Rippled scour depressions add ecologically significant heterogeneity to soft-bottom habitats on the continental shelf

Todd R. Hallenbeck1, Rikk G. Kvitek2,*, James Lindholm3

1West Coast Governors Alliance, Office of the Governor, Salem, Oregon 97301, USA
2Seafloor Mapping Laboratory and 3Institute for Applied Marine Ecology, California State University, Monterey Bay Seaside, California 93955, USA

ABSTRACT: Comprehensive mapping of California’s state waters has revealed rippled scour depressions (RSDs) to be abundant and widespread on the inner continental shelf. Ranging from 100s to 1000s of m² in areal extent, RSDs are 30 to 50 cm deep depressions of coarser sediments and longer period bedforms than found on the surrounding seabed. Although RSDs have been physically described on many continental margins, previous studies have not addressed the ecological influence and associated biological communities of RSDs. Here, we test the hypothesis that there are ecologically important differences in the distribution and abundance of benthic fish and invertebrate groups inside and outside RSDs. A small ROV was used to survey 20 RSDs in 3 depth zones (<15 m, 15 to 30 m, and >30 m) within Monterey Bay, California. Density and richness of benthic communities were determined from the recorded video imagery. Sediment grab samples confirmed significantly larger mean grain sizes inside (0.71 mm) than outside (0.22 mm) the RSDs. Overall mean faunal density (fish and invertebrates) was lower inside RSDs in the shallow, intermediate, and deep zones (0.32, 1.61, and 2.17 ind. 10 m⁻² respectively) than outside (0.38, 3.06, and 4.48 ind. 10 m⁻²). Faunal richness was also lower inside RSDs in each depth zone (0.29, 1.03, and 1.43 taxa 10 m⁻²) than outside RSDs (0.29, 1.72, and 2.48 taxa 10 m⁻²). Surprisingly, RSDs did contain significantly more young-of-the-year rockfishes and small flatfishes than adjacent fine sediments, suggesting a possible nursery function for these otherwise depauperate coarse-grained habitats. These results indicate that RSDs can add a significant and previously undescribed level of ecological patchiness to soft sediment communities on the continental shelf.

KEY WORDS: Rippled scour depressions · Continental shelf · Marine spatial planning · ROV · Soft sediment · Benthic ecology · Sorted bedforms · Sebastes

INTRODUCTION

The role of water depth and temperature, as well as substrate type, in the large-scale distribution and abundance of benthic communities has been well described (Warwick & Davies 1977, Langton & Watling 1990, Barry & Dayton 1991, Snelgrove & Butman 1994, Whitman et al. 2004). Typically, the abundance of fishes and invertebrates increases with water depth in the unconsolidated sediment habitats of the continental shelf (Allen & Moore 1996, Kostylev et al. 2001). Within sediment habitats, the distribution of benthic organisms is mediated by a variety of factors, including grain size (Butman et al. 1988, Snelgrove 1999, Brown & Collier 2008), hydrodynamic forces (Aller 1997, Pillay et al. 2007), bedform type (Auster et al. 2003a, Lindholm et al. 2004), and biogenic structure and biotic interaction (Woodin
Rippled scour depressions (RSDs; Fig. 1) have been identified and physically described in small-scale studies on many of the world’s continental shelves (Garnaud et al. 2005, Gutierrez et al. 2005, Lo Iacono & Guillen 2008, Bellec et al. 2010). These elongate shallow depressions (0.3 to 1 m in depth), oriented normal to isobathic contours, are characterized by long crested bedforms (0.5 to 1 m wavelength), coarse sediment (0.3 to 1 mm), and abundant shell hash. RSDs have been found in water depths of 15 to 160 m with reported dimensions of 30 to 250 m wide and 100 to 3000 m long (Cacchione et al. 1984, Bellec et al. 2010). The sediments surrounding RSDs are typically characterized by finer grain size (0.05 to 0.3 mm), shorter period bedforms, and a sharp edge delineating the coarser-grained RSDs (Green et al. 2004, Goff et al. 2005, Diesing et al. 2006). Further, RSDs can be persistent, with their spatial extent and boundaries remaining stable for decades (Diesing et al. 2006).

Although previous studies provide detailed information on the physical characteristics of RSDs at the local scale, regional-scale assessments of RSD distribution and abundance were not possible with the available data. Now, with the completion of the California Seafloor Mapping Project (CSMP), comprehensive high-resolution multibeam sonar maps indicate that RSDs cover nearly as much of the state’s 1200 km long continental shelf as does rocky habitat (5% and 8% respectively), with RSDs representing the most prominent and widespread soft-bottom feature within the low-relief unconsolidated sediments of California’s coastal margin (A. Davis et al. unpubl.).

Hypothesized mechanisms for the formation of RSDs include tidal scour (Bellec et al. 2010), cross shore currents (Cacchione et al. 1984), storm generated downwelling (Garnaud et al. 2005), and along-shore currents (Murray & Thieler 2004). Bottom currents have been measured as high as 60 cm s\(^{-1}\) inside RSDs (Bellec et al. 2010), compared to current speeds of 20 cm s\(^{-1}\) immediately adjacent to RSDs (Green et al. 2004). It is these strong currents that scour away fine sediment and create large bedforms. The bedforms then help to maintain the RSD by disrupting the laminar flow of water and creating a localized turbulence which resuspends fine sediment (Murray & Thieler 2004).

While RSDs have been described from many parts of the world and are abundant on the inner continental shelf, the majority of research has focused on the physical character and geomorphic dynamics of these habitat features (Garnaud et al. 2005, Gutierrez et al. 2005, Lo Iacono & Guillen 2008, Bellec et al. 2010). As a result, little is known about the biological communities associated with or the ecological importance of RSDs. Knowledge of RSD physical properties and the natural histories of many soft-bottom benthic species, however, can be used to make predictions about the likely influence of RSDs on the distribution, diversity, and abundance of those organisms. Many benthic species actively choose to associate with finer sediments, which can facilitate ease of burial for refuge (Tanda 1990, Burke et al. 1991, Gibson & Robb 1992) and typically support higher abundances of infaunal prey items than coarser sediments (Abookire & Norcross 1998, Brown & Collier 2008). Moreover, those factors thought to be responsible for the maintenance of RSDs once formed (higher bottom currents, bedform-induced turbulent flow and subsequent resuspension and transport of fine sediments; Murray & Thieler 2004) could negatively impact the density and diversity of benthic communities inside RSDs by limiting the settlement and recruitment of organisms (Jumars & Nowell 1984, Kaiser & Spencer 1996, Aller 1997) or interfering with the filter-feeding apparatus of some benthic invertebrates (Rhoads & Young 1970, Pillay et al. 2007).
The goal of the present study was to test the general hypothesis that the physical differences between RSDs and surrounding fine sediment habitats will result in predictable differences in the distribution and abundance of benthic species found inside and outside of RSDs. Specifically, we predicted density and richness of benthic organisms would be lower inside RSDs because coarse sediments generally support less diverse and less abundant communities (Abookire & Norcross 1998, Snelgrove 1999, Brown & Collier 2008). If present, these biotic differences between habitats should further increase with depth because the physical differences between habitats will also increase with depth along the disturbance gradient created by hydrodynamic scouring (Allen & Moore 1996, Kostylev et al. 2001, Green et al. 2004). Additionally, we hypothesized that there will be larger fish inside RSDs because fish length has been correlated with bedform size and RSDs create and maintain larger bedforms than the surrounding fine sediment (Gerstner 1998, Auster et al. 2003a).

**MATERIALS AND METHODS**

The locations of 6 distinct RSD fields were identified in southern Monterey Bay (California, USA) from multibeam and side scan sonar data collected in 2001, 2004, and 2010 by the California State University, Monterey Bay, Seafloor Mapping Lab and compiled as part of the CSMP (Fig. 2). A survey cruise was conducted from 6 to 9 July 2010 aboard the RV ‘Macginitie’ to characterize the sediments and benthic communities associated with RSDs and adjacent areas along transects in these 6 study sites (Fig. 2).

**Collection and analysis of ROV video imagery**

Video transects were conducted using a small acoustically tracked remotely operated vehicle (ROV, LBV 200L, Seabotix) equipped with a digital color camera (560 line wide dynamic range, 0.3 lux), external LED lights, and paired 5 cm sizing lasers. The camera was mounted forward-facing at a 45° angle. Each transect was conducted at a speed over ground of 0.5 to 1.0 knots at an altitude of 0.5 to 1.0 m above the seafloor and with the slightly buoyant tether kept off the bottom and trailing behind the advancing ROV. Survey planning and ROV navigation were conducted using Hypack Hydrographic Survey software, enabling both ROV and vessel position and tracks to be displayed in real time over multibeam bathymetry-derived habitat maps showing the location of previously mapped RSD features. Potential disturbance to fish was minimized by keeping the tether behind the ROV and off the seabed and by maintaining a constant slow vehicle speed and altitude. The position of the ROV was recorded relative to the ship’s position every 0.5 s using an ultra-short baseline acoustic tracking system with ranges of 500 m horizontal and 150 m vertical, and range and bearing accuracies of ±0.2 m and ±3° respectively (Micronav, Tritech). The ROV position fixes were converted to real world coordinates as they were logged by the Hypack software.

A total of 30 ROV transects (10 to 55 m water depth) were conducted to collect continuous video imagery along paths spanning the transition from outside RSDs to inside RSDs (Fig. 2). Transects were stratified by depth, with start locations haphazardly selected in the study area based on wind and current direction at the time. Start positions were chosen to ensure that the vessel and ROV could maintain the desired speed and tether orientation along survey lines crossing comparable amounts of habitat inside and outside RSDs within each of 3 depth zones: shallow (<15 m), intermediate (15 to 30 m), and deep (>30 m). Depth zone ranges were selected a priori based on known faunal depth distributions from previous work conducted the same study areas (Kvitek et al. 2008). Transect lengths ranged from 90 to 1260 m depending on vessel drift and the number of RSDs that could be covered along a single survey line (Fig. 2).

A total of 15 h of ROV survey video imagery was recorded, 14 h of which included usable imagery taken within the prescribed distance from the bottom and used in subsequent analyses. Data were extracted from the imagery using a frame-by-frame approach in which successive non-overlapping video frames were treated as individual quadrats. Frames were ignored where turbidity prevented reliable analysis or the ROV was not within 1 m of the seafloor.

All fish and benthic macro invertebrates that occurred within each sampling frame were counted and identified to the lowest taxonomic level possible. Organisms were then binned into one of 4 faunal groups: infaunal invertebrates, suspension feeders, invertebrate predators, and benthic predatory fishes (see Table 1). Because flatfish can generally be sorted into right-eyed and left-eyed families (Kramer et al. 1995) and this characteristic could be reliably distinguished from the video imagery for all individuals, here we assumed all right-eyed flatfish were...
Fig. 2. Map of southern Monterey Bay study area and Big Sur shelf showing rippled scour depressions and the locations of ROV transects and sediment grab samples.
Pleuronectidae spp. and all left-eyed flatfish were Paralichthyidae spp. We do acknowledge that 2 species of Pleuronectidae known to occur in the study area can be either right or left-eyed (California Halibut and Starry Flounder), but given the ease with which these 2 particular species can be visually distinguished using other discernible morphological features, we concluded the right-eyed/left-eyed binning approach to be useful for the vast majority of the remaining flatfish observations.

The counts for suspension feeders, invertebrate predators, and fish groups were further pooled to provide a coarse measure of total faunal abundance and richness. A fourth group analyzed post hoc was intended to provide a coarse measure of total faunal abundance and richness. A fourth group analyzed post hoc was composed of young-of-the-year (YOY) Sebastes spp. and analyzed separately. Parallel laser beams separated by 5 cm were used to measure total length (TL) of each fish observed, and the width of the sampling frame was used to facilitate calculation of total area covered by each transect. Fish smaller than 2 cm or that occurred only partially within the sampling frame were not counted. We strove to minimize any possibility of ‘herding’ fish into RSDs by maintaining constant ROV speed and altitude, with the tether always oriented aft and off the bottom, as recommended by Stoner et al. (2008). We observed no consistency in the direction taken by fish that swam out of the frame of view. Moreover, the YOY rockfish aggregations observed in the RSD fields did not move toward or away from the ROV. Rather, they first came into view as stationary groups on the far edge of the field of view and remained hovering in place within a meter of the bottom as the ROV passed through, with individuals only moving to the side as needed to get out of the direct path of the vehicle.

Percent cover was used to quantify the abundance of 3 infaunal invertebrate groups that were either too numerous to count as individuals within video frames (Dendraster excentricus) or for which indirect measures were required to estimate abundance (burrows for Urechis caupo and Callianassidae and arms protruding from the sediment for Ophiuroids). Because U. caupo and Callianassidae co-occur on the Monterey Bay shelf (Kvitek et al. 2008) and create similar burrow openings that can be difficult to distinguish from each other in video imagery, these 2 taxa were necessarily treated as a single infaunal group. For these 3 groups, percent cover was measured using a 100 point contact grid for sampling each video frame. This grid was superimposed over each sampling frame, and any point that contacted an organism or burrow opening was tallied. The sum of the tallied points for each faunal group was divided by the total number of points (100) to create a percent cover for each sampling frame.

Physical environment parameters including substrate, relief, bedform type, and water depth were recorded for each sampling frame. Habitat type (inside or outside) was determined using combinations of the physical parameters. Criteria used to define inside RSD habitat included coarse sediment substrate, bedform period > 0.5 m, and amplitude > 0.25 m. Outside RSD habitat was defined as unconsolidated fine sediment substrate, bedform period < 0.5 m, and amplitude < 0.25 m. Bedform period and amplitude were estimated using the paired 5 cm lasers visible in each frame. The bedform period was confirmed using the data from the ROV’s onboard forward-looking sector scanning sonar. Data QA/QC was conducted by randomly selecting and reanalyzing 1.5 h of video imagery.

Variogram analysis of abundance data was used to identify the optimal spatial scale for subsampling transects following the methods of Camarero et al. (2005). Semivariograms plotted the semivariance of total organism abundance in each transect versus distance (m) between observations. Multidirectional (0°, 34°, 90°, and 135°) semivariograms of ROV transects were calculated and compared to a Gaussian model semivariogram with range = 60, psill = 1, and nugget = 1 using a Cressie estimator. Distances where the experimental semivariance exceeded the modeled semivariogram indicated the distance below which spatial autocorrelation was likely present.

While transects crossed multiple habitats (Fig. 2), transects were subsampled post hoc into 10 m sampling units based on the optimal spatial scale indicated by the variogram analysis, with each sampling unit restricted to 1 habitat type (i.e. inside or outside RSD). To create the 10 m sampling units, each transect was apportioned into consecutive 10 m sampling units starting from the RSD transition zone and moving outward to ensure only 1 habitat type within each sampling unit. Any length of transect that could not be apportioned evenly into a 10 m sampling unit was discarded. Organism counts and percent cover measures collected from individual sampling frames were aggregated into this larger 10 m sampling unit for analysis.

Mean density and richness was calculated for individual taxa and groups (with the exception of the infaunal invertebrate group, which was quantified by percent cover) by summing the total number of organisms identified in each sampling frame of each 10 m long sampling unit and dividing by the average area of the sampling unit. The average area of the
sampling unit was measured by the linear distance traveled by the ROV (10 m) multiplied by 0.87 ± 0.02 m, the average width of the sampling frame. To represent density and richness values per 10 m², density and richness values were then multiplied by 1.15.

Welch’s 2 sample t-test was used to test for significant differences in the mean faunal density and richness between habitat types for all enumerated groups. Two-way ANOVAs were used to test for significant differences in the mean density, richness, and percent cover values of individual groups between habitat and water depth. Data were tested for normality and equal variance to meet the assumptions of ANOVA. Density data were right skewed (i.e. Poisson distribution), and a square root transformation was performed prior to statistical analysis.

Collection and analysis of sediment grain size

Sediment grabs taken inside and outside of RSDs were conducted opportunistically along transects using a sediment sampler modeled after a Kahlsico Scoopfish to sample the upper 10 cm of sediment. Samples were only taken in locations where both multibeam imagery and video footage verified habitat type, and only whole samples were retained and analyzed. Coarse fraction analysis was conducted following the methods of Poppe et al. (2000) to calculate a mean grain size for each sample.

To assess how representative the physical characteristics of the Monterey Bay RSDs were of those found on the central California continental shelf, we analyzed an additional 190 archived sediment samples collected in 2006 from inside and outside of RSDs along 45 km of open coast immediately south of Monterey Bay using the same methods described above. A 1-way ANOVA with Tukey’s multiple comparison test revealed mean (±SE) grain size of sediments inside RSDs (0.19 ± 0.01 mm) to be significantly finer than samples collected inside RSDs (0.71 ± 0.03 mm, p < 0.001) but not significantly different from sediments outside RSDs on the Big Sur shelf to the south (0.25 ± 0.01 mm, p = 0.462). The Big Sur shelf RSD mean grain size (0.87 ± 0.02 mm) was significantly coarser than the outside sediments in Big Sur (p < 0.001) and slightly but significantly coarser than the Monterey

**RESULTS**

The ROV video surveys yielded 463 sampling units inside and 609 sampling units outside (each unit 10 m²) a total of 20 individual RSDs in Monterey Bay. A total of 2953 observations of 28 different fish and invertebrate taxa were derived from this video imagery. Individuals were identified belonging to 13 (N = 59), 25 (2034), and 18 (860) separate taxa in the shallow, intermediate and deep depth zones, respectively. A total of 33 sediment grab samples from inside (n = 13) and outside (n = 20) the Monterey Bay RSDs across a 14 to 40 m depth range were collected and analyzed for mean grain size along with the 190 Big Sur shelf archive sediment samples collected in 2006 from inside (n = 103) and outside RSDs (n = 87) spanning a depth range of 12 to 95 m.

**Sediment grain size inside and outside RSDs**

Grain size analysis revealed all sediments collected inside of RSDs to be coarser than all sediments from outside RSDs across all depth zones for Monterey Bay and the Big Sur shelf (Fig. 3). In Monterey Bay, ANOVA and Tukey’s multiple comparison test revealed mean (±SE) grain size of sediments outside RSDs (0.19 ± 0.01 mm) to be significantly finer than samples collected inside RSDs (0.71 ± 0.03 mm, p < 0.001) but not significantly different from sediments outside RSDs on the Big Sur shelf to the south (0.25 ± 0.01 mm, p = 0.462). The Big Sur shelf RSD mean grain size (0.87 ± 0.02 mm) was significantly coarser than the outside sediments in Big Sur (p < 0.001) and slightly but significantly coarser than the Monterey
Bay RSDs ($p = 0.001$). While mean RSD grain sizes in Monterey Bay tended to decline with depth over the 14 to 40 m depth range sampled, the opposite trend was found on the Big Sur shelf when all samples from the much wider 12 to 95 m depth range were included (Fig. 3). However, no trend with depth was found in the Big Sur shelf grain size data over the same 14 to 40 m depth range as sampled in the Monterey Bay study area.

Visual observation of the sediment surface in the ROV video imagery further confirmed the relative differences in grain size inside and outside of all RSDs in all zones (Fig. 4). The coarse nature of the sediment and abundant small shell fragments were consistent features observed inside but not outside the RSDs.

**Effects of habitat on density and richness**

The overall mean ($\pm$ SE) faunal density for all taxa, excluding YOY Sebastes spp., enumerated as individuals across all depth zones outside RSDs ($2.63 \pm 0.14$ ind. 10 m$^{-2}$), was greater than those found inside RSDs ($1.71 \pm 0.09$ ind. 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -4.82$, df = 1068.06, $p < 0.001$). This same pattern was true for the 3 faunal groups of individually enumerated taxa. The density of suspension feeders across all depth zones was nearly 3 times as great outside RSDs ($0.23 \pm 0.04$ ind. 10 m$^{-2}$) as inside RSDs ($0.08 \pm 0.02$ ind. 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -3.84$, df = 926.78, $p < 0.001$). Similarly, the density of invertebrate predators was twice as great outside (0.64 $\pm$ 0.05 ind. 10 m$^{-2}$) as inside RSDs (0.30 $\pm$ 0.03 ind. 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -5.98$, df = 1052.99, $p < 0.001$), and fish were more abundant outside (1.76 $\pm$ 0.11 ind. 10 m$^{-2}$) than inside RSDs (1.33 $\pm$ 0.08 ind. 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -2.28$, df = 1069.99, $p = 0.02$). Significant differences in the abundances of 12 individual taxa were found between habitats in some depth zones, with 10 taxa having higher values outside RSDs and only Dendraster excentricus and YOY Sebastes spp. having higher numbers inside RSDs (Table 1).

Mean richness of individual groups was also generally greater outside RSDs than inside. The mean richness of combined faunal groups outside RSDs ($1.49 \pm 0.06$ taxa 10 m$^{-2}$) was greater than inside RSDs ($1.12 \pm 0.06$ taxa 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -4.54$, df = 1068.72, $p < 0.001$). Mean richness of suspension feeders was greater outside RSDs ($0.14 \pm 0.02$ taxa 10 m$^{-2}$) than inside ($0.07 \pm 0.01$ taxa 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -3.24$, df = 1059.37, $p = 0.001$), as was that of invertebrate predators ($0.48 \pm 0.03$ and $0.25 \pm 0.02$ taxa 10 m$^{-2}$ respectively; Welch’s 2-sample $t$-test, $t = -5.74$, df = 1061.78, $p < 0.001$). However, there was no significant difference in mean richness of fish outside RSDs ($0.86 \pm 0.04$ taxa 10 m$^{-2}$) versus inside ($0.79 \pm 0.04$ taxa 10 m$^{-2}$; Welch’s 2-sample $t$-test, $t = -1.29$, df = 1032.75, $p = 0.20$).

**Effects of habitat and depth on density and richness**

Differences in density and richness of total fauna enumerated as individuals inside and outside of the

![Fig. 4. ROV video frames taken in (A) 54.6 m water depth outside an RSD and (B) 55.3 m water depth inside an RSD at locations 17 m apart along a single deep zone transect. Insets are photographs of inside and outside RSD sediment grab samples from the deep zone with coin (USA dime) shown for scale. Abundant ophiuroid arms protrude from the finer sediment in (A) but are completely absent in (B), where YOY rockfish hover between the prominent and comparatively barren bedforms of the much coarser RSD sediment. Paired laser scaling dots seen in both frames are 5 cm apart](image-url)
RSDs (excluding YOY *Sebastes* spp.) were not consistent across the 3 depth zones sampled. Density and richness of total fauna were greater outside RSDs in the intermediate and deep zones but not the shallow zone (Fig. 5). These differences by depth are reflected in the significant interaction found between habitat and water depth on both density (2-way ANOVA, $SS = 80.19$, $df = 5$, $F = 49.14$, $p < 0.001$) and richness (2-way ANOVA, $SS = 298.91$, $df = 5$, $F = 50.63$, $p < 0.001$) of combined faunal groups.

Interaction between depth and habitat was significant for density and richness of suspension feeders and invertebrate predators ($p < 0.001$; Fig. 5). Density of suspension feeders was greater outside RSDs only in the deep zone ($p < 0.001$), but richness was greater outside in both the intermediate and deep zones ($p = 0.01$ and $p < 0.001$, respectively). Density and richness of invertebrate predators were greater outside RSDs for the intermediate and deep zones ($p < 0.001$). Density of fish was greater outside RSDs in the intermediate zone ($p < 0.001$), and richness of fishes was significantly greater outside RSDs in the intermediate zone.

Differences were also found in the abundance of those fauna for which percent cover was used as a measure (*Dendraster excentricus*, *Urechis caupo*/Callianassidae burrows, and ophiuroids; Table 1, Fig. 4). Two-way ANOVA revealed significant interaction of depth and habitat for *D. excentricus* ($SS = 0.003$, $df = 2$, $F = 6.92$, $p = 0.001$), *U. caupo*/Callianassidae burrows ($SS = 0.04$, $df = 2$, $F = 9.22$, $p = 0.001$), and ophiuroids ($SS = 2.30$, $df = 2$, $F = 1059.71$, $p < 0.001$). *D. excentricus* was found almost exclusively within the shallow zone, where percent cover was significantly higher inside versus outside RSDs ($p < 0.001$). Mean percent cover of *U. caupo*/Callianassidae burrows was significantly greater outside RSDs in both the shallow and intermediate depth zones ($p < 0.001$ and $p = 0.001$, respectively). Ophiuroids were absent from the shallow zone and rarely observed in the intermediate zone but were 3 orders of magnitude more abundant outside RSDs in the deep zone ($p < 0.001$; Table 1, Fig. 4).

### Table 1. Two-way ANOVA summary of mean percent cover and density for individual taxonomic groups that comprise faunal groupings inside and outside of rippled scour depressions (RSD) by depth zone. Asterisks denote p-values (*0.01, **0.001, ***<0.001) of significant differences inside versus outside RSDs within a given depth zone for each taxonomic group. Values in **bold** indicate a significant interaction effect of habitat and depth for a given taxonomic group. Dashes indicate no individuals observed within that depth zone. Number of sampling units (n) are given for each habitat/depth category.

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<td><em>Dendraster excentricus</em></td>
<td>2.0 ± 0.60***</td>
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<td><em>Urechis caupo</em>/Callianassidae</td>
<td>1.29 ± 0.46</td>
<td>6.19 ± 0.66***</td>
<td>1.0 ± 0.22</td>
<td>2.68 ± 0.32**</td>
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<td>Ophiuroids</td>
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<td>0.00 ± 0.00</td>
<td>0.24 ± 0.03</td>
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<td>Suspension feeders (mean ± SE ind. 10 m⁻²)</td>
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<td><em>Metridium</em> spp.</td>
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<td><em>Ptilosarcus</em> spp.</td>
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<td>Invertebrate predators (mean ± SE ind. 10 m⁻²)</td>
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<td><em>Asterina</em> spp.</td>
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<td><em>Pisaster</em> spp.</td>
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<td><em>Pychnodoria</em> sp.</td>
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<td><em>Nassarius</em> spp.</td>
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<td>0.06 ± 0.03</td>
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<td><em>Cancer</em> spp.</td>
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<td>0.06 ± 0.01***</td>
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<td><em>Pagurus</em> spp.</td>
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<td>0.02 ± 0.01</td>
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<td>Octopus</td>
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<td>0.21 ± 0.04</td>
<td>0.23 ± 0.03</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Fish (mean ± SE ind. 10 m⁻²)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Ophiodon</em> sp.</td>
<td>-</td>
<td>-</td>
<td>0.06 ± 0.02</td>
<td>0.04 ± 0.01</td>
<td>0.02 ± 0.01</td>
<td>0.39 ± 0.08***</td>
</tr>
<tr>
<td><em>Sebastes</em> (YOY) spp.</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td><strong>2.81 ± 0.61</strong>*</td>
<td>0.20 ± 0.04</td>
<td>0.95 ± 0.25</td>
</tr>
<tr>
<td><em>Paralichthyidae</em> spp.</td>
<td>0.09 ± 0.05</td>
<td>0.02 ± 0.01</td>
<td>0.63 ± 0.07</td>
<td>1.04 ± 0.08**</td>
<td>0.91 ± 0.10</td>
<td>1.22 ± 0.24</td>
</tr>
<tr>
<td><em>Pleuronectidae</em> spp.</td>
<td>-</td>
<td>0.02 ± 0.02</td>
<td>0.42 ± 0.05</td>
<td>1.05 ± 0.08**</td>
<td>0.81 ± 0.09</td>
<td>0.99 ± 0.21</td>
</tr>
</tbody>
</table>

**Fish size and density**

The mean size of flatfishes for both families combined was greater outside (7.54 ± 0.32 cm) than inside (5.72 ± 0.20 cm) RSDs. However, this pattern...
varied significantly with depth (2-way ANOVA, SS = 862.85, df = 5, \(F = 21.33\), \(p < 0.001\)). Mean total length of combined flatfishes did not differ by habitat in the shallow or intermediate zones but was significantly greater outside RSDs in the deep zone (\(p < 0.001\); Fig. 6). Because the observed Paralichthyidae spp. tended to be larger than Pleuronectidae spp., this difference had the potential of skewing the overall mean flatfish size difference between inside and outside RSDs if there had also been a habitat-specific difference in the abundance of these 2 taxa. However, a paired t-test revealed that there were no significant differences in the abundance of these 2 groups between habitats (Table 1). Flatfish was the only group of fish for which there was an adequate number of reliable measurements made inside and outside RSDs to perform size analyses on.

YOY Sebastes spp. abundance was significantly higher inside RSDs in the intermediate zone, while this group was never observed in the shallow zone for either habitat type (Figs. 4, 7 & 8). Two-way ANOVA revealed a significant interaction between habitat and depth on mean density of YOY in the intermediate zone (SS = 45.07, df = 5, \(F = 27.40\), \(p < 0.001\)), where these fish were an order of magnitude more abundant inside than outside RSDs. While species identification was difficult for every YOY Sebastes spp. observation (N = 875), of the 605 YOY that could be identified to species, 45% were positively identified as S. pinniger (canary rockfish) based on their distinctive dorsal spots. Other positively identified Sebastes species found in these RSD-associated YOY aggregations included S. miniatus (vermilion rockfish), S. saxicola (stripetail rockfish), S. mystinus (blue rockfish), and S. melanops.
(black rockfish). This preference for RSD habitat by YOY *Sebastes* spp. is further confirmed by the spatial distribution of all 875 YOY observations. YOY *Sebastes* spp. were observed in all but one of the intermediate and deep RSDs sampled and never in the shallow depth zone (Fig. 8a), with the overwhelming majority of observed YOY found inside intermediate and deep RSDs (90 and 72% respectively; Fig. 8b).

### DISCUSSION

The present study determined that the biological communities associated with RSDs in southern Monterey Bay were significantly different from those found in the adjacent soft sediments. The RSDs sampled in the present study had sharply defined boundaries separating their coarser grain sediments and longer period bedforms from the 0.3 to 0.5 m higher adjacent and relatively featureless fine sediment plateau (Figs. 1 to 4). As predicted, these coarse sediments tended to support less dense and less diverse biological communities, and the density and richness of most faunal groups were lower inside RSDs for the intermediate and deep zones (Figs. 4 & 5). Also as predicted, the relative difference between biological communities inside and outside RSDs increased with depth. However, the prediction that larger bedforms would support larger fish was not borne out. Indeed, the opposite was true; smaller flatfish were found inside RSDs.

The finding that the densities of suspension feeders, invertebrate predators, and fishes, as well as the richness of suspension feeders and invertebrate predators, were significantly greater outside RSDs is consistent with our *a priori* predictions that RSDs are relatively depauperate habitats. This hypothesis was based on physical descriptions of RSDs as coarse sediment depressions found on fine sediment plateaus (Figs. 1 & 3), combined with the well-established

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**Fig. 7.** Mean (±SE) density of YOY *Sebastes* spp. inside and outside of rippled scour depressions by depth. Error bars represent the standard error, and sample sizes (n) are given. 

**Fig. 8.** YOY *Sebastes* spp. associated with RSD habitats by depth zone. (A) Percentage of individual RSD (inside) and non-RSD (outside) habitats sampled in which YOY *Sebastes* spp. were observed (n = number of individual habitat features sampled). (B) Percentage of observed YOY individuals observed inside versus outside of RSD habitats within each depth zone (N = total number of YOY identified as *Sebastes* spp. within each depth zone)
negative correlation between increasing grain size and species density and diversity (Snelgrove 1999, Brown & Collier 2008). This negative relationship has been documented for many fishes and invertebrates known to actively settle on sediments of a specific grain size (Chia & Crawford 1973, Kurihara 1999, Stoner & Ottmar 2003), with finer sediments for ease of burying (Stoner & Ottmar 2003), abundance of infaunal prey (Abookire & Norcross 1998, Brown & Collier 2008), or to maintain associations with other benthic organisms (Lindholm et al. 2004). While the present work highlights the physical and biological differences inside and outside of RSDs, future work should investigate the possibility of a halo- or ecotone effect at the typically well-defined RSD boundary where physical conditions change abruptly on the sub-meter scale.

As predicted, the effect of grain size alone did not determine the density or diversity of benthic communities (Snelgrove & Butman 1994), which also varied significantly with water depth. The relative difference in density and richness of benthic communities inside and outside of RSDs increased with depth, and significant differences were observed in the intermediate and deep zones that were not observed in the shallow zone (Figs. 4 & 5). Again, this prediction was based on the physical descriptions of both RSDs and the shallow nearshore zone as chronically disturbed by hydrodynamic currents and turbulence (Sternberg 1984, Green et al. 2004, Murray & Thieler 2004).

On the continental shelf, grain size typically decreases with depth as the frequency and magnitude of wave/current disturbance rapidly decreases (Lenihan & Micheli 2001, Allen 2006). However, the presence of RSDs extending beyond the 90 m isobaths on the central California continental shelf (Fig. 3) and to 160 m elsewhere (Bellec et al. 2010) indicate that disturbance inside RSDs is maintained over a broader depth range, thus amplifying the physical differences between habitats as depth increases (Green et al. 2004, Goff et al. 2005). This disturbance has a strong effect on the physical and biological characteristics of RSDs. In particular, hydrodynamic resuspension of sediments may have a negative impact on filter-feeding invertebrate species, whose feeding apparatus may be clogged with suspended sediments (Rhoads & Young 1970, Pillay et al. 2007). Indeed, in the present study, suspension feeders were generally less dense and less rich inside RSDs (Table 1).

Additionally, hydrodynamic scouring may inhibit the settlement of infaunal organisms (Jumars & Nowell 1984), which serve as food for many fish and invertebrate species (Quammen 1984, Edgar & Shaw 1995). Moreover, while the focus of the present study was largely constrained to epifaunal taxa that could be identified in video imagery, the very strong and well-documented influence of grain size on the diversity and abundance of infaunal species (Hall & Hard- ing 1997, Abookire & Norcross 1998, Brown & Collier 2008) suggests that RSDs will likely have an even greater influence on those invertebrate communities.

Exploration of these differences offers a way to examine hypotheses of food availability as a mechanism for the differences in epifaunal and benthic predatory fish described herein. The ROV video imagery revealed turbulent flow generated by even weak currents moving over the RSD bedforms during non-storm periods, sufficient to lift and suspend small particles from the seabed in visibly greater amounts than seen in the water column over non-RSD sediments and thereby potentially increasing prey availability to small fish inside RSDs compared to the otherwise richer surrounding non-RSD sediments.

There were also differences with respect to dominant taxonomic groups found between habitats and depth zones. This was especially true for Dendraster excentricus, Urechis caupo/Caliannassidae burrows, and ophiuroids, which had significant differences in percent cover for the depth zone in which they were most abundant (Table 1, Fig. 4). Burrows (likely formed by U. caupo/Caliannassidae) dominated outside RSDs in both the shallow and intermediate zones, while ophiuroids dominated outside RSD in the deep zone. The dominance of these organisms outside RSDs is especially relevant to the density and diversity of benthic communities, as both U. caupo/Caliannassidae and ophiuroids have been shown to create habitat for other fish and invertebrate species (MacGinitie 1934, Piepenburg & Juterzenka 1994).

While the strong interaction between habitat and depth on the density of organisms and diversity of benthic communities has been well established (Warwick & Davies 1977, Barry & Dayton 1991, Snelgrove & Butman 1994, Whitman et al. 2004), no previous work had examined the distribution and abundance of benthic communities associated with RSDs. The present study offers a first look at the ecological influence of these abundant features, and our results show RSDs conform to the general understanding of how benthic communities respond to fine scale physical differences in soft sediment habitats (i.e. grain size, bedforms, and hydrodynamic disturbance). However, many questions remain as to the larger scale significance of RSDs, including how RSD size,
configuration, and latitudinal distribution mediate the biological relationships observed in the present study and ultimately the significance of RSDs with respect to marine spatial planning (i.e. ecosystem-based management and marine protected area [MPA] design; Browman & Stergiou 2004, Pikitch et al. 2004, Gleason et al. 2010).

Most surprising, and of potentially great significance for marine spatial planning, was the use of RSDs by fishes. While differences in fine scale physical variables allowed us to make reliable predictions about the overall distribution of benthic communities with respect to RSDs, we did not anticipate finding small flatfish and abundant YOY Sebastes spp. inside RSDs (Figs. 4, 6 & 7). The present study revealed flatfish to be significantly smaller inside RSDs (Fig. 6). We had predicted, based on the larger size of bedforms inside RSDs, to find larger fish. This unexpected finding is likely due to a combination of physical and biological interactions. The larger bedforms inside RSDs can act as a refuge for fish small enough to tuck into their lee (Gerstner 1998). As the size of the organism increases, its ability to benefit from refuge between bedforms may diminish. Furthermore, the main predators of juvenile flatfish are larger fishes (Gibson & Robb 1996), which were more abundant outside RSDs. While the mechanism and seasonality of this relationship needs to be investigated further, these physical and biological factors may create an incentive for small fish to seek out RSDs despite potentially harsher physical conditions. This finding suggests a potential ecological role of RSDs as previously undescribed nursery habitat, which has important implications for the design of MPA networks (Beck et al. 2001, Gillanders et al. 2003).

The significantly greater density of YOY Sebastes spp. inside RSDs within the intermediate depth zone and their strong preference for RSDs in both the intermediate and deep zones were surprising discoveries (Figs. 7 & 8). While many species of rockfish are known to recruit in late summer to early fall (Carr 1991, Johnson et al. 2001, Caselle et al. 2010) to a variety of habitats including kelp canopy (Carr 1991, Nelson 2001, Johnson 2006), high-relief hard substrate (Carlson & Straty 1981, Carr 1991, Johnson 2007), deep low-relief rock reefs (Anderson & Yoklavich 2007, Love et al. 2009), and shallow surfgrass beds (Guido et al. 2004), the importance of soft substrata for the young of deep-dwelling rockfish species is relatively unknown (Love et al. 1991, Johnson et al. 2001). The intermediate depth zone RSDs investigated here supported densities of YOY Sebastes spp. comparable to densities found in artificial reefs (0.13 to 0.6 fish 10 m⁻²; West et al. 1994) and kelp canopies (10 to 40 fish 10 m⁻³; Carr 1991). This comparison was particularly true for the most abundant YOY we observed, S. pinniger. Indeed, RSDs may offer ideal conditions as nursery habitats for these fish, with the larger bedforms generating turbulent flow that provides a rich source of suspended organic material and small prey lifted off the bottom, as well as offering refuge from predators or currents.

The potential importance of RSDs as nursery habitat is underscored by the fact that 45% of the identified YOY were canary rockfish Sebastes pinniger, some populations of which are currently classified as overfished (Wallace & Cope 2011). Given the large areal extent of RSD habitat now documented for the California continental shelf (385 km² compared to 616 km² of rocky habitat) and the fact that the majority of these features occur immediately offshore of rocky reefs (A. Davis et al. unpubl.), RSDs may have important implications for the replenishment of adult rockfish on those adjacent reefs and therefore the performance of MPAs in terms of faster than expected population recovery rates. Additionally, the widespread and abundant distribution of RSDs along the California coast may serve as conduits for gene flow between widely spaced reefs, thereby increasing the connectivity of rockfish populations on the west coast of the United States (Miller & Shanks 2004, Hyde & Vetter 2009). If true, this mechanism has significant implications for marine spatial planning efforts, such as California’s Marine Life Protection Act, which mandates that the state’s MPA system be designed to function as a network facilitating connectivity of populations through larval dispersal and ontogenetic migrations (CDFG 2008). The timing of the present study (July) was fortuitous for observing YOY rockfish, but further sampling of these habitats seasonally will be required to determine the term of residency for the young of these and other species throughout the year.

Recently collected high-resolution seafloor maps of the California continental shelf reveal RSDs to be abundant and responsible for making soft bottom habitats much more heterogeneous than previously thought (Morrisey et al. 1992, Kostylev et al. 2001, Brown & Collier 2008, A. Davis et al. unpubl.) Now, with the results presented here, it is also clear that RSD-generated heterogeneity adds a significant level of ecologically important patchiness to the continental shelf, challenging the common assumption within many marine resource management strategies that soft sediments can be treated as homogeneous habitats.
Acknowledgements. We thank the Marine Technology Society, International Women’s Fishing Association, Friends of Moss Landing Marine Labs, and the California State University Monterey Bay (CSUMB) Undergraduate Research Opportunities Center for financial support. We also thank A. Davis, C. Mueller, and J. Carrillo for help with the ROV surveys, J. Vasques, M. Carr, and M. Young for aid in manuscript preparation, T. Laidig for fish identifications, D. Smith for sediment analysis, and C. Garza for statistical advice. This research would not have been possible without the high-resolution seafloor data and maps generated by the California Ocean Protection Council’s California Seafloor Mapping Project and CSUMB Seafloor Mapping Lab.

LITERATURE CITED


Auster PJ, Malatesta RJ, LaRosa SC, Cooper RA, Stewart LL (1991) Microhabitat utilization by the megalauana assemblage at a low relief outer continental shelf site—Middle Atlantic Bight, USA. J Northwest Atl Fish Sci 11: 50−69


Chia FS, Crawford BJ (1973) Some observations on gametogenesis, larval development, and substratum selection of the sea pen, Pilosarcus gurneyi. Mar Biol 23:73−82

Diaz RJ, Cutter GR, Able KW (2003) The importance of physical and biogenic structure to juvenile fishes on the shallow inner continental shelf. Estuar Coast 26:12−20


Kvitek RG, Goldberg JD, Smith GJ, Doucette GJ (2008) Domoic acid contamination within eight representative species from the benthic food web of Monterey Bay, California, USA. Mar Ecol Prog Ser 367:35–47


MacGinitie GE (1934) The natural history of Callianassa californiensis Dana. Am Midl Nat 15:166–177


Biol Annu Rev 32:111–177

Editorial responsibility: Jana Davis, Annapolis, Maryland, USA
Submitted: January 18, 2012; Accepted: July 18, 2012
Proofs received from author(s): November 2, 2012