influence gradient. In particular, we tested the hypothesis that under more intense human pressures, species diversity would be lower, and small-scale spatial variability of assemblages would be altered.

**MATERIALS AND METHODS**

**Study locations**

An extensive survey was carried out along the coasts of Tuscany (NW Mediterranean Sea) in summer 2008. According to prevailing human activities, 4 levels of decreasing human pressure were identified: (1) urban reefs, (2) extra-urban reefs, (3) off-shore reefs, and (4) MPAs off islands. Two locations were randomly selected for each of these conditions (Fig. 1).

Urban reefs were sampled in front of the cities of Livorno and Rosignano Solvay, which are located about 25 km apart along the coast of Tuscany and are considered pollution hotspots in European coastal waters (OECD 2002). Livorno has about 180,000 inhabitants and hosts an oil refinery and one of the
The urban locations included in this study are densely populated (Fig. 2B; Livorno 1507.73 people km$^{-2}$, Rosignano Solvay 341.33 people km$^{-2}$), located in proximity to industrial or port areas (Fig. 2A) and exposed from high to maximum levels of organic (Fig. 2C), nutrient (Fig. 2D), and inorganic pollution (Fig. 2E). Extra-urban locations were selected in areas showing similar population densities (Fig. 2B), but lower urbanization, as shown by the proximity to natural areas in the land use map (Fig. 2A). Organic, nutrient, and inorganic pollution indices show lower values in extra-urban compared to urban locations.
Off-shore reefs are exposed from high to medium levels of organic, nutrient, and inorganic pollution (Fig. 2C–E). In contrast, MPAs represent relatively pristine locations, with low population densities (Fig. 2G,L; Capraia 16.91 people km\(^{-2}\), Pianosa 48.71 people km\(^{-2}\)) and the majority of the territory occupied by natural areas (Fig. 2F,K). Organic and nutrient pollution are nearly absent across the islands (Fig. 2H,I,M,N), while inorganic pollution levels are very low (pollution index <0.006; Fig. 2J,O).

**Sampling of habitat structure**

At each location, 8 transects of 30 m length and hundreds of meters apart, were randomly selected within an area of about 1 km\(^2\), at depths varying from 3 to 13 m (Fig. 1). Off islands, transects were selected on the west coast to allow comparisons with mainland locations, in terms of geographical orientation and wave exposure. The relative proportion of different habitats was visually quantified by recording the habitats' linear extent and calculating their percentage cover along each transect. Seven habitat types could be identified according to the presence of dominant habitat formers or prevalent features of the substratum: (1) patches occupied by sand or gravel, (2) barren grounds (species-poor patches dominated by encrusting corallines and deprived of erect macroalgae), (3) pre-coralligenous assemblages (a hard substratum of biogenic origin that is mainly produced by the accumulation of calcareous encrusting algae growing in dim light conditions and hosting diverse macroalgae and invertebrates), (4) seagrass *Posidonia oceanica* meadows, (5) *P. oceanica* dead matte, (6) stands of canopy-forming macroalgae (mostly *Halopithys incurva* and *Cystoseira* spp.), and (7) turf-forming macroalgae (mostly filamentous and articulated coralline forms).
In order to examine patterns of variability in habitat distribution at the landscape scale, we calculated a contagion index (CI; Parresol 2011), which quantifies composition (i.e. number and relative proportions) and configuration (i.e. spatial arrangement and fragmentation) of different habitats along each transect. Each linear transect was divided into 300 cells of 10 cm length (our resolution for habitat mapping), and the CI was calculated as:

$$\sum_{i=1}^{n} \sum_{j=1}^{n} \frac{P_{ij}^2 \times \ln(P_{ij})}{(P_{ij} - 1)}$$  

(1)

where n is the total number of habitat types in a particular transect, $P_{ij}$ is the product between the probability of a cell to be occupied by habitat i ($P_i$) and the probability to be adjacent to a cell occupied by habitat j ($P_j$). This CI is a function of an evenness index; it ranges from 0 to 1 and measures the extent to which patch types are aggregated or interspersed. Greater values of contangion occur in the presence of few large, contiguous patches, whereas many small and scattered patches generate lower values. In addition, we calculated mean patch size for each type of habitat and the global mean patch size among all habitats, in order to examine patterns of variability in habitat fragmentation (Zhang et al. 2011).

In order to account for potentially confounding effects due to substratum heterogeneity, we measured topographic complexity according to the chain method (Luckhurst & Luckhurst 1978). One end of a 10 m long stainless-steel chain was randomly placed along the transect and allowed to sink to the bottom, while released in a straight line. Care was taken to ensure that the chain conformed to the contour of the substratum. The relative difference between the real length of the chain and the measured distance between the 2 ends when conforming to the substratum provides a good estimate of complexity. This quantity was divided by the original length of the chain, in order to obtain values ranging from 0 to 1, with smaller values indicating increasing complexity. Two measures of complexity were taken along each transect and subsequently averaged, to be included in multivariate analysis as a covariate.

**Sampling of rocky reef assemblages**

At each location, a minimum of 5 transects were placed entirely on rocky substrata. Thus, at each location, the structure of rocky benthic assemblages was quantified in 6 quadrats (20 × 15 cm) randomly placed along 5 transects. Five transects were randomly selected at locations where a greater proportion of transects lying on rocky substrata was available. Quadrats were photographed by divers using a digital camera (Canon PowerShot S70) equipped with a waterproof case and a metal spacer (28 cm long) and framer. When present, canopy-forming macroalgae were set aside to take pictures of understory assemblages. The percentage cover of macroalgae and sessile invertebrates was visually estimated on a PC screen. A grid of 25 sub-quadrats was superimposed onto each image, a score from 0 to 4% was given to each taxon in each subquadrat, and the percentage cover was obtained by summing over the entire set of sub-quadrats. Sessile invertebrates, which represented 1.044 ± 0.138% (mean ± SE per plot), were sampled estimating their percentage cover and not the number of individuals, as they were mainly (75%) composed of encrusting and/or colonial organisms (e.g. sponges, bryozoans, didemnids, encrusting vermetids). Organisms could generally be identified to the species level, except for encrusting and filamentous algal forms, which were treated as morphological groups.

**Statistical analyses**

**Habitat structure**

The relative proportion and the mean patch size of each habitat type, the global average patch size and the CI were analyzed by means of 2-factor analyses of variance (ANOVA), including the Condition (fixed, with 4 levels: Urban, Extra-Urban, Off-shore, MPA) and the Location (random, with 2 levels, nested within Condition). Cochran’s C-test was used before each analysis to check for homogeneity of variance, and data were ln(x + 1) transformed when necessary (Underwood 1997). The data were analyzed even when homogeneity of variance was not achieved by transformation, since ANOVA is robust for the departure from this assumption when there are many independent replicates (n = 64) and sizes of samples are equal. When necessary, pooling procedures were applied according to Underwood (1997). Student-Newman-Keuls (SNK) tests were used for a posteriori comparisons of the means. The analysis was also repeated for a second, less conservative CI (CI2) proposed by Parresol (2011), which is calculated as:

$$\sum_{i=1}^{n} \sum_{j=1}^{n} [P_{ij} \times \ln(P_{ij})]$$  

(2)

The results did not differ from those obtained for CI and are therefore not reported.
Structure of rocky reef assemblages

Differences in the structure of rocky benthic assemblages among areas subjected to different human influences were tested by means of a permutational multivariate analysis of variance (PERMANOVA; Anderson 2001) performed on a Bray-Curtis dissimilarity matrix of untransformed data. The model included 3 factors: Condition (fixed, with 4 levels: Urban, Extra-Urban, Off-shore, MPA), Location (random, nested within Condition, with 2 levels) and Transect (random, nested within Location, with 5 levels), and substratum heterogeneity as a covariate. The same analysis was repeated also including the depth of each plot as a covariate. Since depth did not significantly contribute to explain variability in assemblage structure, the analysis is not reported. Permutational analysis of multivariate dispersion (PERMDISP, Anderson 2006) and pairwise a posteriori comparisons were performed to evaluate whether differences among conditions were due to diverse dispersion and, hence, heterogeneity of assemblages. Multivariate patterns were displayed by plotting centroids of each transect in a 2-dimensional non-metric multidimensional scaling (nMDS) plot.

Percent cover of the most abundant species or morphological groups (classified according to Steneck & Dethier 1994), species richness (number of taxa) and evenness (calculated as the reciprocal of Simpson’s index/number of species) of assemblages were analyzed using a 3-factor ANOVA, with the same design described for PERMANOVA.

RESULTS

Habitat structure

Stands of canopy-forming macroalgae were nearly absent in urban, extra-urban, and off-shore reefs, while they represented the dominant habitat in MPAs (Table 1, Fig. 3f). In accordance, the availability of habitats formed by turfing macroalgae was significantly lower in MPAs than off the mainland (i.e. urban and extra-urban) or on off-shore reefs (Table 1, Fig. 3g). The ANOVA showed significant differences between conditions in the availability of *Posidonia oceanica* beds and dead matte, but the SNK tests could not rank the means (Table 1, Fig. 3D,E). There was, however, a trend for a greater extension of *P. oceanica* beds and dead matte on off-shore and urban reefs, respectively (Fig. 3d,e).

The relative proportions of coralligenous, barren, and sandy habitats (Table 1, Fig. 3a−c), as well as the spatial configuration of habitats (estimated through the CI) and the global mean patch size did not vary among conditions (Tables 1 & 2). In contrast, the degree of fragmentation of coralligenous, dead matte, and macroalgal canopy habitats (estimated as the mean patch size) significantly varied among conditions (Table 2). Patches dominated by coralligenous and macroalgal canopy were significantly larger in MPAs than in other conditions (Table 2, Fig. 4), while the SNK test failed to provide an alternative to the null hypothesis for dead matte. Mean patch size of barrens, algal turfs, *Posidonia oceanica* beds, and sandy habitats did not vary among conditions (Table 2).

Structure of rocky reef assemblages

After accounting for the variability due to topographic complexity, the structure of rocky reef assemblages differed among conditions, locations, and transects (Table 3, Fig. 5). In addition, the PERMDISP analysis showed significant differences among conditions in terms of dispersion and, hence, in small-scale variability of assemblages (Table 3). Variability among replicates was greatest in MPAs and progressively decreased along the gradient of human influence, with assemblages of urban areas being more homogeneous (Fig. 5).

There was no effect of the level of human influence on the diversity or evenness of benthic assemblages (Table 4). In accordance to patterns of habitat availability, *Cystoseira* spp. dominated space in MPAs, while they were scarcely abundant in off-shore areas and absent off the mainland (Table 4, Fig. 6a). *Peyssonnelia* spp. were more abundant in MPAs than in all other conditions of human influence (Table 4, Fig. 6c). In contrast, the percentage cover of filamentous algae was greater on reefs off the mainland (e.g. urban and extra-urban) than in off-shore or MPA reefs (Fig. 6b). The ANOVA did not detect significant differences in the cover of articulate and encrusting coralline macroalgae (Table 4).

DISCUSSION

Only a few habitat types (coralligenous, dead matte, and canopy stands) varied in terms of habitat fragmentation among areas exposed to different human influences, and there were no significant dif-
ferences in the extent to which habitat patches are aggregated or interspersed. The scarce evidence of fragmentation suggests that the characteristics of prevailing human activities do not necessarily result in smaller and more interspersed patches of habitats, as demonstrated by other studies (Boström et al. 2011). This could be due to the fact that heterogeneity in the spatial arrangement of habitat patches can be a natural feature of shallow reefs. Habitat diversity and fragmentation can, in fact, be generated by natural processes like grazing or the hydrodynamic regime (Bell et al. 1999 and references therein) or by physical features of the bottom, such as topographic complexity or the alternation of sedimentary and rocky substrata.

Rather, human influence could determine gradual changes in habitat composition when moving from urban through extra-urban areas to off-shore banks, while generating a more abrupt shift between these 3 continental areas and islands. For instance, habitats formed by algal canopies (Cystoseira spp.) were found nearly exclusively on island MPA reefs, and, not surprisingly, mean patch size of this habitat type was larger there than at other locations where canopy-formers were sparsely present. Loss of Cystoseira spp. canopies is strongly correlated with urbanization (Benedetti-Cecchi et al. 2001, Mangialajo et al. 2008, Sales & Ballesteros 2009). The decline of these macroalgae can, in fact, be promoted by eutrophication (Soltan et al. 2001), increase in water turbidity, and sedimentation (Airoldi & Beck 2007).

Factors such as orientation and wave exposure can influence patterns of distribution of Cystoseira spp. (Sales & Ballesteros 2009). However, they likely played only a small role in determining regional variations in the abundance of these macroalgae, since sampling was carried out at locations with similar orientation and hydrodynamic conditions. Indeed, the presence of Cystoseira spp. has been historically documented along the Tuscan coast (Cinelli 1969), suggesting a case of habitat loss. In other regions of the Mediterranean Sea, the genus Cystoseira is still pre-

Fig. 3. Mean ± SE percentage cover of habitats (n = 64) in the different conditions of human influence. Letters above the columns illustrate the ranking of the means as from the SNK tests. Y-axis scale varies among panels. MPA: marine protected area.
sent in moderately or scarcely urbanized areas (Man-

gialajo et al. 2008). The decline of Cystoseira spp. has

been far more dramatic in our study region, where

these species have completely disappeared from

mainland reefs, and a few scattered individuals can

be occasionally found on off-shore reefs that, similar
to the mainland reefs, are characterized by poor

water quality. The canopy-forming macroalga Halopit-

hyis incurva is relatively resistant to anthropogenic

disturbance (e.g. sedimentation, Bulleri et al. 2010),

but densities were too low to form a distinct habitat

on urban and extra-urban reefs.

When comparing the relative proportions of habitat
types on rocky bottoms (i.e. excluding transects lying
entirely on Posidonia oceanica beds or sandy areas),
stands of algal turfs represented the dominant habi-
tat in coastal areas, with covers up to 90% in urban,

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Deviation from centroid

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Denominator = 0.345 Lo(C) + 0.655 Tr(Lo(C)) + 0.007 Res,
Denominator = 0.981 Lo(C) + 0.019 Tr(Lo(C))
Denominator = Tr(Lo(C))

Table 1. Analysis of variance (PERMANOVA), permutational analysis of multivariate dispersion (PERMDISP), and a posteriori comparisons between different conditions of human influence. MPA: marine protected area

Table 2. Analysis of variance of the contagion index (CI) and of the percentage cover of habitats between different conditions of human influence. Matte data were ln(x + 1) transformed. *p < 0.05, **p < 0.01, ***p < 0.001, Den = denominator

Table 3. Permutational multivariate analysis of variance (PERMANOVA), permutational analysis of multivariate dispersion (PERMDISP), and a posteriori comparisons between different conditions of human influence on benthic assemblages of rocky reefs, after accounting for substratum heterogeneity as a covariate. MPA: marine protected area
extra-urban, and off-shore reefs. In contrast, they represented only a small proportion of habitat (<30% cover) in off-island MPAs. Dominance of algal turfs is positively correlated with poor water quality due to coastal runoff, increasing sedimentation, and nutrient loads (Gorgula & Connell 2004). Our results suggest that land-based human activities have the potential to alter the structure of algal assemblages also on reefs several kilometers off the coast.

There is a global trend for a decline and loss of seagrass beds along continental coasts, as a consequence of mechanical damage (e.g. trawling or anchoring) and poor water quality (Duarte 2002, Airoldi & Beck 2007). Seagrass bed regression and disappearance generally occur over temporal scales of tens to hundreds of years (Airoldi & Beck 2007) and can hardly be documented by short-term studies. However, a considerable decline of this habitat along the coast of Tuscany emerges clearly from the comparison of maps of *Posidonia oceanica* beds produced in 1990 and 2009 (Padovani 2009). This would suggest that the smaller extension of seagrass beds on mainland than off-shore or island reefs, documented by this study, would not reflect natural variation in the distribution of this habitat-forming plant, but rather variations in the intensity of human disturbance across the region.

In addition, our results suggest that the loss of seagrass meadows and the increase in the extent of dead matte patches do not necessarily occur under the same type and intensity of human pressures. For instance, off-shore banks are characterized by extensive *Posidonia oceanica* beds, being probably less influenced by land-based human activities than coastal reefs. However, greater cover and larger average patch size of dead matte in these areas compared to extra-urban areas likely reflect detrimental effects of recreational boating and artisanal fishing activities (e.g. anchoring, entanglement of nets) that can be more severe here than on coastal reefs.

In contrast to our expectation, the distribution of encrusting coralline barrens did not vary according to prevailing human influences. Barren habitats are common in overfished areas, where a reduced predatory pressure causes an increase in urchin density and, ultimately, overconsumption of erect macroalgae (Guidetti 2006). Therefore, we expected a lower extent of barren habitats within island MPAs, although previous studies did not detect a direct correlation between barren cover and protection regime at these sites (e.g. Micheli et al. 2005). The lack of shift towards canopy-dominated habitats has been recognized as a common phenomenon in poorly enforced protected areas (Guidetti et al. 2008). At all our study locations, patches of encrusting corallines represented a small fraction of total habitat (average cover <5%) and presented a small average size (0.96 ± 0.12 m). Thus, variation in natural processes, such as wave exposure (Micheli et al. 2005) and recruitment rates (Behrens & Lafferty 2004), might be

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**Fig. 4.** Mean ± SE patch size of habitats (n = 64) in the different conditions of human influence. Letters above the columns illustrate the ranking of the means as from the SNK tests. MPA: marine protected area

**Fig. 5.** Non-metric multidimensional scaling plot of benthic assemblages on rocky reefs representing centroids for each transect across different conditions of human influence. △: urban reefs, ■: extra-urban reefs, ○: off-shore reefs, ▼: marine protected areas
more important than prevailing human influences in determining the extent of coralline barrens.

Finally, the extent of coralligenous formations did not vary among areas exposed to different human influences. This is in accordance with the findings of Piazzi et al. (2004), who found no differences in coralligenous assemblages among coastal, off-shore, and island reefs at depths greater than those encompassed by our study. These patterns are unexpected, since coralligenous assemblages have been shown to be susceptible to several kinds of human disturbances (i.e. nutrient enrichment: Hong 1983; sediment deposition: Balata et al. 2005; trawling: Palanques et al. 2001; recreational diving: Coma et al. 2004).

Coralligenous habitats were, however, more fragmented in coastal areas and off-shore reefs than off islands. At shallow depths like those encompassed by our study, coralligenous formations are mainly composed of green macroalgae (like Halimeda tuna or Flabellia petiolata) and articulated or encrusting corallinales, with a smaller component of sessile invertebrates (on average <1.5% per plot) that generally become dominant at greater depths (Balasteros 2006). Human disturbances could further favor the dominance of macroalgae given their greater tolerance to stress in comparison to sessile invertebrates (Airoldi & Virgilio 1998, Benedetti-Cecchi et al. 2001).

In general, the structure of rocky benthic communities markedly differed among areas subjected to different human influences. Species more susceptible to the degradation of environmental conditions (e.g. Cystoseira spp. and Peyssonnelia spp.) were dominant off islands (Soltan et al. 2001). In contrast, filamentous algae flourished in coastal areas, likely by virtue of their ability to conquer and retain space under enhanced sediment and nutrient loads (Airoldi & Virgilio 1998, Gorgula & Connell 2004). Our coarse taxonomical resolution for filamentous species, however, does not allow us to assess whether filamentous algal assemblages differed in species composition along the gradient of disturbances examined. Finally, some algal groups (i.e. articulated and encrusting corallines or sheet-like algae) did not vary along the gradient of disturbances. Articulated and encrusting corallines tolerate stressful conditions (Airoldi & Virgilio 1998).

Small-scale variability was greatest on island reefs and decreased along the gradient of human influence, with assemblages off urban areas displaying the greatest homogeneity. Reduction in small-scale variability has been identified as an outcome of

### Table 4. Analyses of variance on the effects of different conditions of human influence, location, and transect on the number and evenness of taxa and percentage cover of the most abundant taxa. Data on encrusting corallines were ln(+1) transformed. *p < 0.05, **p < 0.01, ***p < 0.001

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>Number of taxa</th>
<th>Evenness</th>
<th>Filamentous</th>
<th>Cystoseira</th>
<th>Encrusting</th>
<th>Articulated</th>
<th>Corallines</th>
<th>Peyssonnelia</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>df</td>
<td>MS</td>
<td>F</td>
<td>df</td>
<td>MS</td>
<td>F</td>
<td>df</td>
<td>MS</td>
</tr>
<tr>
<td>Condition = C</td>
<td>3</td>
<td>60.92</td>
<td>0.42</td>
<td>1</td>
<td>1.61</td>
<td>0.08</td>
<td>1</td>
<td>1.20</td>
</tr>
<tr>
<td>Location = Lo(C)</td>
<td>4</td>
<td>145.19</td>
<td>8.32</td>
<td>8</td>
<td>346</td>
<td>3.57</td>
<td>32</td>
<td>343</td>
</tr>
<tr>
<td>Transect = Tr(Lo(C))</td>
<td>32</td>
<td>17.80</td>
<td>2.48</td>
<td>32</td>
<td>279</td>
<td>0.01</td>
<td>200</td>
<td>284</td>
</tr>
<tr>
<td>Residual = Resc</td>
<td>200</td>
<td>4.89</td>
<td>0.01</td>
<td>1</td>
<td>289</td>
<td>0.17</td>
<td>100</td>
<td>0.67</td>
</tr>
<tr>
<td>Cochran’s test</td>
<td>0.075</td>
<td>0.01</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Denominator for C: *Denominator for Lo(C): **Denominator for Tr(Lo(C))
chronic human influences (Chapman et al. 1995). Such homogenization did not, however, correspond to a reduction in species diversity or evenness. Biotic homogenization can in fact be paralleled by an increase in local diversity as a result of the replacement of vulnerable species by others that are more disturbance-tolerant (Abadie et al. 2011). Vulnerable species tend to be habitat specialists, while disturbance-tolerant species are often habitat generalists, thus resulting in the biotic homogenization of assemblages at a small scale (Abadie et al. 2011).

In summary, our assessment of human impacts at different levels of ecological complexity has disclosed similar responses at the landscape and at the assemblage scale, suggesting that disturbances might favor the shift to degraded assemblages through the displacement of habitat-forming species. Although supported by historical and contemporary literature and by the mapping of habitats and human pressures, our work is purely correlative. Descriptive studies over regional scales have the merit to depict complex patterns of variability along a gradient of multiple human pressures, but it is important to bear in mind that the correlative approach does not allow us to identify cause and effect relationships. Experimental studies can provide insight into the effects of multiple stressors (Crain et al. 2008), but due to logistic constraints, they are often limited to simplified settings (laboratory or mesocosms) and focused on the response of single species (Crain et al. 2008). Even more rarely, the experimental approach has been applied to investigate the effects of habitat loss or fragmentation on biota homogenization (Boström et al. 2011) or ecosystem functioning (Bulling et al. 2008, Godbold et al. 2011).

Previous studies suggested that habitats represent an efficient proxy to quantify biodiversity, especially over broad scales (Ward et al. 1999). Having to choose a set of representative sites for a marine reserve network, Ward et al. (1999) compared investigations at the habitat and assemblage scale in a pristine region. Their findings suggest that habitat categories can efficiently be used to identify high-priority areas deserving protection. Our study supports the conclusions of Ward et al. (1999), showing that focusing on habitat categories can provide reliable estimates of the effects of multiple human influences and can provide information for management strategies over regional scales.

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