Ectosymbiosis is a critical factor in the local benthic biodiversity of the Antarctic deep sea

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ABSTRACT: In deep-sea benthic environments, competition for hard substrates is a critical factor in the distribution and diversity of organisms. In this context, the occurrence of biotic substrates in addition to mineral substrates may change the characteristics of sessile fauna. We tested this hypothesis at different localities of the Weddell Sea (Antarctica) by studying the diversity of ectosymbionts living on the spines of cidaroids (echinoids). The presence of cidaroids promoted a higher total specific richness and increased sessile species abundance, but did not change the diversity. Analyses of species distribution suggested that the cidaroids are a favourable habitat for sessile organisms, compared to rocks, but are colonized by relatively specialist sessile species, leaving the unfavourable rock habitat to more generalist species. Therefore, our study highlights the role of some living organisms, such as cidaroids, as key species increasing Antarctic benthic deep-sea species richness through the niche they provide to symbiotic species.

KEY WORDS: Symbiosis · Species diversity · Species abundance · Null models · Cidaroids · Weddell Sea

INTRODUCTION

Interactions between organisms have an extensive influence on their distribution and abundance. Historically, competition, predation and ecosystem engineering were considered to be important factors controlling community structure (Jones et al. 1994, Begon et al. 1996). More recently, the importance of symbiosis, mostly the role of parasitism or mutualism, has been identified in changing other interactions (e.g. Clay & Holah 1999, Clay 2001, Hatcher et al. 2006). However, few studies show that symbiosis can modify community structure.

Antarctic marine biodiversity is of particular interest because of the relative isolation of the Southern Ocean, its high endemism and its sensitivity to perturbations (Brey et al. 1994, Battaglia et al. 1997). Extensive investigations of Southern Ocean benthic biodiversity have taken place only within the last few decades, thanks to the ever more numerous explorations, mostly consisting of photographic surveys at all depths including the deep sea (Brey et al. 1994, Gutt & Starmans 1998, Starmans et al. 1999, Brandt et al. 2007). The forces shaping benthic biodiversity in the Antarctic are not yet fully understood, but communities seem to be structured by bathymetry (Gutt 2000) as well as by the geography of the shelf (Clarke et al. 2007). Communities are dominated by sessile organisms, but biomass and diversity distributions are discontinuous, with patches of high abundance and diversity surrounded by quasi-deserts (Arntz et al. 1994, Gutt & Starmans 1998, Starmans et al. 1999, Gutt 2000). The nature and accessibility of hard mineral substrates appear to be important factors affecting the distribution of benthic species (Gutt 2000). In the Southern Ocean, large areas of the deep-sea bottom are covered with poorly sorted sediments. Most of the sediments consist of mud derived from the decomposition of pelagic organisms (Tyler 1995), while hard substrates include pebbles as well as large and small rocks dropped by icebergs, the rocks having been removed from the continent by glacial erosion (Andrews et al.
Rocks are usually partly buried in the sediment, limiting the availability of these hard mineral substrates to sessile benthic organisms (Knox 1994, Tyler 1995, Andrews et al. 1997). Biotic interspecific interactions, such as competition for access to limited substrates (Gutt 2000), have been proposed as another key factor to explain species distribution. In addition to such competition, Gutt & Schickan (1998) suggested that epibiosis is quite common among species of the Antarctic benthos, and may govern the distribution of species richness. In case of scarcity of mineral hard substrates, biotic substrates (such as shells) can provide suitable sites for attachment and therefore new ecological niches for sessile species. Ectosymbiosis (i.e. sessile life on biotic substrate, whatever the nature of the relationship: mutualism, commensalism, or parasitism) is commonly observed in marine ecosystems (Key et al. 1996, Williams & McDermott 2004). In the Antarctic, numerous species have been documented as epibions or substrates, respectively (Barnes & Clarke 1995, Gutt & Schickan 1998); however, these studies were not able to establish the importance of ectosymbiosis in the biodiversity of sessile organisms, mainly because relative proportions of species fixed on living organisms versus those fixed on abiotic substrates were not compared at geographically restricted, homogenous sites. Therefore, it is difficult to determine if ectosymbionts are specific to their living substrate or if they are opportunistic sessile organisms, selecting other organisms as substrate because of the rarity of mineral substrates.

Among biotic substrates available in the Antarctic, a clade of echinoids (sea urchins), the Cidaroida, appears to be useful in addressing questions concerning ectosymbiosis. Echinoderms represent 55% of the benthic biomass in the Antarctic, but the cidaroids are rare and have a patchy distribution. They can be locally abundant, however, reflecting the general pattern of benthic species distribution (Jacob et al. 2003). The shaft of their spines is not covered by an epithelium and lacks an anti-fouling mechanism (Märkel & Röser 1983), a characteristic that allows colonization by a large number of epibions. While several studies have described epibiosis on cidaroids, recent (e.g. Brey et al. 1993, Gutt & Schickan 1998, Massin & Hétérier 2004, David et al. 2005a) and fossil (Schneider 2003) analyses of epibiont diversity and specificity remain rare (Hétérier et al. 2004, Hopkins et al. 2004). Cidaroids have rarely been taken into account in exploration of epibenthic biodiversity (Ragua-Gil et al. 2004). Hétérier et al. (2004) have shown that 2 species of Antarctic cidaroids, Ctenocidaris spinosa and Rhynchocidaris triplopora, are colonized by 60 and 30 morphological types of ectosymbionts, respectively, suggesting that the presence/absence of cidaroids on the sea floor might significantly affect local biodiversity, by providing new substrates for sessile organisms. However, epibiotic assemblages on cidaroids have not been compared with assemblages living on mineral substrates.

Here, we test the role ectosymbiosis involving cidaroids plays in establishing local biodiversity of the sessile fauna in the Weddell Sea. The present study aims to answer 2 main questions: (1) What is the role played by ectosymbiosis in abundance and diversity of sessile organisms at a local scale? (2) Are organisms involved in epibiosis found only on cidaroids, or are they generalists also found on non-living substrates? To answer these questions, samples were collected, using an identical sampling procedure, at different sites of the deep Weddell Sea, and the faunas were compared according to their sites of attachment: rocks or cidaroids.

**MATERIALS AND METHODS**

Samples were taken between 26 January and 30 March 2005 during the expedition ANDEEP III of the RV ‘Polarstern’ (ANT XXII/3) in the Southern Ocean (Linse et al. 2007). Of the 15 deep-sea benthic stations sampled in the Weddell Sea area, 10 were selected for analysis in the present study. From each of these 10 stations, samples were taken using an Agassiz trawl (opening 5 m). The location and depth of the stations, as well as the area and substrate sampled are given in Fig. 1 and Table 1, respectively. The stations

![Fig. 1. Sampling stations in the Weddell Sea (stars indicate the 2 stations with cidaroids from the western group)](image-url)
Table 1. Depth, surface sampled and nature of sediments at the 10 stations sampled during ANDEEP III. Cat: categories of station location (E: eastern, W: western part of the Weddell Sea without cidaroids; W*: station located in the western part of the Weddell Sea with cidaroids); nr: number of rocks analyzed; nc: number of cidaroids collected; Cr and Cc: rates of colonization by sessile organisms on rocks and cidaroids, respectively; Dr: approximation of mean diameter of rocks \[(length + width) / 2\]; Hr: mean height of rocks; Hc: mean height of cidaroids; nr: number of rocks analyzed; nc: number of cidaroids collected; Cr and Cc: rates of colonization by sessile organisms on rocks and cidaroids, respectively. Dr: approximation of mean diameter of rocks \[(length + width) / 2\]; Hr: mean height of rocks; Hc: mean height of cidaroids; nr: number of rocks analyzed; nc: number of cidaroids collected; Cr and Cc: rates of colonization by sessile organisms on rocks and cidaroids, respectively.

<table>
<thead>
<tr>
<th>Station no.</th>
<th>Cat</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Depth (m)</th>
<th>Surface sampled (m²)</th>
<th>Main sediments</th>
<th>nr</th>
<th>Cr (%)</th>
<th>Dr (mm)</th>
<th>Hr (mm)</th>
<th>nc</th>
<th>Cc (%)</th>
<th>Hc (mm)</th>
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<td>5°19.68’W</td>
<td>1822</td>
<td>1515</td>
<td>Mud</td>
<td>100</td>
<td>7.0</td>
<td>51.8 ± 10.4</td>
<td>44.1 ± 10.4</td>
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<td>–</td>
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<td>0°4.34’E</td>
<td>4648</td>
<td>2360</td>
<td>Mud–Rocks</td>
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<td>71.4 ± 6.4</td>
<td>42.5 ± 4.5</td>
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<td>–</td>
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<td>1610</td>
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<td>41.4 ± 7.7</td>
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<td>2780</td>
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<td>1310</td>
<td>Mud</td>
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<td>47°28.64’W</td>
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<td>1400</td>
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<td>28.5 ± 5.8</td>
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</table>

*aAll Aporocidaris milleri, bAll Ctenocidaris speciosa*

are all below the continental shelf (>1000 m, Clarke & Johnson 2003), deeper than the 1000 m limit below which a depth gradient vanishes in different taxonomic groups (e.g. Allcock et al. 2001, Hilbig 2001, Linse et al. 2006). The sampled stations were combined into an eastern group (‘E’) and a western group (‘W’). Cidaroid echinoids were only found at 2 stations of the western group (subset W*). Cidaroids and rocks, the only available abiotic hard substrates suitable for sessile organisms, were specifically sorted, measured and counted. The area of the rocks that was originally buried versus the area that was exposed (i.e. available for settling by invertebrates) could not be easily determined after the rocks were brought up to the surface, so their total height was measured. As the number of rocks always surpassed 500 in each sample, sub-samples ranging from 100 to 250 (proportional to a rough estimate of the total number) were taken for analysis.

Two species of cidaroids were found in our samples: Ctenocidaris speciosa Mortensen 1910 and Aporo-
cidaris milleri Agassiz 1898, and they occurred only at 2 stations (Table 1). Specimens with >5% broken spines were removed from the analyses to ensure reliability of epibiosis estimates. Cidaroids and rocks carrying at least 1 sessile organism were fixed in 70% ethanol for further analysis, and were considered as ‘colonized’. The prevalence of sessile organisms was calculated as the ratio of colonized individuals to total individuals. Identification and counting of sessile species was done in the laboratory under a stereo microscope. Abundance data were obtained for each colonized individual (cidaroid or rock). For abundance data of colonial species (e.g. sponges, bryozoans, hydrozoans), for which the notion of ‘individual’ is ambiguous, it is conservative to consider that a single colony was founded by 1 propagule only. Therefore, 2 colonies were considered as 2 ‘individuals’ when they were spatially discrete (e.g. attached to different spines of a given echinoid, or attached at several separated points of a given spine or rock). Attached organisms were identified at least to the class level, and to a lower taxonomic level (e.g. genus or species) when possible. Some unknown specimens were sent to specialists for identification.

Data were analysed using indices for diversity, richness and similarity as suggested by Chao & Lee (1992), Lee & Chao (1994) and Chao et al. (2005), using the SPADE software (Chao & Shen 2005) and following recommendations from the software user guide for the choice of indices. These indices, most of which are non-parametric, are an extension of classical indices used in community ecology (Krebs 1989), taking into account ‘unseen’ species in estimates and using replicates to compute confidence intervals, allowing statistical comparisons. To obtain these indices for a given sampling station, we treated individual cidaroids as replicates for describing the ‘cidaroid assemblages’, as suggested by Shaw & Dobson (1995) for host–parasite relationships, and single rocks as replicates for describing the ‘rock assemblages’. The estimate of S, the species richness index of a given assemblage, was obtained using the abundance-based coverage estimator \( S_{JAC} \). The estimate of \( H \), the species diversity index of a given assemblage, was obtained using the index \( H \) of Chao & Shen (2003). Finally, the estimate of \( J \), Jaccard’s similarity index between assemblages, was computed from abundance-based data \( (J_{JAC}) \), according to the approach of Chao et al. (2005).
The species abundance distributions (SADs) of sessile assemblages were also analysed according to methods reviewed by McGill et al. (2007). Such methods are independent of taxon identity and prevail for comparison of communities that do or do not share species. The rank abundance plot captures most of the information (McGill et al. 2007). Abundance data on a log_{10} scale were plotted against the rank of the species, from the most abundant (Rank 1) to the least abundant, and the distributions were inspected visually.

In order to find co-occurrence patterns of sessile organisms in the colonization of hard substrates (i.e. to determine whether the observed co-occurrence of epibionts departs from that expected by chance), we compared the observed species distribution with that generated by null models. Because the choice of a null model may influence the results (Gotelli 2000), the presence/absence matrices were analysed with 2 co-occurrence indices. Following Gotelli (2000) and Gotelli & Rohde (2002), we chose C-score and V-ratio indices. Both are single-number measuring patterns for an entire presence/absence matrix. The C-score quantifies the degree of species co-occurrences; the larger the index, the less co-occurrence of species pairs. The V-ratio measures the community structure: this ratio is >1 for strong positive covariance between species pairs and <1 for strong negative covariance (see Gotelli 2000 for more details). The indices were calculated for each matrix (i.e. each sampling station) and compared with indices obtained for 5000 null communities randomly generated from the observed matrix. Again following Gotelli (2000), 2 null algorithms were chosen, mainly for their good statistical properties when used in association with the 2 indices (they are not prone to Type I and Type II errors). The first is the ‘fixed–equiprobable’ (f–e) algorithm: during the randomization process generating null matrices, the symbiont occurrences are fixed, but all sites have the same potential to be colonized. The second null algorithm is the ‘fixed–fixed’ (f–f) algorithm, where both symbiont occurrences and settlement sites are fixed. Here, the algorithm preserves differences among sites in the number of symbionts they contained. Therefore, in the f–f model, the empty observed sites are not used to generate the null matrices, whereas, in the f–e model, all sites, including the empty ones, are considered suitable for colonization (Gotelli & Rohde 2002). The difference between the 2 null models allowed us to test symbiont distributions considering either all sites or only the already colonized sites as available for attachment, a factor that was impossible to observe in situ due to the great depth of the sampling sites. Analyses and comparisons of observed versus simulated indices were conducted with the software EcoSim 7.0 (Gotelli & Entsminger 2007) with a sequential swap algorithm creating the null matrices (Gotelli & Rohde 2002). Finally, we applied another kind of null model, based on species density distributions (Janovy et al. 1995). For each sampling station, the observed sessile species density distribution (i.e. the observed number of hosts harbouring 0 to n sessile taxa, n being the maximum number of sessile taxa observed) was compared to the null model distribution (i.e. distribution independent of each colonizing event). This null model distribution was obtained using a multiple-kind lottery model (see Janovy et al. 1995 for details). This algorithm is a kind of f–e model, as the observed frequencies of symbionts are used to generate the probabilities of colonization, but all fixation sites are equiprobably free for colonization. Comparisons of observed versus simulated distributions were made with a $\chi^2$ test (Janovy et al. 1995).

Comparisons of the sessile communities attached to rocks and on cidaroid spines were made at hierarchical spatial scales: inter-regions, intra-region but inter-stations, and intra-station. At the scale of the Weddell Sea (about 2000 km wide), we compared the ‘rock assemblages’ among 3 groups of stations: the eastern set (E), the western subset of stations without cidaroids (W) and the western subset of stations with cidaroids (W*) (Table 1). Within the western region, we compared assemblages at the scale of 100s of kilometers by distinguishing the W and W* stations, and, among the W* subset, by distinguishing the rock assemblages (W*r) from the cidaroid assemblages (W*c). At this level of analysis, the sampling stations were individualized. The last order of comparisons concerned ‘cidaroid assemblages’ and ‘rock assemblages’ at the scale of 10s of meters within the 2 W* stations.

Statistical analyses were carried out using the software JMP v.6 (SAS Institute).

**RESULTS**

Rock size did not significantly differ between geographic groups of stations (E vs. W + W*; ANOVA with ‘group’ as fixed factor and ‘station’ as random factor nested within ‘group’: $F_{5,128} = 1.51$, $p = 0.15$ for diameter and $F_{5,128} = 1.23$, $p = 0.28$ for height), allowing comparison of their colonization. Cidaroids, with their spines, were not significantly different in height from rocks at each station where they were found in common (W* stations) (Table 1; $t = –1.11$, $p = 0.30$ at Stn 142; $t = 0.86$, $p = 0.39$ at Stn 151).

Within the 2 W* sites (Stn 142 harboured only Aporocidaris milleri, while Stn 151 harboured only Clenocidaris speciosa), cidaroids were less numerous than rocks, but the number of sessile animals they carried was higher than that of rocks (Table 1; Fisher...
exact test, 2-tailed p < 0.005 in all cases). The proportion on colonized rocks was also variable between stations (Table 1; Pearson \( \chi^2 = 60.80, p < 0.0001 \)). Across stations, there was no correlation between average rock size and the rate of colonization (Spearman \( \rho = 0.19, p = 0.59 \)). The distributions of sessile taxa colonizing each station are given in detail in Appendix 1, Table S1 available in MEPS Supplementary Material at www.int-res.com/articles/suppl/m364p067_app.pdf).

Global analysis of the rock assemblages at a large spatial scale revealed that SADs were comparable on both sides of the Weddell Sea. In addition, in the western samples, the presence or absence of cidaroids did not substantially modify the SAD of rock assemblages (Fig. 2A). The general pattern of these SADs is a lack of dominant species (none was found with a total exceeding 100 individuals) and a slight excess of rare species. Diversity (Fig. 2B) and richness (Fig. 2C) did not significantly differ between the different groups of stations. In addition, rocks from E and W stations harboured very similar sessile faunas, sharing about 75% of taxa (Fig. 2D). However, in the western region, only 30% of the taxa were shared between rocks from stations with and without cidaroids (Fig. 2D). Generally, while diversity and distribution of organisms attached to rocks are relatively homogenous across the Weddell Sea, their species composition seems to be affected by the local presence of cidaroids. Therefore, ‘rock assemblages’ might be echinoid-sensitive. Indeed, the SADs for the W* stations, when including the cidaroids, departed strongly from those for rocks only (Fig. 2A).

Comparison of stations within the western Weddell Sea confirmed the general pattern. Stations without cidaroids (W) exhibited rock assemblages with a comparable magnitude of species richness (Fig. 3A), the only significant difference being between Stns 152 and 121. Total species richness of W* stations (with cidaroids) was significantly higher than that of W stations (Fig. 3A), with the exception of comparisons involving Stn 152, for which the confidence interval was very large. The high species richness in W* stations was not due to a higher species richness on cidaroids, but rather due to the combination of the 2 sub-assemblages (rock fauna plus cidaroid fauna, right part of Fig. 3A). Species diversity was not found to be altered or enhanced by the presence of cidaroids, the values of \( \hat{H} \) for total fauna (rocks plus cidaroids) falling within the same range in W or W* stations (Fig. 3B). Moreover, at Stn 142, species diversity of the cidaroid sub-assemblage was lower than that found on rocks (Fig. 3B), probably because of the presence of numerically predominant taxa (see Fig. 4A). Among the W and W* stations, the similarities of rock assem-
blages were very variable (Fig. 3C). The rocks of the 3 W stations shared most of their sessile fauna (>80% taxa in common). However, rocks from W and W* shared a significantly lower proportion of their sessile fauna (20 to 30%) (Fig. 3C). Rocks from the 2 W* stations showed an intermediate pattern and shared about half of the taxa.

At a local scale, the SADs of sessile fauna at W* stations were split into 2 sub-samples: the fauna found on rocks and the fauna found on cidaroids. The 2 SADs obtained were different, as abundant and medium-

abundant taxa were, relatively, most frequent on cidaroids but not on rocks (Fig. 4A). Within each W* station, the average similarity between cidaroid and rock assemblages was low. This was especially true at Stn 142, with <5% of species shared by the 2 types of substrates (Fig. 4B). Similarity was significantly different and higher at Stn 151, where about 50% of species were shared between substrates, with large variation around this estimate (Fig. 4B). However, we can hardly assume that sessile assemblages living on cidaroids are similar to those living on rocks. In addition, the species shared by cidaroids and rocks were not the more common species found at both W* stations (Appendix 1, Table S1): the prevalence distributions of shared versus unshared species were not significantly different on rocks (Wilcoxon non-parametric test: Z = 0.47, p = 0.65) or on cidaroids (Z = 0.13, p = 0.89). The 2 stations were pooled for this analysis, because there was no difference between stations in the prevalence of sessile organisms on rocks (Z = −0.71, p = 0.47) or on cidaroids (Z = −0.77, p = 0.44). For example, at Stn 151, the shared 'Foraminifera A' was one of the most prevalent and abundant taxa on both the cidaroid Ctenoci-


**DISCUSSION**

Our results suggest that the presence of cidaroid echinoids modifies the local diversity of the deep-sea sessile benthos in the Weddell Sea. It is worth noting that the assemblages of sessile faunas were quite similar on rocks across the Weddell Sea (Fig. 2), but very different between cidaroids and rocks within the same local sampling stations (Fig. 4). The observed pattern is that the presence of cidaroids generally promoted a higher total species richness (i.e. species fixed on both rocks and cidaroids), and allowed some species already present to become dominant (e.g. *Foraminifera A*), but also allowed the, sometimes abundant, settlement of additional species (e.g. *Hydrozoa A*). In addition, while different species were attached to rocks and cidaroids, respectively, the overall species diversity remained approximately the same.

The cidaroid endosymbiotic fauna can be richer or poorer than that on rocks, but it seems generally less diverse, with more very abundant species. Our results suggest that sessile communities in the Weddell Sea are not composed of generalist species able to settle on all available substrates. Such a situation is reminiscent of the fauna attached to hermit crab shells in the Mediterranean Sea (Bick 2006), where some specificity was observed. We propose that the presence of cidaroids provides specific ecological niches for sessile organisms, thereby strongly influencing local biodiversity at this level. This confirms observations by Gutt & Schickan (1998), but at a local scale and relying on consistent comparisons within the same sampling sites (the same micro-habitats).

Differences in sessile assemblages were found between the 2 stations with cidaroids. This suggests that cidaroid-specific characteristics are likely to also influence endosymbiotic diversity and distribution. Clearly, more studies are needed to investigate what these characteristics could be (morphological, chemical, or otherwise). For instance, cidaroid spine shape and roughness not only vary between species, but also
along the shaft within a given species (Hétérier et al. 2004, David et al. 2005a,b). Spine morphologies could thus have a discriminative effect on the settlement and development of associated species and could explain some differences in the distribution of ectosymbionts, a hypothesis amenable to manipulative experiments. Behavioural differences between hosts could also explain such differences, as observed between encrusting communities on gastropods shells and shells used by hermit crabs (Bell 2005).

At stations where both cidaroids and rocks were present, comparison of their respective sessile fauna indicates that the abundance of species on echinoids is generally higher than the abundance of species on rocks. In addition, the SAD of taxa on cidaroids showed no excess of rare species, while the distribution is left-skewed on rocks (they host more rare species). Recently, several community subdivisions (also called deconstructions), either temporal (Maguran & Henderson 2003, Ulrich & Ollik 2004) or spatial (Ulrich & Zalewski 2006), into ‘core’ species versus transient or satellite species have shown that left-skewed SADs can often be attributed to the fact that species that are non-permanently present in the community occur only occasionally, thus generating an overall disproportionate number of rare species. The pattern recorded for the deep Weddell Sea is similar, but, following the abovementioned interpretation, one would consider that core sessile species in the Weddell Sea are the cidaroid ectosymbionts, while the satellite fauna would be the one colonizing rocks. Such a situation is unexpected, since rocks as settlement sites are much more available and predictable than cidaroids. Therefore, the observed distribution pattern remains to be fully understood, but a tentative explanation can be proposed. Most of the epibionts found on rocks and cidaroids were filter-feeding organisms that need to be in the water flow to optimize their food uptake (Gutt & Schickan 1998). Although they can sometimes be as high as cidaroids, rocks can sink into and become covered with mud, while cidaroid spines offer sessile organisms a higher position in the water column. Rocks can be seen as a less favourable habitat for sessile fauna than cidaroids. Prevention from burial has been shown to be one of the benefits gained by sessile species. Several community subdivisions (also called deconstructions), either temporal (Maguran & Henderson 2003, Ulrich & Ollik 2004) or spatial (Ulrich & Zalewski 2006), into ‘core’ species versus transient or satellite species have shown that left-skewed SADs can often be attributed to the fact that species that are non-permanently present in the community occur only occasionally, thus generating an overall disproportionate number of rare species. The pattern recorded for the deep Weddell Sea is similar, but, following the abovementioned interpretation, one would consider that core sessile species in the Weddell Sea are the cidaroid ectosymbionts, while the satellite fauna would be the one colonizing rocks. Such a situation is unexpected, since rocks as settlement sites are much more available and predictable than cidaroids. Therefore, the observed distribution pattern remains to be fully understood, but a tentative explanation can be proposed. Most of the epibionts found on rocks and cidaroids were filter-feeding organisms that need to be in the water flow to optimize their food uptake (Gutt & Schickan 1998). Although they can sometimes be as high as cidaroids, rocks can sink into and become covered with mud, while cidaroid spines offer sessile organisms a higher position in the water column. Rocks can be seen as a less favourable habitat for sessile fauna than cidaroids. Prevention from burial has been shown to be one of the benefits gained by symbionts of hermit crabs (Williams & McDermott 2004). Gutt & Schickan (1998) proposed that sessile species would opportunistically colonize living substrates, rather than rocks. Nevertheless, our results do not support an opportunistic colonization of cidaroids, since their ectosymbiotic fauna differs from epibionts of rocks. However, because cidaroids are much rarer than rocks, colonization of the favourable micro-sites (spines) could be achieved by some specialists reaching high abundance, leaving the unfavourable rocks sites to generalists unable to reach high abundances. In the Antarctic, the rock–cidaroid differences in sessile fauna composition and abundance could therefore be due to differences in colonizing dynamics of specialist versus generalist species.

Comparison of co-occurrence patterns of sessile organisms between rocks and cidaroids tend to support this point of view. Settlement on rocks deviated from the null models at all stations when algorithms considered all sites equiprobably available for fixation. The observed excess of co-occurrence on rocks confirms that the majority is not suitable for sessile organisms. This could be due to the fact that some of these rocks were partially buried in the mud and therefore not fully accessible to colonization. However, if the non-colonized rocks were not considered in the analysis (f–f model), no aggregative pattern was found. This means that settlement on available rocks by a given species was hardly influenced by other species. Contrasting with the pattern recorded on rocks, species distribution on cidaroids always agreed with the null model distribution, suggesting that there is no species interaction for cidaroid colonization. This could indicate favourable attachment sites, where the occurrence of a given sessile organism is independent of the presence of other species. Finally, the strong segregation pattern observed, using the f–f null model, between rock andcidaroid assemblages at the same sampling stations strengthens the hypothesis of cidaroid colonization by specialist organisms.

To conclude, our study suggests that the presence of cidaroids may modulate benthic biodiversity in the Antarctic by providing attachment sites for ectosymbionts on their naked spines. Although more studies, investigating more cidaroid species and more biotopes, particularly in shallower water, are needed to confirm this assumption, the present study contributes to the understanding of the effects of species interactions on biodiversity (see also Clay 2001). It generates hypotheses that can be tested with other organisms in the Antarctic (see Gutt & Schickan 1998 for other organisms acting as substratum for sessile forms). Finally, it supports the hypothesis that cidaroids might be key species of the benthic Antarctic deep-sea ecosystem, facilitating the settlement of many others.

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