

Scaling the influence of topographic heterogeneity on intertidal benthic communities: alternate trajectories mediated by hydrodynamics and shading

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ABSTRACT: Scale-dependent influence of environmental complexity has become a central issue in ecology. We quantified the impact of artificial reefs on community characteristics (biomass, density) and on individual mussel growth, and we tested the relative importance of physical processes (i.e. flow velocity, substratum temperature) as intermediate factors mediating the scale-dependent influence of topographic heterogeneity on benthic communities. Twelve concrete reefs (cylinders) of 3 different sizes (52, 76 and 115 cm) were placed on randomly selected sites along a rocky intertidal platform. The area around each reef and 4 control sites were divided into 24 sampling cells (6 orientation and 4 distance categories). Hydrodynamic patterns around reefs and control sites were determined using the dissolution of plaster cylinders. Flow velocity was simulated around reefs using a finite-element hydrodynamic model. Substratum temperature was also measured. The biomass and density of benthic community adjacent to the reefs was sampled using 10 × 10 cm quadrats before and 1 yr after installation. Growth of individual *Mytilus edulis* attached to experimental panels was measured. A flow index revealed a strong scale-dependent gradient of decreasing water motion intensity with distance from the reefs, and the hydrodynamic model showed a reduction of flow velocity on the downstream side of large reefs. Substratum temperature was lower closer to reefs, and shaded areas increased with reef size. Maximum *M. edulis* biomass was around large reefs, while the biomasses of other dominant species were not positively influenced by reef size. Biomass and density patterns close to the reefs were significant only around large reefs, with the downstream side having the lowest *M. edulis* biomass. Growth of *M. edulis* decreased significantly with distance away from the reefs. Biomass patterns were best explained by the flow velocity around the large reefs ($R^2 = 0.27$), while mussel growth was best correlated with substratum temperature close to the medium reefs ($R^2 = 0.66$). Our study shows that the spatial structure of the benthic community studied and its scaling with topographic heterogeneity depends on dominant mediating physical factors (i.e. hydrodynamic processes or substratum temperature).

KEY WORDS: Benthic community · Spatial scale · Intertidal · Environmental complexity · Finite element model

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INTRODUCTION

A central issue in ecology is spatio-temporal scaling of community structure and dynamics (Wiens 1989,

Levin et al. 1997). The goal of spatial ecology is to determine how space and spatial scales influence population and community dynamics (Tilman & Kareiva 1997). Theoretical studies have suggested that internal (biotic) properties of individuals and populations interact to produce spatio-temporal complexity in homogeneous environments (Deutschman et al. 1993, Bascompte & Solé 1995). Potentially, environmental

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complexity interacts with biotic processes and influences spatial patterns (Roughgarden 1974, McLaughlin & Roughgarden 1992, Pascual & Caswell 1997). In marine ecosystems, many empirical studies have determined correlation scales between environmental and biological patterns (Seuront et al. 1996, Archambault et al. 1998, Legendre et al. 1997, Schneider et al. 1997, Blanchard & Bourget 1999), but few have quantified scale-dependent processes and determined scaling rules to link environmental complexity to spatial structure in benthic communities. We experimentally quantified scale-dependent influences of meter-scale topographic heterogeneity on biomass, density and spatial structure in benthic communities. We have defined the relative importance of physical (flow velocity, substratum temperature) and biological (growth, aggregation) processes involved in a cascade of events leading to spatial structure.

Topography, measured by substratum elevation above the zero level, describes landscapes at any given point in space. Topography influences community structure in intertidal habitats by creating desiccation stress gradients (Norton 1974) and in terrestrial ecosystems by temperature and precipitation gradients (Bartha et al. 1995, Hofer et al. 1999). Topographic heterogeneity, the scale-dependent measure of topographic variability, also provides refuges against harsh physical (wind blow-down [Boose et al. 1994], fire disturbance [Takaoka & Sasa 1996], desiccation [Garrity 1984], ice scouring [Bergeron & Bourget 1986]) and biotic (competition and predation [Wetthey & Walters 1986, Gosselin & Bourget 1989]) conditions. In marine ecosystems different ecological processes have been linked to topographic heterogeneity over a wide range of spatial scales from micron (Bourget 1988, Le Tourneux & Bourget 1988, Thompson et al. 1996) to kilometer (Wolanski & Hammer 1988, Archambault & Bourget 1996) scales. In intertidal rocky shores, topography can influence the spatial structure of environmental variables, which in turn affect biological processes during both low and high tidal periods. Through the measurement of height above zero level, topography can be linked to the immersion period and desiccation stress. Topographic heterogeneity results in specific hydrodynamic patterns during high tide and shading and air flow patterns during low tide. At the meter scale, topographic heterogeneity from boulders has a scale-dependent influence on hydrodynamic patterns and benthic fish recruitment (Breitburg et al. 1995), as well as soft-bottom (Cusson & Bourget 1997) and rocky-bottom (Guichard & Bourget 1998) intertidal communities.

Mussel aggregation influences food depletion and growth of individuals. Mussels within mussel beds have lower growth rates than solitary individuals

(Okamura 1986, Svane & Ompi 1993). Aggregated mussels also have lower body temperatures than solitary individuals during emersion (Helmuth 1998), thus suggesting that aggregation affects desiccation stress during low tide. By driving both flow velocity and shading patterns, topographic heterogeneity could therefore interact with density-dependent physical processes that influence mussels. Scaling the influence of topographic heterogeneity on benthic communities would depend on the dominant mediating factors (i.e. shading or hydrodynamics) involved in this cascade of events.

We experimentally manipulated topographic scales by adding topographic heterogeneity by way of different-sized concrete cylinder reefs in order to measure scale-dependent patterns of flow velocity, substratum temperature, abundance (biomass and density) and of *Mytilus edulis* growth. We more precisely tested the following hypotheses: (1) Reef size influences the change in dominant species biomass between sampling years. Reef size and position around reefs influence (2) hydrodynamics (flow velocity and water motion index), (3) substratum temperature, (4) the biomass and density of dominant species, and (5) *M. edulis* shell and mass growth. We also explored how the relative importance of hydrodynamics and shading as mediating factors explaining the influence of reefs on biological patterns changes with reef size. Our results support most of these hypotheses and establish further evidence of the scaling of biological responses to environmental heterogeneity found in intertidal ecosystems.

METHODS

Study area. A 1 km stretch of shore along the south shore of the St. Lawrence Estuary, Sainte-Flavie, Québec, Canada (Fig. 1), was chosen as a study site. The site met 2 major criteria: First, it was located along a relatively straight section of the coastline with no large-scale topographic features such as headlands or bays that would affect the hydrodynamic flow regime. Second, the shore, which was a rocky platform, was weakly inclined, and complex hydrodynamic patterns were less likely to develop. Overall, the Estuary shoreline was oriented along the 240°/60° axis relative to magnetic north. The macrobenthic community was dominated by *Mytilus* spp. and *Fucus* spp. In the Estuary *M. trossulus* and *M. edulis* (hereafter *Mytilus*) are morphologically similar, and the newly settled fucoid species are small and difficult to distinguish to the species level. *Littorina saxatilis* and *Littorina obtusata* are also abundant (Guichard & Bourget 1998). No known predators of the mussel have been observed in

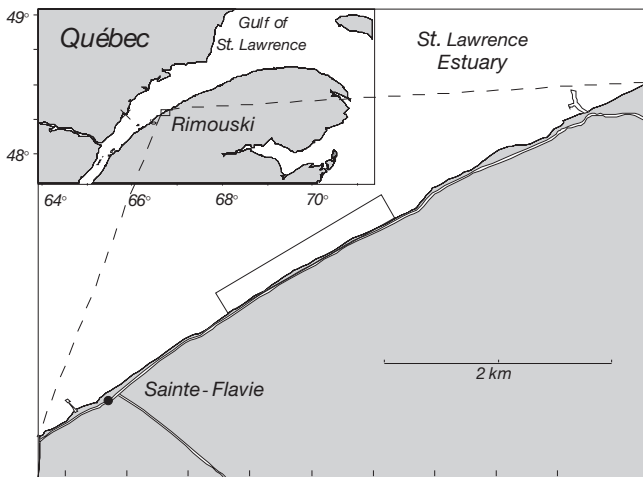


Fig. 1. Study area along the south shore of the St. Lawrence Estuary, Québec, Canada

the mid-intertidal zone of this sub-arctic area. The semi-diurnal tide ranges from about 2 (neap tides) to 4.5 m (spring tides). In spring, ice scouring is an important disturbance factor, and it dislodges most exposed organisms (Archambault & Bourget 1983, Bergeron & Bourget 1986).

Artificial reefs. The experimental design used 12 cylinder-shaped artificial reefs made from reinforced silicium-added concrete which could resist salt water dissolution. The 12 included 4 of each of 3 sizes of cylinders with equal height and diameter: small cylinders were 52 cm, medium cylinders were 76 cm and large cylinders were 115 cm. We preselected thirty 5 × 5 m potential sites in 1994. These sites were free from topographic irregularities such as boulders and crevices larger than 20 cm, and at least 60% of the total area was free from sediments thicker than 2 cm, and from tide pools >2 cm deep. Sixteen of the suitable preselected sites were randomly assigned as either 1 of 4 control sites or a site where 1 of the 12 (3 sizes, 4 replicates) cylinders would be placed. These cylinders (artificial reefs) were installed on 29 June 1994. As we aimed to test the influence of reefs on benthic community development over the growing season, ice scour was allowed. Ice lifts and transports erratic boulders along the shore (Drapeau 1990). During winter the unattached reefs were displaced by ice and the benthic community, unprotected by the reefs, was disturbed by ice. Since sites were selected to be completely exposed, ice disturbance was maximum, removing most exposed organisms. Reefs were reinstalled at their earlier position in April 1995 and not further displaced by wave action. At selected sites 24 sampling cells were assigned by dividing the area around the cylinders into six 60° sectors relative to magnetic north and 4 dis-

tance classes from the cylinders (0–40, 40–80, 80–120 and 120–160 cm). At the control sites we structured the sampling around a medium-size circle (76 cm) and obtained the same number of sampling cells.

Flow measurement. Dissolution of plaster cylinders: Plaster cylinders were anchored in the center of each sampling cell (24 cells site⁻¹; n = 384; Fig. 2) over 24 h (2 full semi-diurnal tidal cycles), which gave a relative measure of flow velocity. Plaster mass loss is linearly related to mean water velocity (Komatsu & Kawai 1992) and has been used previously at our study site (Guichard & Bourget 1998). The plaster cylinders (8 cm long × 2.5 cm diam.) were made from plaster of Paris (LepageTM, Brompton, Ontario, Canada). The ends were covered with epoxy, which prevents dissolution; this means that a near-constant area was exposed during the immersion period. Before installing, the cylinders were dried for 21 d at room temperature and then weighed. After retrieval they were again dried for 21 d and reweighed (± 0.005 g). The plaster cylinders were immersed on 15 August 1995. They were kept in place 2 cm above the substratum by threading metal rods through them. These rods were firmly anchored into the intertidal bedrock. Dry mass loss (initial – end weight) was used as a relative index of hydrodynamic parameters (flow velocity and/or turbulence intensity).

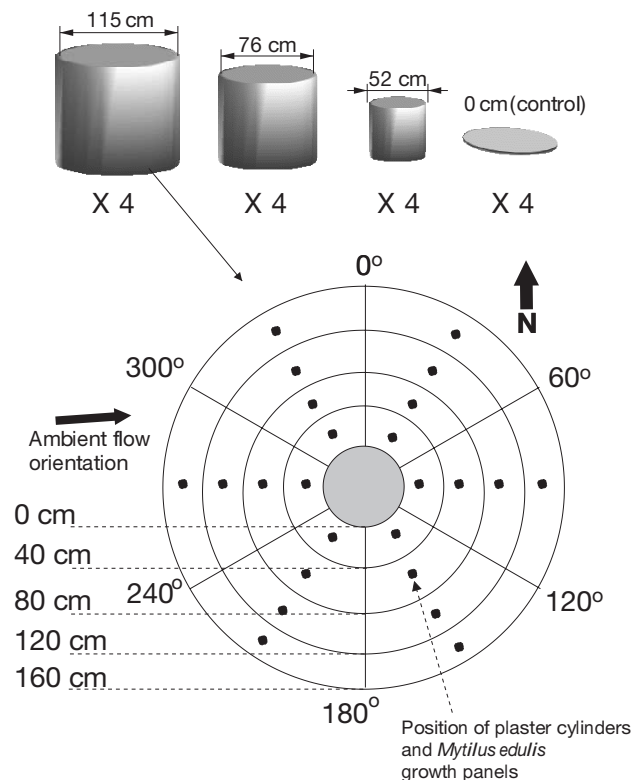


Fig. 2. Schematic of the sampling design

Current velocity measurements: The flow at the scale of the study area was measured using 1 S4 current meter moored at 30 cm from the bottom at one end of the 1 km long study area, and 2 Marsh-McBirney electromagnetic current meters (one at 10 cm and the other at 60 cm from the bottom) moored at the other end of the study area. The current meters were moored from 14 July to 11 August 1995. Dataloggers recorded the average of 60 velocity measurements for the Marsh-McBirney meters and the average of 120 velocity measurements measured over 1 min by each probe, at 10 min intervals, when water depth was greater than 50 cm.

Finite-element hydrodynamic model: Flow around cylinders on an intertidal inclined platform was simulated using a 2D finite-element hydrodynamic model. The model consisted of building three 200×300 m grids each corresponding to an inclined and regular intertidal platform that included a cylinder at its center. The cylinder size was set to equal one of our artificial reefs (52, 76 or 115 cm). Grids were defined by triangular elements. On each element, the 2 horizontal vectors of flow velocity and water level were computed at 6 nodes (corners and center of each side of the element) at each time step (1 min). The model uses 2D (3D integrated over the water column) Navier-Stokes equations for incompressible flows (Robert & Hamed 1995, Robert et al. 1998). The equations express momentum conservation, mass conservation, and boundary conditions for flow velocity and water level, and the solution of the non-linear system was approximated using the Newton-Raphson variational approximation method (Hadjji & Dhatt 1997). While building the grid, elements had variable sizes, and spatial resolution increased near the cylinder up to half the cylinder size. We ran the simulation over 1 tidal cycle for each cylinder size and retained output from one time step where ambient flow velocity matched current meter data (0.05 m s^{-1}). Flow orientation was then adjusted to also match current meter data. We resampled simulation data according to our experimental design (Fig. 2) to obtain mean flow velocity in each experimental sampling cell.

The model does not incorporate wind effect or wave action and reflects current meter data for water depth >0.5 m, as opposed to the plaster cylinders which were immersed for 2 tidal cycles and submitted to wave action in shallow water.

Substratum temperature. An infrared surface thermometer (Everest Interscience, Fullerton, CA, USA) was used to measure bare rock substratum temperature ($\pm 0.1^\circ\text{C}$) in the center of each sampling cell at 1 randomly selected cylinder of each size and 1 control site. The substratum and the probe were not exposed to the sun during measurements. The temperature was

measured 6 times between 25 and 30 August 1997, at the end of low tide during the day. The time of low tide varied from 07:55 to 19:40 h meaning that the data integrated changes in position of the sun. Throughout the sampling period, ambient air temperature was near the average growing season temperature, with daily maxima between 16 and 22°C .

Community characteristics. The benthic community was sampled 2 times at each experimental site: the first one between 5 and 20 July 1994 just after the artificial reefs were installed, and the second time the next year, between 17 July and 13 August 1995. The sampling periods were as short as possible to minimize temporal change bias.

In the 1994 samples, one 10×10 cm quadrat was positioned using random polar coordinates in each sampling cell at each experimental site, except for cells in the 80–120 cm distance category ($n = 288$). All nonencrusting organisms (invertebrates and algae) were removed from each quadrat. The encrusting species could not be efficiently collected but the percentage cover was visually estimated. Samples were kept frozen at -18°C until analysis. In the laboratory the samples were sieved using 1 mm mesh sieves and all sessile or sedentary individuals were sorted and identified to either the species, or the lowest taxonomic level possible. All organisms were counted and the blotted mass of all species in each quadrat was measured using a Mettler 360 balance (± 0.005 g).

In 1995, we followed a modified sampling procedure. Specifically we sampled 4 quadrats in each sampling cell of all distance categories ($N = 1101$ due to loss of data). Random coordinates which overlapped 1994 quadrat positions were excluded. To test the influence of artificial reefs on body size structure in benthic species, we randomly selected 1 quadrat within each sampling cell of the 60 to 120° category and the opposite 240 to 300° category. The 80–120 cm distance category was excluded. Selected samples were sieved using 3 successive mesh size (1, 2 and 8 mm) and the number of individuals and blotted mass of each species were determined for 3 body size categories (1–2, 2–8 and >8 mm) in each quadrat. The orientation categories chosen for the body size structure analysis were based on flow velocity which is oriented along the $270^\circ/90^\circ$ axis at the study site (Guichard & Bourget 1998).

Mytilus growth. The growth rate of *Mytilus* around artificial reefs was determined by anchoring 4 mussels on a 10×10 cm Vexar panel in the center of each sampling cell, except for the 100 cm distance ($n = 288$). Mussels (17 to 20 mm length) were selected from an intertidal site 3 km from our study site. Initial shell length (± 0.1 mm) and blotted mass (± 0.005 g) were recorded. One mussel valve was glued using epoxy to a corner of a Vexar panel. The panels were placed

through metal rods anchored onto the rock bottom as for plaster cylinders. The mussels were set out between 5 June and 21 August 1995. At the end of the experiment the mussels were remeasured for changes in length and blotted mass ($N = 1033$ after mortality and loss of individuals from growth panels).

Data analysis. The first null hypothesis tested was that reef size would not affect the changes in dominant group (i.e. *Mytilus*, *Fucus* sp., *Littorina obtusata* and *L. saxatilis*) biomass between years. The hypothesis was tested using a mixed factorial ANOVA model (see Tables 3 & 4) and analysed using the SAS statistical package (SAS Institute 1988). We compared the changes between the means of the 1995 samples from each cell and the corresponding quadrat sampled in 1994. The model included among-boulder variability [reef(reef size)] as the error term which tested reef size effect. The same ANOVA model was used for abundance data of dominant species sampled in 1995 (see 'Community characteristics').

Other null hypotheses tested were that reef size, orientation and distance would not affect plaster cylinder dry mass loss, *Mytilus* shell and mass growth or the biomass of dominant species. A mixed factorial ANOVA model (see Tables 1, 5, 6 & 7) that included among reef variability [reef(reef size)] as an error term which tested reef size effect, and within reef variability [orientation \times distance \times reef(reef size)] which tested the effect of other main treatments. Residuals were visually examined, and square root transformations were applied to *Mytilus* biomass and growth (shell and mass) and to *Fucus* biomass to satisfy homoscedasticity and normality requirements of the model.

Temperature was measured around 1 reef of each size. This means that substratum temperature effects were tested using an unreplicated factorial ANOVA model that was missing both among and within reef variance estimates. Thus reef size effect was confounded with site effect for these analyses.

Table 1. ANOVA showing the effect of reef size (RSize), orientation (Ort), distance (Dst), and among reef variability [R(RSize)] on velocity flow index (plaster erosion rate, mg s^{-1})

Source of variation	df	MS	F	p
RSize	3	1.33×10^{-6}	0.096	0.9605
R(RSize)	12	1.409×10^{-5}	47.059	0.0001
Ort	5	1.981×10^{-6}	6.614	0.0001
Dst	3	1.352×10^{-5}	45.130	0.0001
RSize \times Ort	15	9.241×10^{-7}	3.086	0.0001
RSize \times Dst	9	2.536	8.468	0.0001
Ort \times Dst	15	1.062	3.5471	0.0001
RSize \times Ort \times Dst	45	3.584	1.197	0.1966
Error	256	3.30×10^{-7}		
Corrected total	363			

The strength of the linear relationships between physical (i.e. flow velocity, flow index, substratum temperature) and biological (i.e. biomass of dominant species, growth of *Mytilus*) variables as a function of reef size was tested using multivariate regression models. Least-square multivariate regression was used to fit each biological variable to the 3 physical parameters for each reef size category. The contribution of each physical variable to the total fit was examined using squared semi-partial correlation coefficients giving the percentage of variability of the dependent variable explained by each physical variable entered into the model.

RESULTS

Hydrodynamics

Flow regime

The mean velocity of current meter flow at the western end of the study site was 0.05 m s^{-1} at 10 cm from the bottom and 0.111 m s^{-1} at 60 cm from the bottom. The current mean orientation relative to magnetic north was 94.0° at 10 cm and 67.6° at 60 cm. At the eastern end of the study site, the mean flow velocity was 0.464 m s^{-1} and mean orientation was 81.2° at 30 cm from the bottom. Flow reversal, common in the St. Lawrence Estuary during a tidal cycle, was never recorded at the study site from June to August 1995. Flow was always oriented eastward, approximately parallel to the shore, and usually varied between 60 and 100° relative to magnetic north. These data agree with a hydrodynamic model of the whole Estuary, showing a permanent gyre circulating across our study site.

Flow index

The rate of plaster loss (flow index; mg s^{-1}) was significantly influenced by reef size interacting with orientation and distance from reefs (Table 1). The smallest reef induced a significantly lower flow velocity at 100 cm away from the reef than at 20 cm (Fig. 3B). The 76 cm reef had an effect at 60 cm and flow remained significantly less than at 20 cm at 100 and 140 cm (Fig. 3C). The 115 cm reef had a greater influence on flow and the flow index continued to drop significantly between 60 and 100 cm from the reef; this lower index was still seen 140 cm from the reefs (Fig. 3D). In summary, the larger the reef, the greater the effect of orientation and distance, indicating that scale affected the amplitude of the flow index patterns. There was a symmetrical flow index pattern around large reefs,

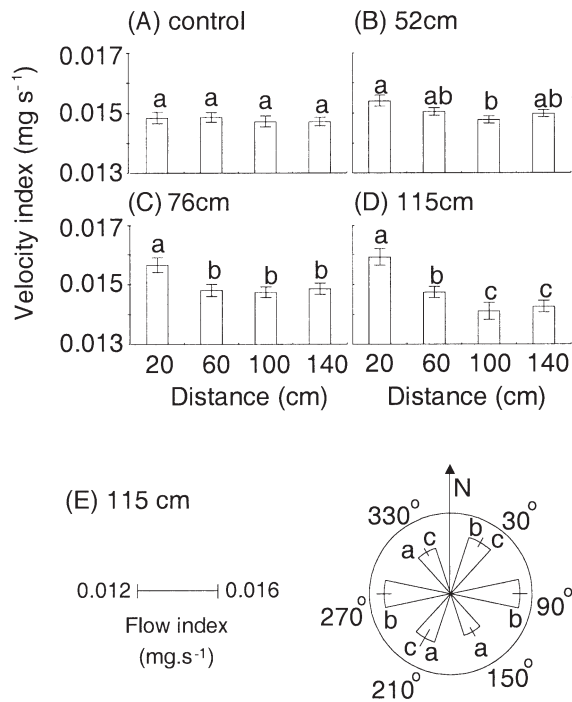


Fig. 3. Flow index (plaster dissolution rate, $\text{mg s}^{-1} \pm \text{SE}$, $N = 24$ [A–D] or $N = 16$ [E]) as a function of distance from the edge of control sites (A) and small (B), medium (C) and large (D) reefs. (E) Flow index at 6 orientations (all distances combined) around large reefs (115 cm). Letters above bars show significant differences among distances (A–D) or orientations (E) according to a least-squares means multiple comparisons test ($\alpha = 0.01$)

with the index significantly higher along the 270°/90° axis compared to along the 330°/150° axis (Fig. 3E). Mean flow orientation as measured by the current meters when the water depth was >0.5 m was 83.4° during the period plaster cylinders were in place. The symmetrical plaster loss pattern around the large reefs suggests that plaster erosion was imposed by a flow along the 330°/150° axis, and this was also the dominant wave direction which created a shoreward flow in shallow water through the interaction with topography at the scale of the study area (F.G. pers. obs.).

Hydrodynamic finite-element model

We simulated a mean flow velocity of 0.067 m s^{-1} over an intertidal platform at time $t = T/4$, where T is the tidal period (12.4 h). The cylinder reefs induced 2 major features. There was a velocity decrease on the downstream side of the reefs (60 to 100° according to current meter data) and a symmetrical flow acceleration on both sides along the main flow axis, near the edge of cylinders (Fig. 4). The model uncovered strong

scaling of these features as a function of reef size. The ratio of accelerated flow velocity over minimum downstream velocity was 1.57 around the small reef, 2.15 around the medium reef and 2.99 around the largest.

Substratum temperature

There was a significant interaction between reef size and the distance from reefs on mean substratum temperature during low tide (Table 2). There was no dis-

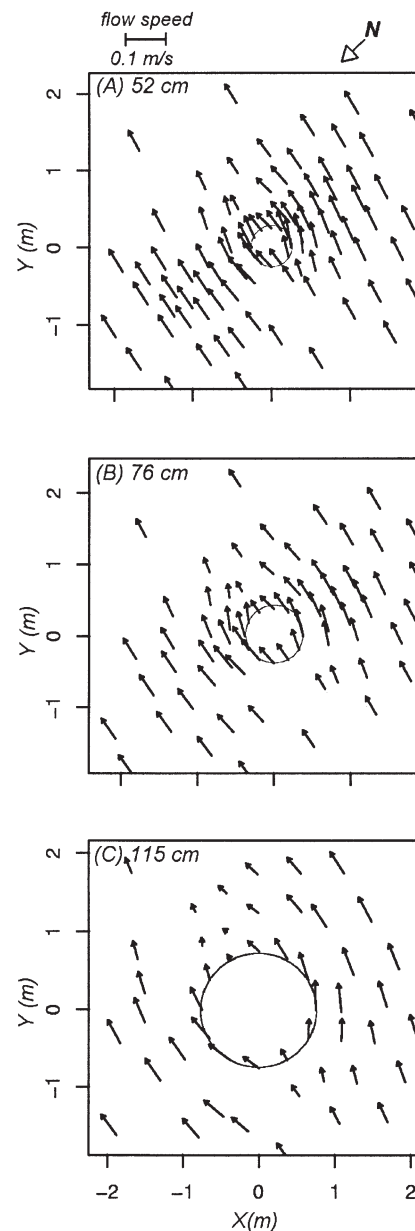


Fig. 4. Finite-element hydrodynamic model outputs showing flow velocity (m s^{-1}) and orientation around a small (52 cm; A), a medium (76 cm; B) and a large (115 cm; C) reef

Table 2. ANOVA showing the effect of reef size (RSize), orientation (Ort), and distance (Dst) on substratum temperature (°C)

Source of variation	df	MS	F	p
RSize	3	5.9345	51.74	0.0001
Ort	5	0.6030	5.26	0.0007
Dst	3	2.1186	18.47	0.0001
RSize × Ort	15	0.2134	1.86	0.0550
RSize × Dst	9	0.3320	2.89	0.0087
Ort × Dst	15	0.1210	1.05	0.4220
Error	45	0.1147		
Corrected total	95			

Table 3. ANOVA showing the effect of reef size (RSize), year (Y), orientation (Ort), distance (Dst), and among-replicate reef variability [R(RSize)] on the biomass (g) of *Mytilus*

Source of variation	df	MS	F	p
RSize	3	70.4851	3.8673	0.0411
R(RSize)	11	18.2561	3.6606	0.0001
Y	1	402.4029	80.6886	0.0001
Ort	5	4.2947	0.8612	0.5073
Dst	2	3.0386	0.6093	0.5443
RSize × Y	3	1.8911	0.3792	0.7681
RSize × Ort	15	9.5032	1.9055	0.0214
RSize × Dst	6	3.0725	0.6161	0.7175
Y × Ort	5	6.1043	1.2240	0.2971
Y × Dst	2	1.4777	0.2963	0.7437
Ort × Dst	10	2.0481	0.4107	0.9414
RSize × Y × Ort	15	6.3652	1.2763	0.2139
RSize × Y × Dst	6	7.2397	1.4517	0.1938
RSize × Ort × Dst	30	6.2526	1.2538	0.1723
Y × Ort × Dst	10	3.5755	0.7169	0.7086
RSize × Y × Ort × Dst	30	4.1869	0.8395	0.7116
Error	380	4.9871		
Total	534			

tance effect on substratum temperature at control sites (Fig. 5A) but temperature was significantly lower at the first distance for the small and medium reefs compared to other distances (Fig. 5B,C). Around large reefs, temperature decreased from the farthest to the closest distance categories (Fig. 5D). In contrast to flow index patterns, the spatial limits but not the amplitude of the temperature gradient were influenced by reef size, despite some variability in amplitude among reef sizes and unreplicated reef size treatment.

Community characteristics

There were significant annual biomass differences for all dominant species. Reef size significantly interacted with annual changes in the biomass of *Littorina saxatilis* and *L. obtusata* but *Mytilus* was independently affected between years and by reef size. Spatial

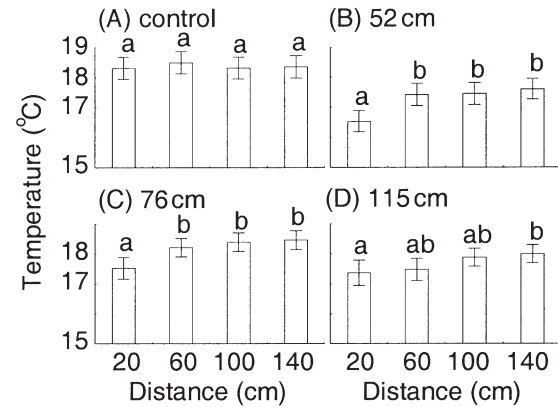


Fig. 5. Substratum temperature (°C ± SE, N = 6) as a function of distance from the edge of a control site (A) and a small (52 cm; B), a medium (76 cm; C) and a large (115 cm; D) reef. Different letters above bars show significant differences between distance categories according to a least-squares means multiple comparisons test ($\alpha = 0.01$)

patterns of *Mytilus* and *L. saxatilis* biomass and density around large reefs were significant and depended on body size. There was also a significant *Fucus* biomass distance gradient around large reefs.

Impact of reef size

There were significant year and scale effects on *Mytilus* biomass with no interaction (Table 3). Biomass was significantly higher in 1995 than in 1994 and higher around larger reefs compared to controls (Fig. 6A). *Mytilus* biomass was the dominant around large reefs in 1995 and appeared to be responsible for both significant effects (Fig. 6A).

Littorina saxatilis biomass was significantly influenced by interactions between year and scale (Table 4). There was an increase in *L. saxatilis* biomass between 1994 and 1995 at control sites, but this interannual difference did not hold and was progressively reversed with increasing reef size (Fig. 6B). Around large reefs, *L. saxatilis* biomass was less in 1995 than in 1994.

There was a significant interaction between year and reef size on *Littorina obtusata* biomass (Table 4). While maximum biomass was observed around medium-size reefs in 1994, biomass was at a minimum and significantly lower around medium-size reefs compared to control sites a year later (Fig. 6C).

Spatial patterns of biomass and density

In 1995, differences in spatial patterns of biomass and density were significant only around large reefs

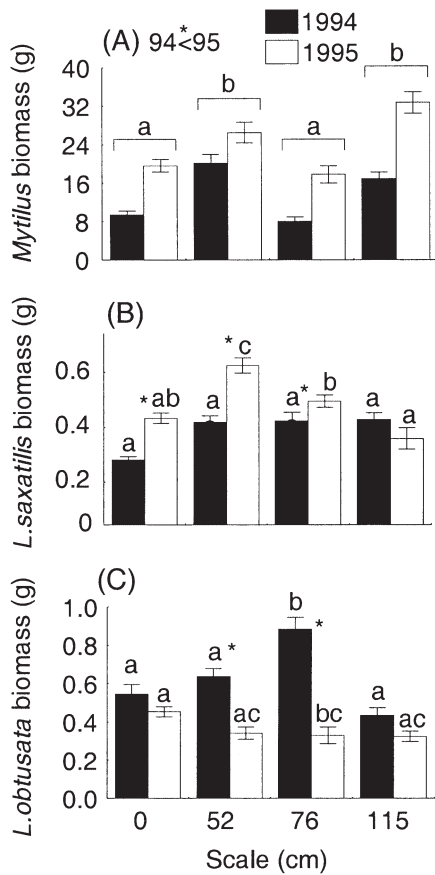


Fig. 6. Biomass ($g \pm SE$) of dominant species before (1994; black bars) and 1 yr after (1995; white bars) installation of reefs of different sizes (Scale). Different letters above bars show significant differences among scale categories and stars represent significant differences between years according to a least-squares means multiple comparisons test ($\alpha = 0.01$)

and were symmetrical along the $270^\circ/90^\circ$ axis. There was a significant interaction between scale and orientation on *Mytilus* biomass (Table 5), which was significantly less between 60° and 120° compared to other orientations around large reefs (Fig. 7A). The density of individuals >2 mm was linked to this pattern (Fig. 8A,B). Large (>2 mm) *Mytilus* density around the large reefs was less between 60° and 120° compared to between 240° and 300° ; however, this pattern was significant for individuals between 2 and 8 mm, but not for individuals >8 mm (Table 6, Fig. 8A,B). *Littorina saxatilis* biomass distribution was similar with the 0° to 60° and 120° to 180° orientation categories, having lower values than orientations between 240° and 270° (Table 5, Fig. 7B). *L. saxatilis* density, however, was significantly greater around medium cylinders, but with no oriented density pattern (Table 6, Fig. 8C). There was a significant interaction between reef size and distance from reefs on *Fucus* biomass (Table 5).

Fucus biomass increased with distance from large reefs (Fig. 7C) and was significantly lower around reefs compared to controls (Fig. 7D). Reef size had a significant effect on *L. obtusata* density (Table 6), and *L. obtusata* was significantly less abundant around cylinders compared to control sites (Fig. 8D).

Growth of *Mytilus*

There was a significant influence of distance from the reefs on both shell length and weight growth (Table 7, Fig. 9A). There was no significant influence of reef size on growth, but shell growth at 20 cm was 1.36 times greater than growth at 140 cm on control

Table 4. ANOVA showing the effect of reef size (RSize), year (Y), orientation (Ort), distance (Dst), and among-replicate reef variability [R(RSize)] on the biomass (g) of *Littorina obtusata* and *Littorina saxatilis*

Source of variation	df	MS	F	p
<i>Littorina obtusata</i>				
RSize	3	0.0863	0.1412	0.9332
R(RSize)	11	0.6125	5.0632	0.0001
Y	1	1.9976	16.5126	0.0001
Ort	5	0.1063	0.8784	0.4955
Dst	2	0.4164	3.4424	0.0330
RSize \times Y	3	0.8606	7.1139	0.0001
RSize \times Ort	15	0.1036	0.8567	0.6136
RSize \times Dst	6	0.1863	1.5399	0.1639
Y \times Ort	5	0.0510	0.4215	0.8337
Y \times Dst	2	0.0050	0.0410	0.9598
Ort \times Dst	10	0.0859	0.7103	0.7149
RSize \times Y \times Ort	15	0.1074	0.8882	0.5778
RSize \times Y \times Dst	6	0.2387	1.9731	0.0685
RSize \times Ort \times Dst	30	0.0730	0.6031	0.9531
Y \times Ort \times Dst	10	0.0898	0.7426	0.6841
RSize \times Y \times Ort \times Dst	30	0.0970	0.8017	0.7643
Error	380	0.1210		
Total	534			
<i>Littorina saxatilis</i>				
RSize	3	0.2965	0.9356	0.4562
R(RSize)	11	0.3175	6.9364	0.0001
Y	1	2.2920	50.0768	0.0001
Ort	5	0.0318	0.6948	0.6276
Dst	2	0.0611	1.3358	0.2642
RSize \times Y	3	0.3169	6.9231	0.0002
RSize \times Ort	15	0.0774	1.6908	0.0504
RSize \times Dst	6	0.0175	0.3829	0.8899
Y \times Ort	5	0.0346	0.7564	0.5818
Y \times Dst	2	0.0645	1.4100	0.2454
Ort \times Dst	10	0.0137	0.2986	0.9813
RSize \times Y \times Ort	15	0.0585	1.2777	0.2131
RSize \times Y \times Dst	6	0.0413	0.9019	0.4934
Csize \times Ort \times Dst	30	0.0344	0.7511	0.8279
Y \times Ort \times Dst	10	0.0387	0.8447	0.5858
RSize \times Y \times Ort \times Dst	30	0.0293	0.6400	0.9309
Error	380	0.0458		
Total	534			

sites, while this ratio was 1.61 around 76 cm reefs (Fig. 9B). Orientation around reefs did not influence *Mytilus* growth.

Flow and biological variables

The regression model outcome indicated that flow index, flow velocity, and substratum temperature explained up to 77.2% of *Mytilus* shell growth around medium reefs and 51.7% of *Mytilus* biomass around large reefs. Relationships between physical and biological variables were highly dependent on reef size. Regression models were not significant at control sites and small reefs; however R² values increased with reef size, with a maximum for medium and large reefs

Table 5. ANOVA showing the effect of reef size (RSize), orientation (Ort), distance (Dst), among-reef [R(RSize)] and within-reef [Ort × Dst × R(RSize)] variability on *Littorina saxatilis*, *Mytilus* and *Fucus* sp. biomass (g)

Source of variation	df	MS	F	p
<i>Littorina saxatilis</i>				
RSize	3	2.6702	3.1852	0.0631
R(RSize)	11	1.0148	5.8665	0.0001
Ort	5	0.1274	0.7373	0.5963
Dst	3	0.0310	0.1798	0.9100
RSize × Ort	15	0.3439	1.9877	0.0172
RSize × Dst	9	0.0416	0.2407	0.9881
Ort × Dst	15	0.1308	0.7542	0.7270
RSize × Ort × Dst	43	0.1018	0.5862	0.9807
Ort × Dst × R(RSize)	209	0.1739	1.2522	0.0175
Error	787	0.1389		
Total	1100			
<i>Mytilus</i>				
RSize	3	45.3995	0.8717	0.4830
R(RSize)	11	63.4382	6.6469	0.0001
Ort	5	25.0089	2.6258	0.0250
Dst	3	5.2260	0.5509	0.6481
RSize × Ort	15	17.6479	1.8482	0.0299
RSize × Dst	9	9.3625	0.9815	0.4562
Ort × Dst	15	9.2819	0.9679	0.4903
RSize × Ort × Dst	43	7.4034	0.7697	0.8460
Ort × Dst × R(RSize)	209	9.6460	1.6492	0.0001
Error	787	5.8491		
Total	1100			
<i>Fucus</i> sp.				
RSize	3	1.9489	0.1066	0.9545
R(RSize)	11	22.7605	17.0283	0.0001
Ort	5	4.4444	3.3335	0.0064
Dst	3	2.5630	1.9316	0.1254
RSize × Ort	15	1.9350	1.4468	0.1278
RSize × Dst	9	3.7760	2.8268	0.0037
Ort × Dst	15	0.8520	0.6337	0.8452
RSize × Ort × Dst	43	1.2580	0.9323	0.5952
Ort × Dst × R(RSize)	209	1.3541	1.9200	0.0001
Error	787	0.7052		
Total	1100			

Table 6. ANOVA showing the effect of reef size (RSize), orientation (Ort), distance (Dst), and among-replicate reef variability [R(RSize)], on *Littorina obtusata*, *Littorina saxatilis* and 2–8 mm *Mytilus* density

Source of variation	df	MS	F	p
<i>Littorina obtusata</i>				
RSize	3	2077.5660	6.2176	0.0100
R(RSize)	11	334.1418	6.6871	0.0001
Ort	1	2.0920	0.0419	0.8386
Dst	2	54.5524	1.0917	0.3428
RSize × Ort	3	61.3855	1.2285	0.3081
RSize × Dst	6	24.8867	0.4981	0.8071
Ort × Dst	2	43.6943	0.8744	0.4228
RSize × Ort × Dst	6	13.9762	0.2797	0.9442
Error	55	49.9679		
Total	89			
<i>Littorina saxatilis</i>				
RSize	3	1785.3535	2.1084	0.1572
R(RSize)	11	846.7687	8.9418	0.0001
Ort	1	0.0062	6.59E-5	0.9936
Dst	2	189.3974	2.0000	0.1451
RSize × Ort	3	268.7808	2.8383	0.0463
RSize × Dst	6	119.3516	1.2603	0.2908
Ort × Dst	2	198.8257	2.0996	0.1322
RSize × Ort × Dst	6	140.2458	1.4810	0.2018
Error	55	94.6982		
Total	89			
<i>Mytilus</i>, 2–8 mm				
RSize	3	274.0545	0.0713	0.9741
R(RSize)	11	3846.2087	7.3532	0.0001
Ort	1	2534.3092	4.8451	0.0319
Dst	2	100.3973	0.1919	0.8259
RSize × Ort	3	1321.6342	2.5267	0.0668
RSize × Dst	6	144.4834	0.2762	0.9458
Ort × Dst	2	571.4271	1.0925	0.3426
RSize × Ort × Dst	6	470.5051	0.8995	0.5020
Error	55	523.0683		
Total	89			

(Fig. 10A,B). For medium-size reefs, where the *Mytilus* growth gradient was maximum (Fig. 9B), temperature alone explained 66.4% of *Mytilus* shell growth variability (Fig. 10B,D). Around large reefs, *Mytilus* biomass variance was best explained by flow velocity (27%; Fig. 10A,C). The regression model was not significant for *Mytilus* density.

DISCUSSION

Topographic heterogeneity in the form of concrete cylinder reefs had a significant and scale-dependent influence on both surrounding physical (hydrodynamics, substratum temperature) and biological (biomass, density, growth) variables. There were significant differences in the scaling of hydrodynamic and substratum temperature patterns as a function of reef size

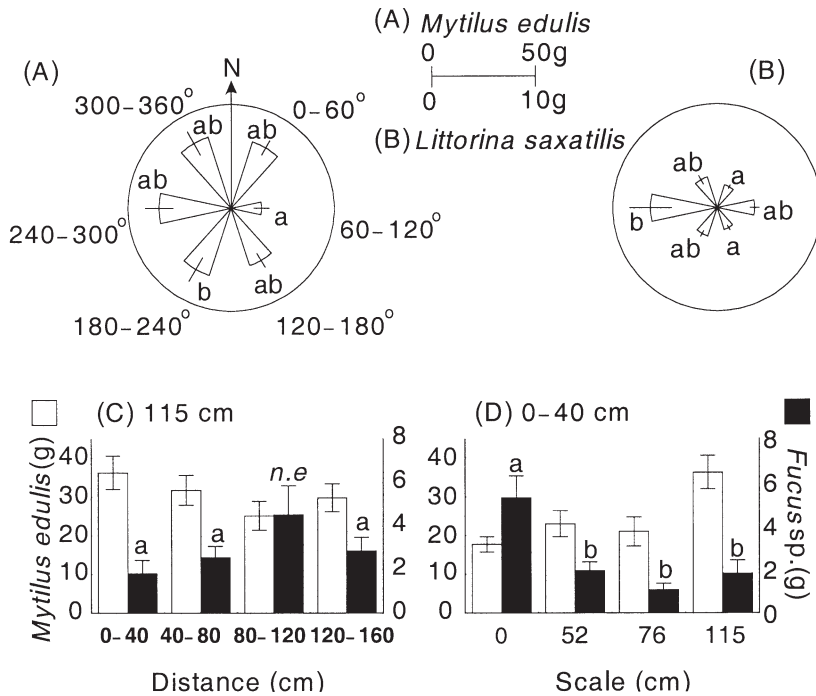


Fig. 7. Biomass (g ± SE) of *Mytilus* (A) and *Littorina saxatilis* (B) as a function of orientation around large (115 cm) reefs. (C,D) *Mytilus* and *Fucus* sp. biomass as a function of distance from the edge of large reefs (C) and within 40 cm from the edge of reefs as a function of their size (D). Different letters above bars show significant differences among orientation (A–B), distance (C) or scale (D) categories according to a least-squares means multiple comparisons test ($\alpha = 0.01$, n.e = nonestimable)

(topographic scale). Reef size also determined the form of biological patterns in the vicinity of the artificial reefs and had a significant effect on the dominant species between sampling years. We found a scale-dependent relationship between abiotic and biotic parameters and identified dominant biological and physical variables that interact in the vicinity of medium and larger reefs.

Scaling topographically generated environmental patterns

The scaling of hydrodynamic patterns created by reefs was investigated theoretically and experimentally (Lighthill 1986). This scaling can best be described using the nondimensional Reynolds number ($Re = DU/\nu$), relating ambient flow velocity (U), reef size (D) and fluid viscosity (ν) to the spatial structure of flow over any object or surface. The Re number can describe the increase in turbulence around a cylinder. When flow velocity and viscosity are held constant, scaling up the topography results in an increase in turbulence levels above the substratum. More precisely,

high Re values ($>10^5$) are associated with a fully turbulent boundary layer and low shear stress over the surface of the cylinder. These properties are responsible for an area of low flow velocity on the downstream side of cylindrical structures (Tritton 1988, Granger 1995). Flow acceleration on 2 sides of a cylinder is linked to increased pressure around the cylinder. From these considerations, we can expect hydrodynamic properties to scale nonlinearly with topographic heterogeneity. Specific hydrodynamic patterns around a cylinder, such as a decrease in flow velocity, depend on scale thresholds, since their strength changes nonlinearly as a function of scale. The predicted result is that the spatial extent and amplitude of hydrodynamic properties will increase with topographic scale.

Plaster erosion patterns around large reefs were characteristic of shoreward-oriented ambient flow, along the 330°/150° axis (Fig. 3), but the current meters showed that flow in >0.5 m water depth was oriented eastward, along the 270°/90° axis. This suggests that hydrodynamic conditions are different close to the bottom

in shallow water compared to >0.5 m water. The flow index determined by plaster erosion integrates all

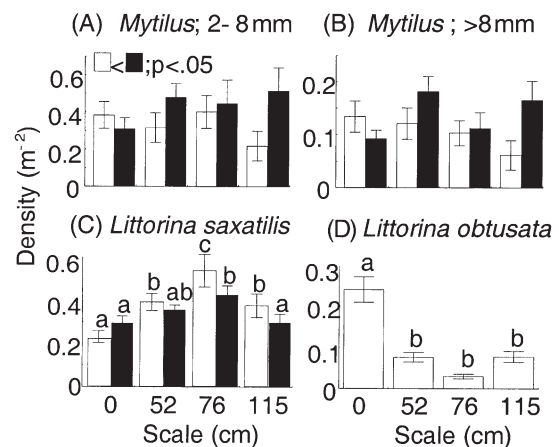


Fig. 8. Density (number of individuals, ±SE) of the dominant species measured upstream (240° to 300°; black bars), or downstream (60° to 120°; white bars) of reefs or at both orientations combined (D) as a function of reef size (Scale; cm). Different letters above bars show significant differences among scale categories according to a least-squares means multiple comparisons test ($\alpha = 0.01$)

Table 7. ANOVA showing the effect of reef size (RSize), orientation (Ort), distance (Dst), among-reef [R(RSize)] and within-reef [Ort × Dst × R(RSize)] variability on *Mytilus* weight (g) and shell (mm) growth

Source of variation	df	MS	F	p
Weight growth				
RSize	3	0.0684	1.6173	0.2373
R(RSize)	12	0.05201	14.6673	0.0001
Ort	5	0.0041	1.4950	0.1924
Dst	2	0.0711	18.7170	0.0001
RSize × Ort	15	0.0042	1.2975	0.2050
RSize × Dst	6	0.0011	0.4664	0.8328
Ort × Dst	10	0.0036	1.2810	0.2423
RSize × Ort × Dst	30	0.0037	1.0804	0.3622
Ort × Dst × R(RSize)	203	0.0033	1.2626	0.0159
Error	746	0.0027		
Total	1032			
Shell growth				
RSize	3	2.2522	1.2044	0.3500
R(RSize)	12	1.8807	19.8200	0.0001
Ort	5	0.1188	1.2533	0.2853
Dst	2	2.1903	23.1246	0.0001
RSize × Ort	15	0.1223	1.2877	0.2113
RSize × Dst	6	0.0721	0.7602	0.6020
Ort × Dst	10	0.1320	1.3913	0.1855
RSize × Ort × Dst	30	0.0883	0.9273	0.5796
Ort × Dst × R(RSize)	203	0.0965	1.4316	0.0004
Error	746	0.0674		
Total	1032			

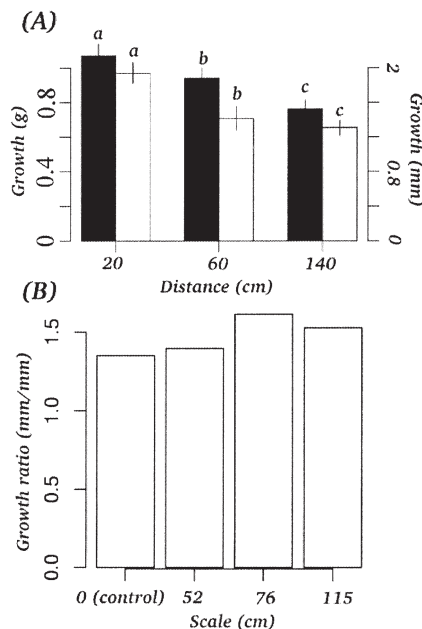


Fig. 9. (A) *Mytilus* weight (g ± SE, black bars) and shell (mm ± SE, white bars) growth as a function of distance from reefs (cm). Different letters above bars reveal significant differences among distances. (B) Ratio of shell growth at 20 cm over shell growth at 140 cm from reefs as a function of reef size. Different letters above bars show significant differences between distance categories according to a Bonferoni multiple comparisons test ($\alpha = 0.01$)

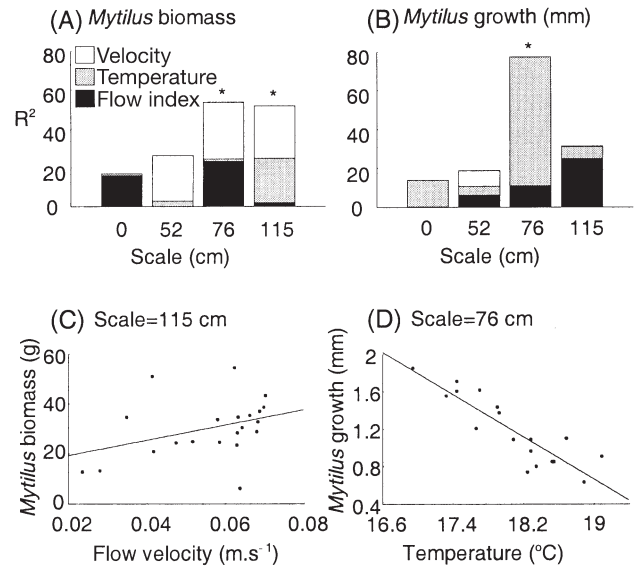


Fig. 10. (A,B) Percentage of variability in *Mytilus* biomass (g; A) and shell growth (mm; B) explained by flow velocity (white bars), substratum temperature (gray bars) and flow index (black bars), represented as the contribution of each independent variable (squared semi-partial correlation coefficients) to the total R^2 of a multivariate regression model. Stars above bars show statistically significant regression models. (C,D) Influence of flow velocity on *Mytilus* biomass around large reefs (C) and of substratum temperature on *Mytilus* shell growth around medium reefs (D)

factors acting on exposed surfaces (turbulence, wave action in shallow water) and may be considered as an index of turbulent shear stress or ‘intensity of water motion’ (Komatsu & Kawai 1992).

The flow index confirmed the expected scaling of both the spatial extent and amplitude of patterns with reef size. The main pattern of water motion intensity around reefs was a distance gradient that suggested greater water motion near the reef’s edge. This gradient grew in amplitude and spatial extent with topographic scale in agreement with the theory and observations from fluid mechanics.

Temperature data integrated over a 6 d temporal scale averaged out short-term differences caused by the interactions between the time of low tide and sun angle. There were temperature gradients around the artificial reefs, but the scaling of the substratum temperature with topographic scales was different from that of hydrodynamics. There was a clear distance gradient in substratum temperature from the reef edges. We found evidence that reef size linearly influenced the shaded area but did not affect the degree of shading, i.e. the temperature difference between shaded and nonshaded areas. Hydrodynamic scaling resulted in the creation of significant oriented patterns around large reefs, but no such scale threshold was observed

for substratum temperature. These scaling differences for different factors show that in the intertidal zone, both high and low tide phases were characterized by environmental cascades of events induced by topographic heterogeneity, each with a specific set of scaling rules.

Individual versus population level response

In intertidal ecosystems, physical factors can control many aspects of benthic communities. During low tide, temperature and desiccation stress are major factors limiting benthic species distribution in the high intertidal (Suchanek 1985). During high tide, hydrodynamic properties can influence key biological processes. For example, flow velocity determines larval and food flux for suspension feeders, thus influencing recruitment and growth (Fréchette & Bourget 1985, Wildish et al. 1987). Turbulent diffusion influences settlement (Harvey et al. 1995) and feeding rates (Nowell & Jumars 1984), while wave force influences maximum body size (Gaylord et al. 1994), fertilization (Denny & Shibata 1989) and post-settlement mortality (Connell 1961). The dominant species at our study site, *Mytilus*, is known to experience mortality from desiccation and high temperature in the upper intertidal (Seed & Suchanek 1992), and individuals usually aggregate to form 3D matrices (mussel beds). Within the matrix, desiccation stress may be less as moisture is retained (Helmuth 1998), but food becomes limiting within the boundary layer and may limit growth (Fréchette et al. 1989). Increasing the flow velocity over a *Mytilus* bed could prevent food depletion by increasing vertical turbulent diffusion within the boundary layer (Butman et al. 1994). Solitary individual mussels could be more limited by desiccation stress than by flow velocity, since they are not protected from desiccation and are less likely to compete for food with congeners than those individuals within a mussel bed.

When the growth rate of individual mussels (17 to 20 mm initial shell length) was measured as a function of position around the artificial reefs, there was a linear decreasing gradient with distance away from reefs which was greatest around the medium reefs. The growth patterns were similar to substratum temperature patterns and were not strongly linked to reef size or influenced by orientation. Moreover, growth rate of individuals with shell length used in our experiment was best explained around medium reefs by the negative influence of substratum temperature (squared semi-partial correlation = 0.78; Fig. 10B,D). Our results thus suggest that substratum temperature was an indication of shading. In turn, shading has a primary influ-

ence on desiccation stress and was a major factor mediating individual response of *Mytilus* to topographic heterogeneity. However, wind velocity patterns are also directly influenced by topographic heterogeneity during low tide and may also influence desiccation stress.

Naturally occurring *Mytilus* biomass was significantly less downstream of the large reefs. There was also a decrease of large mussel density downstream of large reefs. Patterns of abundance for *Mytilus* therefore suggest a topographic scale threshold which results in the appearance of variations in abundance only above a minimum (threshold) topographic scale. Around large reefs, flow velocity best explained *Mytilus* biomass, which was positively influenced by flow velocity as simulated by the finite-element model. In natural populations individuals are mostly found aggregated in mussel beds, and flow velocity, as a determinant of boundary-layer structure, is the dominant mediating factor of mussel response to topographic heterogeneity. The dominant contribution of large individuals (>2 mm) to the biomass patterns suggests that increased size by way of growth was likely the biological link between flow velocity and mussel biomass. Changes in food availability would create the emergence of patterns, although active migration could also influence patterning.

Other abundant species, *Littorina saxatilis* and *Fucus* sp., also showed spatial patterns with a significant influence of orientation or distance, but these patterns were significant only around large reefs and were opposite to the *Mytilus* patterns. Mussel beds are reported to increase colonization by soft substratum fauna that develop within beds (Tsuchiya & Nishihira 1995, Svane & Setyobudiandi 1996), but *Fucus* sp. interacts directly with mussels for primary substratum colonization (McCook & Chapman 1991). Since the dominance pattern between *Mytilus* and *Fucus* sp. depends on environmental conditions (McCook & Chapman 1991, Petraitis 1987), the same topographically induced areas that favor mussel development and dominance could decrease *Fucus* sp. abundance.

Scaling the influence of topography on communities

Increased environmental heterogeneity has often been linked to increased species diversity (Simpson 1964, August 1983). The mechanisms invoked usually involve multiplication of habitat types allowing more species to cohabit without being competitively excluded or by creating local environmental conditions allowing viability of species otherwise unable to survive or reproduce (King & Pimm 1983, Tilman & Pacala 1993). Environmental heterogeneity can also

generate complex biotic-abiotic interactions leading to complex spatial patterns. There is often nonlinear scaling of environmental heterogeneity that influences benthic communities, with specific scale ranges matching key physical or biological processes at the individual, population or community level (Archambault & Bourget 1996, Cummings et al. 1997, Johnson et al. 1997, Schneider et al. 1997). In intertidal habitats, patch and disturbance dynamics result in a positive influence of mussel patch size on species diversity and resilience (Paine & Levin 1981, Kim & Dewreede 1996). Topographic heterogeneity increases benthic invertebrate abundance and recruitment with scales from 10^{-3} to 10^{-1} m (Archambault & Bourget 1996, Lemire & Bourget 1996) and from 10^2 to 10^3 m (Archambault & Bourget 1996, Blanchard & Bourget 1999). Scales of topographic heterogeneity between 0.5 and 2 m also facilitate benthic fish recruitment (Breitburg et al. 1995) and decrease the nearby soft-bottom species richness (Cusson & Bourget 1997). Guichard & Bourget (1998) showed that the presence of natural boulders had a negative effect on *Mytilus* and *Fucus* sp. biomass as a function of boulder size. Boulders are more spherical than cylinders. In the earlier study, areas under boulders, and thus current acceleration zones near the edge of boulders, were not sampled, and this may explain the reported inverse influence of topographic scale on *Mytilus* biomass (Guichard & Bourget 1998). However, there may be a more complex dependence of scaling rules on external forcing factors. Here we found that the biomass of all numerically dominant invertebrate species (*Mytilus*, *Littorina saxatilis* and *L. obtusata*) was affected by artificial reefs. Reef size had a negative effect on all species, but *Mytilus* biomass was maximum around large reefs. Flow acceleration on the reef's sides had a positive influence on *Mytilus* biomass. The overall increase in *Mytilus* around large reefs was only near the edge of reefs, while there was a decrease in *Mytilus* biomass and density of large individuals downstream, where flow velocity was lowest. While both substratum temperature and hydrodynamic patterns were influenced by artificial reefs, we found that flow velocity was the dominant factor mediating the scaling of benthic communities to topographic heterogeneity, while substratum temperature was the dominant factor for solitary individuals (*Mytilus* growth). These alternate trajectories lead to major changes in scaling rules. For *Mytilus* biomass there was a scale threshold (large reefs) and symmetrically oriented patterns along the main flow orientation. However, the scaling of solitary *Mytilus* growth with reef size resulted in patterns that were maximum around medium reefs with no scale threshold and oriented patterns.

Conclusion

Biological scaling rules, changes in biological patterns with topographic scale, depend on observation level, individual versus population, and can have major impacts on pattern formation and population dynamics. Temporal changes in abiotic external forcing, such as increased air temperature, could lead to a switch from hydrodynamics to temperature as the main factor mediating the influence of topographic heterogeneity on benthic communities. Since local scaling rules can influence large-scale population dynamics (Pacala & Levin 1997), switching local scaling rules and resulting local patterns could impact the large-scale dynamics of benthic communities. These results further suggest that the relationship between topographic heterogeneity and benthic communities can take alternate trajectories depending on the dominant mediating physical factors such as hydrodynamics or shading.

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