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PRIMARY RESEARCH PAPER

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Distinct or similar? Soft bottom polychaete diversity in Arctic and Antarctic glacial fjords

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Abstract The main aim of this study was to compare the polychaete communities in two similar polar areas: an Arctic fjord, Hornsund (Svalbard) and an Antarctic fjord, Ezcurra Inlet (South Shetlands). This is the first attempt to compare Arctic and Antarctic diversity based on fully comparable datasets. Forty van Veen grab samples were collected in each fjord: twenty replicates were taken in each of two fjord areas characterized by a different level of glacial disturbance-in the inner (glacial bay) and outer (fjord mouth) region of both fjords, from depths of about 100 m in 2005 (Hornsund) and in 2007 (Ezcurra Inlet). In the glacial bays, species richness and diversity were significantly higher in Ezcurra Inlet than in Hornsund due to higher rate of glacial disturbance in the latter one. In the outer areas, species richness was similar in

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National Marine Fisheries Research Institute, ul. Kołłataja 1, 81-332 Gdynia, Poland both fjords, although diversity values were higher in Ezcurra Inlet. Polychaete species richness in the habitats characterized by similar level of disturbance (outer areas of the fjords) was the same in both polar regions. At this small scale, where community drivers are very similar, the species richness seems to be independent from the local or regional species pool.

Keywords Polychaeta · Fjords · Polar regions · Diversity · Svalbard · South Shetland Islands

Introduction

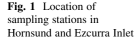
Until recently, it was commonly assumed that the benthic diversity in the Antarctic is substantially higher than in the Arctic (Dayton et al., 1994; Knox & Lowry, 1977; Crame, 1997; Brandt, 2001). Different geological age and different level of geographic isolation of those two regions and thus their evolutionary history were the main basis for this assumption. Arctic marine fauna is relatively young. The present environmental characteristics of the Arctic region were shaped by Pliocene cooling about 3 million years ago and some later Quaternary events (Dayton, 1990; Piepenburg, 2005). However, the current benthic fauna probably colonized the Arctic shelf only after the last glacial maximum, no more than 13,000 years ago (Clarke & Crame, 2010). In contrast, the Southern Ocean benthic fauna evolved in

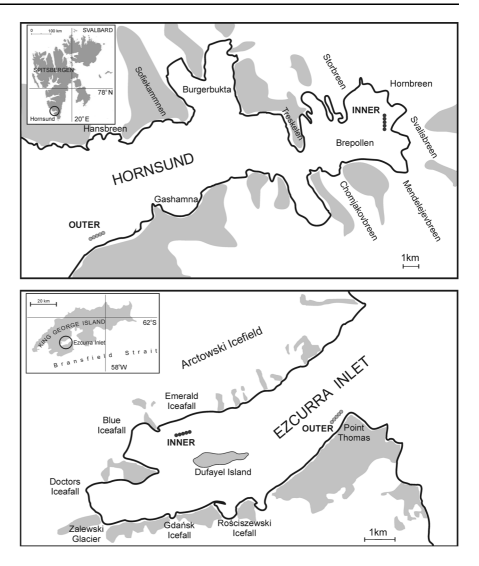
stable conditions and isolation since the establishment of the Drake Passage and Antarctic Circumpolar Current 34 million years ago (Barrett, 1999). Antarctic benthic fauna is often described as more diverse and characterized by higher level of endemism than the Arctic fauna (e.g., Knox & Lowry, 1977; Dayton et al., 1994). The homogeneity of the Arctic benthic habitats and higher rates of disturbance were also pointed as possible reasons for differences in diversity between both regions (Dayton, 1990). Yet, these common assumptions have recently been questioned and considered as a result of overgeneralization (Piepenburg, 2005). Recent taxonomic inventories of both polar regions demonstrated that in general the number of species is similar. De Broyer et al. (2011) listed 5,800 benthic species in the Southern Ocean, while the number of Arctic species was estimated to be about 4,800 (Sirenko, 2001; Piepenburg et al., 2011). Moreover, some studies suggested that both Arctic and Antarctic diversity are still underestimated (e.g., Carr, 2012; Pabis et al., 2014). Nevertheless, these totals can provide only a very superficial knowledge about differences and similarities between those two polar regions, and diversity may further vary depending on the taxonomic or ecological group. So far, there have only been few attempts to compare benthic diversity on Arctic and Antarctic shelves, and those included only large scale syntheses comparing either the Arctic and Antarctic as a whole (Knox & Lowry, 1977), or large basins, like the Laptev Sea and Weddell Sea (Piepenburg et al., 1997; Starmans et al., 1999; Starmans & Gutt, 2002; Sirenko, 2009). Moreover, those analyses were mostly focused on mega-epibenthic fauna, not always collected with quantitative sampling gear. Even single studies on the intermediate scale of polar fjords (Błażewicz-Paszkowycz & Sekulska-Nalewajko, 2004; Włodarska-Kowalczuk et al., 2007) were based on not fully comparable datasets since samples were collected from different substrates and/or different depths. Thus, rigorous comparisons of similar depths and types of habitats are essential as the only way to provide any further meaningful inventories (Clarke & Crame, 2010).

Glacial fjords are hosting a very rich and diverse fauna with many species associated only with this type of habitats (Włodarska-Kowalczuk & Pearson, 2004; Siciński et al., 2011; Włodarska-Kowalczuk et al., 2012; Grange & Smith, 2013). Those semi-closed inlets are affected by chronic disturbance processes associated with the influence of subglacial streams, resulting in high inflow of inorganic suspension and silting of bottom sediments. This is considered as the main reason of the fjords' vulnerability to environmental changes associated with on-going climate warming (Smale & Barnes, 2008; Węsławski et al., 2011). In this study, we compared two similar polar inlets: Hornsund, Svalbard, and Ezcurra Inlet, South Shetlands. Hornsund is an All Taxa Biodiversity Inventory site (ATBI) and is an important area for monitoring the effects of climate change (Warwick et al., 2003). Ezcurra Inlet is a part of Admiralty Baythe most comprehensively studied basin in the rapidly changing region of the Western Antarctic Peninsula (WAP) (Siciński et al., 2011). Both fjords have been intensively sampled for the last 35 years and can be treated as model polar fjord ecosystems. Information about the benthic diversity of those basins were recently summarized by Kędra et al. (2010a) and Siciński et al. (2011) as a result of two large polar marine programs: Marine Biodiversity and Ecosystem Functioning (MarBEF) and Census of Antarctic Marine Life (CAML).

Polychaetes are an ecologically diverse group that often dominate in terms of species richness, abundance, and biomass in marine benthic communities all over the world (e.g., Knox, 1977; Ansari et al., 1991; Hutchings et al., 1993; Mackie & Oliver, 1993; Smith, 2000), also in polar soft bottom communities (Kendall et al., 2003; Piepenburg, 2005; Clarke & Crame, 2010; Pabis et al., 2011). They are considered excellent surrogates of overall biodiversity and indicators of ecosystem response to changes in environmental conditions (Olsgard & Somerfield, 2000; Olsgard et al., 2003; Giangrande et al., 2005; Włodarska-Kowalczuk & Kędra, 2007). Currently, there are 590 polychaete species included in the Register of Antarctic Marine Species (De Broyer et al., 2011), while the number of species recognized in the Arctic reaches almost 660 (Sirenko, 2001). For those reasons, polychaetes were chosen as a model group in this study.

The aim of this study is to compare the soft bottom polychaete abundance, species richness, diversity, and functional groups in Arctic and Antarctic soft bottom habitats affected by different level of glacial disturbance: glacial bays and fjord mouth. Trustworthy diversity assessment of those two fjords based on the comparable material should bring new insights into





the debate on diversity of the marine seafloor in polar regions. It could also substantially influence creation of the future climate warming scenarios of changes in benthic community structure of polar fjords.

Materials and methods

Study area

Hornsund

Hornsund is a medium-sized glacial fjord located on the southwest coast of Spitsbergen (Svalbard, Arctic, 77°N) with a maximum depth of about 260 m. It is influenced by cold Arctic waters of the Sørkapp Current and

warmer Atlantic waters of West Spitsbergen Current (Swerpel, 1985). The inner glacial bay, Brepollen, is partially isolated by the Treskelen Peninsula (Fig. 1; Table 1) and is affected by meltwater inflow from the five large tidewater glaciers (Weslawski et al., 1995). High values of mineral suspension in this area result in high water turbidity (Gorlich et al., 1987; Piwosz et al., 2009) and a reduced euphotic layer (Urbański et al., 1980). Compared to the outer part of the fjord, the sedimentation in Brepollen is much higher (35 vs. 0.1 cm year^{-1}), while the mineral suspension content in Hornsund waters was estimated to be up to 1,000 and about 20 mg dm⁻³, respectively (Gorlich et al., 1987). Mud dominates bottom sediments but sandy deposits and dropstones often occur in the outer fjord area (Rudowski & Marsz, 1996; Szczuciński & Schettler,

| Table 1 Sampli | Sampling sites and their characteristics | | | | | |
|---|---|--|---|---|--|---|
| | Hornsund | | References | Ezcurra Inlet | | References |
| | Inner | Outer | | Inner | Outer | |
| Location | Glacial bay isolated from the rest of the fjord by Treskellen Peninsula, area influenced by five tidewater glaciers | Outlet area of the fjord not influenced by glaciers | Gorlich et al. (1987), Rudowski and Marsz (1996) | Glacial bay located behind the submerged sill influenced by two large tidewater glaciers—Emerald Icefall and Blue Icefall | Outlet area of the fjord located close to the rocky cliffs and not influenced by glaciers | Rudowski and Marsz (1996), Siciński et al. (2011), Campos et al. (2013) |
| Size and surface area | Fjord axis: 27 km, surface area: 275 km^2 | km ² | Gorlich et al. (1987) | Fjord axis: 9 km, surface area: 19 $\rm km^2$ | 19 km ² | Rakusa- Suszczewski 1993 |
| Sediments | Mud | Mud with sand and gravel, dropstones | Rudowski and Marsz (1996), Szczuciński and Schettler unpublished results | buM | Sandy mud, dropstones | Siciński (2004), Siciński et al. (2011), Campos et al. (2013), Own data |
| Organic matter content in the sediments | 1.75–1.99% | 1.37–1.84% | Szczuciński and Schettler unpublished results | No data | 2.58-4.83% | Own data |
| Hydrology | No influence of the warm waters of the West Spitsbergen Current, large outflow of ablation waters in the summer, cooling of the water by glaciers | Influence of outer waters of the West and South Spitsbergen Current | Swerpel (1985) | Very weak currents | Area influenced by strong current circulation which brings the strong water flows from the Bransfield Strait | Campos et al. (2013) |
| Surface water suspension concentration | Up to 1,000 mg/dm ³ | 10-20 mg/dm ³ | Gorlich et al. (1987) | >100 mg/dm ³ | 15 mg/dm ³ | Pęcherzewski (1980), Siciński (2004), Siciński et al. (2011) |

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unpublished results). All those factors affect primary production and organic matter content in the fjord resulting in reduced amounts of total organic carbon (TOC) in the areas located in the vicinity of glaciers (Gorlich et al., 1987; Eilersten et al., 1989; Piwosz et al., 2009). TOC reaches values between 1.75 and 1.99% in the inner basin and between 1.37 and 1.84% in the outer part of the fjord (Schettler & Szczuciński, unpublished data).

Ezcurra Inlet

Ezcurra Inlet is a part of Admiralty Bay. It is a semiclosed fjord of tectonic origin located at King Georg Island (South Shetlands, Antarctic, 63°S) with maximum depth of about 280 m (Marsz, 1983). The central basin of Admiralty Bay and outer areas of the inner inlets (Ezcurra, Martel, and MacKellar) are influenced by warmer and less saline waters from the Bellingshausen Sea and colder and more saline Weddell Sea waters (Gordon & Nowlin, 1978; Tokarczyk, 1987). The Ezcurra Inlet is a narrow fjord with tidewater glaciers distributed along its north-west coastline with the largest glaciers located in the inner region (Fig. 1). The amount of mineral suspension is decreasing from the inner glacial bays (>100 mg dm⁻³) toward the outlet area (about 15 mg dm⁻³) (Pęcherzewski, 1980; Siciński et al., 2011). High water turbidity was recorded in the inner region of this fjord contrasting with the lower turbidity in the fjord mouth (Lipski, 1987). Those differences in sedimentation processes are causing differences in sediment type: highly disturbed glacial bays are characterized by silt and clay sediments, while the outer region has sandy mud, poorly sorted deposits with larger amount of skeletal fractions (Siciński, 2004; Siciński et al., 2011; Campos et al., 2013). The differences between the inner and outer part are enhanced by bottom topography (Table 1): both parts of the fjord are separated by a submerged sill which partially isolates two areas (Marsz, 1983). The glacial sedimentation inflow is influencing the primary production in the waters of this inlet resulting in low concentrations of phytoplankton especially in the inner part of Ezcurra Inlet (Tokarczyk, 1986).

Sampling

Sampling was conducted in Hornsund and Ezcurra Inlet in July of 2005 and March of 2007, respectively, with standardized sampling methods. Two sampling areas were selected in regions of the fjords characterized by different levels of glacial disturbance. The inner area (glacial bay) and outer region (fjord mouth) of both fjords were sampled at depths of about 100 m (Ezcurra Inlet: inner 90-130 m, outer 100-130 m, Hornsund: inner 100-120 m, outer 100-115 m). In each area, 20 van Veen grabs (0.1 m²) were collected (5 stations in each area, 4 replicates at each station) (Fig. 1). Altogether 80 quantitative samples were gathered, 40 samples in each fjord. Samples were sieved through 0.5-mm mesh and fixed in 4% buffered formalin solution. An additional van Veen grab for sediment analysis (granulometry and TOC) was collected at each station. Those results are included in habitat descriptions provided in Table 1.

Laboratory and data analysis

In the laboratory, samples were sorted and polychaetes were picked, identified to the lowest possible taxonomic level and counted. Almost all specimens were identified to species level and only cirratulids to generic level. The closely related genera Aphelochaeta and Chaetozone, morphologically very similar taxa, are often destroyed, lack many important diagnostic features and require special collecting techniques. In order to compare functional groups diversity between the habitats and fjords, each species was assigned to the functional group following the classification proposed by Fauchald & Jumars (1979) which include feeding modes, mobility type, and type of buccal organ. This method of describing polychaete functional diversity is known as good tool for environmental assessment especially in characteristic of communities affected by various disturbance processes (Pagliosa, 2005).

Species accumulation curves with 95% confidence intervals were created for each area of both fjords using the formulae of Colwell (2009). Indices of species richness (*S*—number of species per sample, Margalef—*d*), evenness (Pielou—*J*), and diversity (Shannon index log e H and Hurlbert rarefaction index (ES[n], where n = 50)) were calculated (Hurlbert, 1971; Magurran, 2004). Levene's test was used to estimate the homogeneity of variance. Since the variances were not homogenized, differences between *S*, *d*, Shannon, and ES[50] indices in the outer areas of both fjords were tested with nonparametric Mann–

Hornsund Ezcurra Inlet Hornsund Ezcurra Inlet inner inner outer outer Total number of 6,079 10,742 17,024 13,379 individuals Total number of species 21 58 81 83 Number of singletons 3 (14%) 12 (20%) 21 (26%) 22 (26%) Number of doubletons 2 (9%) 5 (8%) 3 (3%) 6 (7%) Species richness (S) 9.4 ± 1.5 19.5 ± 3.3 34.3 ± 3.9 35.2 ± 5.0 Margalef Index 1.4 ± 0.2 $2.9\,\pm\,0.6$ 4.9 ± 0.5 5.3 ± 0.5 Shannon Index 1.1 ± 0.1 1.1 ± 0.2 1.9 ± 0.2 2.3 ± 0.2 ES [50] 5.6 ± 0.6 7.6 ± 1.3 11.1 ± 1.1 13.8 ± 1.1 0.5 ± 0.04 0.3 ± 0.07 0.5 ± 0.07 0.6 ± 0.04 Pielou Index Density [ind. 0.1 m²] 319.9 ± 83.8 537.1 ± 140.1 851.2 ± 126.7 668.9 ± 236.9

Table 2Comparison ofdiversity indices in thestudied areas (for singletonsand doubletons percentagein the total number ofspecies in each area isgiven)

Whitney U test. For all other indices, differences were tested with one-way ANOVA (STATISTICA 6 package). Information about species rarity is important in diversity estimates of benthic communities and was pointed as an important element of habitat-based analysis (Ellingsen et al., 2007; Fontana et al., 2008). Therefore, the number and percentage of rare species, defined as singletons (represented by only one individual) or doubletons (represented by only two individuals), were calculated for each area.

Results

Altogether 23,103 specimens were collected in Hornsund (HD), and 24,121 in Ezcurra Inlet (EI) (Table 2). The total number of polychaete species was similar in both fjords: 88 species from 30 families were found in Hornsund and 90 species from 26 families in Ezcurra Inlet. The most speciose families were: Syllidae (HD: 6, EI: 11), Terebellidae (HD: 8, EI: 10), and Sabellidae (HD: 6, EI: 9) (Fig. 2). The total number of species differed between the inner areas (21 species in HD, 58 in EI) but was very similar in the outer regions (81 in HD and 83 in EI) (Table 2). Those differences were reflected in the number of species from each functional group recorded in each of the studied sites. All functional groups were found in the inner area of the Ezcurra Inlet while in the inner area of Hornsund only six functional groups were present (Table 3). The number of species of sessile, surface deposit feeding (SST) and motile non-jawed burrowing (BMX) polychaetes was higher in the glacial bays in Ezcurra Inlet

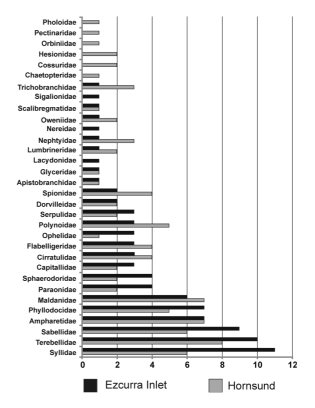


Fig. 2 Number of species in each family at each of studied sites

than in Hornsund (SST 11 vs 1 species, BMX 11 vs 5 species). All 7 functional groups were found in both outer areas and the total number of species in almost all categories was very similar (Table 3). Percentage of rare species was very similar in both fjords. In the inner areas singletons were slightly more numerous in Ezcurra Inlet (20%) than in Hornsund (14%) while in

| Functional group | Hornsund inner | Ezcurra Inlet inner | Hornsund outer | Ezcurra Inlet outer |
|------------------|---------------------|----------------------|---------------------|----------------------|
| BMX | 5 (105.8 ± 50.7) | 11 (60.6 \pm 24.2) | 10 (167.2 ± 43.1) | 10 (95.3 ± 34.4) |
| BSX | 0 | 3 (14.3 ± 13.5) | 7 (52.2 ± 17.6) | $6~(65.7\pm40.1)$ |
| CDJ | 0 | $1 (0.3 \pm 0.4)$ | $1 (0.05 \pm 0.2)$ | $1 (0.2 \pm 0.4)$ |
| СМЈ | 8 (7.2 ± 2.4) | $10~(25.3~\pm~7.1)$ | $22~(124.5\pm36.3)$ | 19 (16.0 \pm 7.9) |
| FST | $2~(15.4\pm7.1)$ | $7 (4.2 \pm 2.3)$ | $6 (4.0 \pm 3.0)$ | 10 (61.7 ± 70.0) |
| HMJ | 0 | 5 (1.1 ± 1.02) | $2~(0.8\pm 0.6)$ | 8 (29.0 ± 30.5) |
| SDT | $5~(13.0\pm 6.5)$ | 3 (22.5 ± 12.1) | 12 (88.4 ± 49.9) | 7 (18.6 ± 14.6) |
| SMT | $2(178.2 \pm 43.6)$ | 3 (398.2 ± 138.9) | 4 (386.7 ± 128.4) | 3 (203.1 ± 100.7) |
| SMX | 0 | $4 (4.6 \pm 3.9)$ | $2 (4.9 \pm 5.3)$ | 4 (165.2 \pm 77.6) |
| SST | $1 (0.1 \pm 0.3)$ | $11 (5.9 \pm 3.9)$ | $18 (22.2 \pm 7.9)$ | 16 (13.8 ± 10.8) |

Table 3 Polychaete functional groups in selected areas of both fjords-total species number and mean density with standard deviation (in brackets)

CMJ carnivore, motile, jawed; *CDJ* carnivore, discretely motile, jawed; *HMJ* herbivore, motile, jawed; *SMX* surface deposit feeding, motile, non-jawed; *SDT* surface deposit feeding, discretely motile, tentaculate; *SMT* surface deposit feeding, motile, tentaculate; *SST* surface deposit feeding, sessile, tentaculate; *SDJ* surface deposit feeding, discretely motile, jawed; *SMJ* surface deposit feeding, motile, jawed; *BMX* burrowing, motile, non-jawed; *BSX* burrowing, sessile, non-jawed; *FST* filter-feeding, sessile, tentaculate

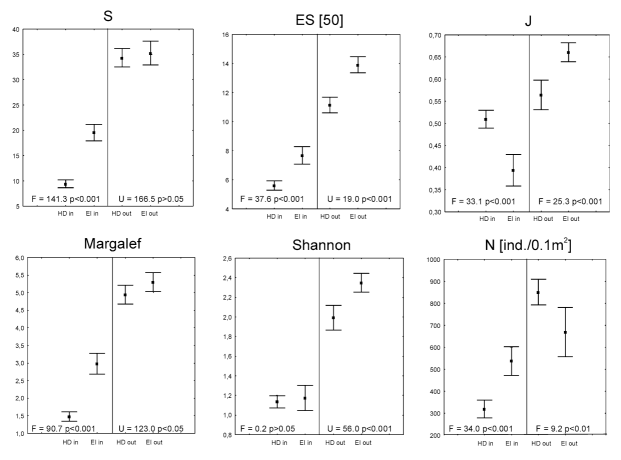
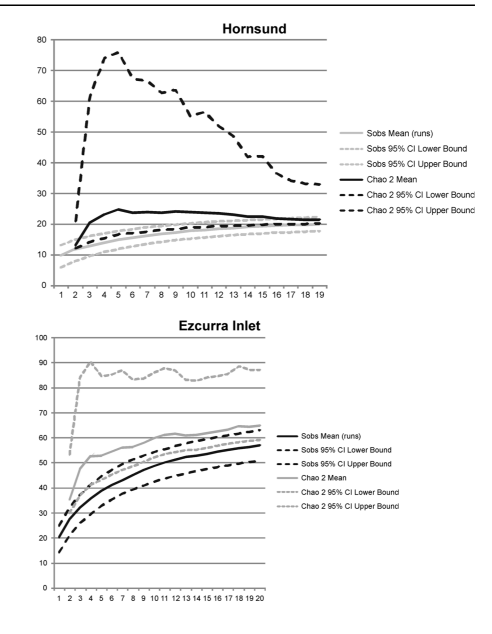


Fig. 3 Mean with 0.95 confidence intervals for: density (N number of individuals per 0.1 m²), species richness (S number of species per sample, Margalef Index), diversity

(*ES*[50] Hurlbert rarefaction index, Shannon Index) and evenness for each site (*HD* Hornsund, *EI* Ezcurra Inlet)

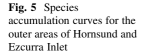
Fig. 4 Species accumulation curves for the inner areas of Hornsund and Ezcurra Inlet

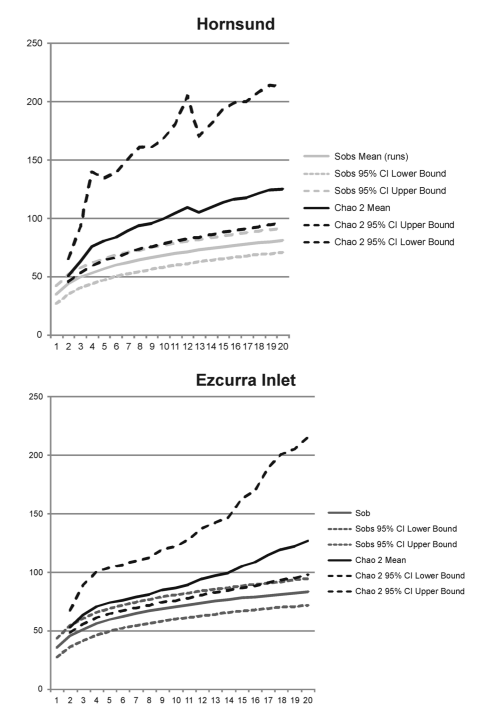


the outer areas of both fjords singletons comprised up to about 26% of all species (Table 2).

There were significant differences in density and all species richness and diversity indices except from the Shannon index between the inner areas of both fjords (one-way ANOVA, P < 0.05). Species richness (*S*) and ES[50] in Ezcurra Inlet's glacial bay were higher than in the inner area of Hornsund. On the contrary, the evenness values in the inner area of Ezcurra Inlet was lower than in Hornsund. Shannon index was very similar in both fjords (Table 2; Fig. 3).

In the outer areas, diversity (ES[50], Shannon index) was significantly higher in Ezcurra Inlet than in Hornsund (Man Whitney *U* test P < 0.05). There were no differences in species richness (*S*), however, the Margalef index was significantly higher in the outer area of Ezcurra Inlet (Man Whitney *U* test P < 0.05). Differences for evenness and density were also noted between the outer areas. Evenness was significantly higher in Ezcurra Inlet, while density was higher in Hornsund (one-way ANOVA, P < 0.05) (Table 2, Fig. 3). Species–accumulation curves were close to





asymptotic level in the Sobs and Chao2 richness estimators in the inner area of Hornsund (Figs. 4, 5). The curves were steeper in the other three areas.

Inner and outer areas of both fjords differed in terms of functional groups density. In the glacial bays, the most important differences were found for burrowers (BMX and BSX), carnivores (CMJ), and suspension feeders (FST). In the fjord mouth, the most important differences concerned carnivores (CMJ), suspension feeders (FST), but also surface deposit feeders (SDT, SMX) and herbivores (HMJ) (Table 3).

Discussion

The discussion about the differences and similarities between Arctic and Antarctic diversity remains open. Our study did not demonstrate an unequivocal result. Significant differences were found mostly for the inner areas of both polar fjords. Higher species richness and diversity in Ezcurra Inlet glacial bay are most probably related to much lower rate of disturbance in Ezcurra Inlet than in Hornsund (Table 1). The mineral suspension content in inner areas of Hornsund can be even an order of magnitude higher than in Ezcurra Inlet (Gorlich et al., 1987; Siciński et al., 2011). The meaningful conclusions for the regional differences can, therefore, be only drawn from the diversity comparisons between the outer areas, where the level of disturbance was similar in both fjords. Values of the most important disturbance agents like sediment type and sedimentation inflow were very similar despite the differences in size of both fjords (Table 1). In the fjord mouth, the total species number, the species richness of each functional group, and the number of species per sample were similar in both fjords, however, the diversity values were significantly higher in Ezcurra Inlet. On the other hand, it is worth mentioning that widely used diversity indices like Shannon and Hurlbert index might be influenced by the dominance of certain species (Gray, 2000; Salas et al., 2004) so the basic diversity measures like number of species per sample could be more useful in such bipolar comparisons.

Earlier studies conducted at the intermediate scale are also inconsistent. For example, higher α diversity of tanaidaceans (Shannon index, number of species per sample) was found in Admiralty Bay than in Kongsfjorden (Błażewicz-Paszkowycz and Sekulska-Nalewajko, 2004), but this group of crustaceans plays a minor role especially in the Arctic benthos (Brandt, 1997) and cannot be treated as surrogate of overall diversity. Nevertheless, the maximal values of tanaidacean species richness per sample were similar in both fjords. The only study of infaunal macroinvertebrates showed similar polychaete diversity between Admiralty Bay and two Arctic fjords: Kongsfjorden and van Mijenfjord (Włodarska-Kowalczuk et al., 2007). Results of both studies can be, however, biased due to incomparable datasets since samples from different habitats were taken into account in the analysis. Moreover, the results might be influenced by to the thirteen to even twenty years time span between the sampling campaigns, especially when rapid climate changes, that occurred during this period, are taken into account (Walsh, 2009).

Other attempts of comparison between Arctic and Antarctic diversity of benthic fauna also revealed some inconsistencies. The total number of species in various groups of macroepifauna in the methodologically standardized studies was higher in the Antarctic (Piepenburg et al., 1997; Starmans et al., 1999; Starmans & Gutt, 2002) while regional (Sirenko, 2001; De Broyer et al., 2011) and local (Weddell Sea vs Lazarev Sea) (Sirenko, 2009) comparisons of species lists showed similar species richness in both polar regions. It might suggest that some of the Arctic taxa have a higher share of rare species. However, α diversity (single sample results) differs depending on the groups studied and/or sampling gear used. For example, Piepenburg et al. (1997) demonstrated that echinoderm diversity was higher in the Weddell Sea than on the north-eastern Greenland shelf, yet that study was based on Agassiz trawl samples. It included stations from a depth range of 200-1,200 m with both shelf and slope fauna and did not take into account large differences between the depth of the Arctic (200-500 m) and Antarctic (500-1,000 m) shelf. On the contrary Starmans et al. (1999) and Starmans & Gutt (2002) did not find any significant differences in megaepibenthic α diversity between Arctic and Antarctic seas at shelf depths (underwater video survey). Brandt (2001) demonstrated higher densities of peracarids in the Arctic compared to the Antarctic and linked this observation with different age of those communities and lower maturity of the Arctic ecosystem which could therefore be dominated by fewer species occurring in higher densities. Our results did not confirm this opinion. In the outer areas, the total number of species, number of species per sample, and number of singletons were almost the same in both investigated fjords.

Higher habitat heterogeneity associated with the presence of large sponges, ascidians, and bryozoans, which are creating a complex habitat for other epifaunal organisms, was pointed as the most probable reason of higher diversity of Antarctic macro- and mega-epibenthic fauna (Barnes, 1994; Gutt & Schickan, 1998; Gutt & Piepenburg, 2003). Włodarska-Kowalczuk et al. (2007) emphasized that spatial heterogeneity of sedimentary habitats is in fact

comparable between Arctic and Antarctic shelf. However, the number of soft bottom microhabitats seems to be higher in Ezcurra Inlet than in investigated Arctic fjords (Kongsfjorden and Hornsund). Studies of polychaete diversity in Ezcurra Inlet and in Admiralty Bay as a whole showed high number of polychaete assemblages associated with different microhabitats (Siciński, 2004; Pabis & Siciński, 2010), while in Kongsfjorden the character of bottom deposits was less diversified resulting in more simple assemblage differentiation into three main faunal groupings associated with the inner, middle and outer areas of this inlet (Włodarska-Kowalczuk & Pearson, 2004). A similar pattern was also observed in Hornsund (Kędra et al., 2013). Our small scale study was focused on two clearly defined similar habitats to allow for more reliable comparisons. At this scale, where community drivers (sedimentation inflow, character of bottom sediments, organic matter content) influencing the fauna are very similar, we can expect similar species diversity. Those patterns could be independent of the evolutionary history of the whole regions (Hillebrandt & Blenckner, 2002). On the other hand, the regional species pool can be reflected at the local scale (Cornell & Lowton, 1992). Also, when the local area is smaller in proportion to the whole region it is less influenced by drivers shaping the regional species pool (Hillebrandt & Blenckner, 2002). Moreover, the Arctic and Antarctic species pools are independent and there is no overlap in species composition which was suggested as very important element in local-regional diversity comparisons (Ricklefs, 2000). Some of the recent molecular studies demonstrated the presence of bipolar species in some of the benthic invertebrate groups like amphipod crustaceans (Havermans et al., 2013), however, this result should be considered rather an exception from the rule (Brandt et al., 2012). There is a growing number of examples of polychaete species previously considered bipolar or cosmopolitan taxa turning out to be in fact species complexes (Scaps et al., 2000; Bleidorn et al., 2006; Barroso et al., 2010), and detailed taxonomic studies resulted in re-descriptions and changes in their taxonomic status (Parapar & Moreira, 2008). Nevertheless, Witman et al. (2004) showed that the factors operating at different spatial scales can be responsible for the diversity patterns. Therefore, scrutinizing the influence of the local conditions (including competition, level of disturbance, environmental factors) and their possible role in shaping regional species pools in the Arctic and the Antarctic is necessary.

The significance of the local and regional drivers in polar marine faunas still needs further investigations. Witman et al. (2004) demonstrated that 70% of local richness could be explained by the richness of the regional species pool. Our study showed contradictory results. The discrepancy between local (habitats) and intermediate (fjords) species richness is evident. The total number of polychaete species in Hornsund and Ezcurra Inlet differs strongly. 162 species were recorded in the whole Admiralty Bay (Siciński et al. 2011), 138 species were recorded in Ezcurra Inlet (Siciński, 1986, Hartmann-Schoder & Rosenfeldt, 1988, 1989; Siciński 2004; Pabis & Siciński, 2010; Siciński et al., 2012), while 93 species were found in Hornsund (Kędra et al., 2010a). Similar differences in general number of species were also highlighted in comparisons of peracarid diversity of those two fjords (Jażdżewski et al., 1995; Błażewicz-Paszkowycz & Sekulska-Nalewajko, 2004). A recently updated list based on more comprehensive studies showed similar results (Hornsund: Amphipoda 70 species, Tanaidacea 3 species, Admiralty Bay: Amphipoda 172 species, Tanaidacea 14 species (Kędra et al., 2010a; Sicinski et al., 2011). Those results are still quite similar for the Ezcurra Inlet alone (at least 110 species of Amphipoda Siciński et al. 2012, A. Jażdżewska unpublished results, and 8 species of Tanaidacea M. Błażewicz-Paszkowycz unpublished results). Svalbard polychaete diversity was estimated to be about 259 species (Palerud et al., 2004; Voronkov et al., 2013). The number of polychaete species known from South Shetlands was never summarized, however, basing on combined information from various sources (Hartman, 1964, 1966, 1967; Averincev, 1972; Hartman-Schroder, 1986; Hartman-Schroder & Rosenfeldt, 1988; Gallardo et al., 1988; Orenzsanz, 1990; Parapar & San-Martin, 1997; San Martin & Parapar, 1997; San-Martin et al., 2000; Lovell & Trego, 2003) it can be assumed that there are at least 230 species known from this area, so the result is similar to the one recorded in Svalbard waters. Total number of species in both fjords and archipelagos was not equally reflected in the richness on the habitat level in our small scale study. At large time scale almost all species from a regional species pool are expected to invade every community (Hillebrand & Blenckner, 2002). It could be justified especially for the animals

with relatively high dispersal ability like polychaetes (mostly planktonic larval stage) (Rouse & Pleijel, 2001). On the other hand, Bhaud (1998) demonstrated that polychates with long-lasting planktonic larval development do not necessarily have a larger geographic range which is further confirmed by the large differences in species composition between neighbouring fjords located along the Western Antarctic Peninsula (Grange & Smith, 2013).

Our data, including the large number of singletons and doubletons found in both fjords as well as species accumulation curves suggest that there are more species that can be potentially found in the investigated habitats, especially in the outer areas. Rare species are important for supporting stability of ecosystem functioning and at the same time are often highly vulnerable to environmental changes and disturbance events (Ellingsen et al., 2007). Their number as well as their role in the polar marine habitats still need more scientific attention especially in the context of ongoing rapid climate change which could result in an increased rate of disturbance in polar fjords. Węsławski et al. (2011) postulated that climate change may cause a homogenization of benthic habitats. It is possible that such changes may be reflected at first in the distribution of rare species which have lower environmental tolerance and high habitat specificity (Ellingsen et al., 2007). In our study, the number of rare species was much lower in the glacial bays, especially in Hornsund where level of glacial disturbance is higher than in Ezcurra Inlet (Table 1).

Our comparison of diversity within the same habitats between Ezcurra Inlet and Hornsund pose also important queries in the discussion on climate influence on benthic fauna in polar fjords. The Arctic as well as the region of the Antarctic peninsula (WAP) are amongst the fastest warming areas on Earth (Walsh, 2009). Influence of rapid warming on the community structure and diversity of Arctic benthic fauna were already documented (e.g., Kędra et al., 2010b; Grebmeier, 2012), but there are no similar studies in the Southern Ocean. Węsławski et al. (2011) highlighted that Arctic warming might influence not only diversity of benthic communities but cause also the functional simplification (with increasing dominance of non selective deposit feeders). Such changes could be especially pronounced in the inner areas of the fjords. Increasing sedimentation associated with warming in the WAP region could result in modification in community structure. It could be assumed that inner areas of Ezcurra Inlet (with still relatively high functional diversity) might in a future resemble the current state of the Hornsund glacial bays (area with clearly lower diversity of functional groups).

On the other hand, it was recently emphasized that the predictions and scenarios presented for the Arctic regions are not directly applicable to the Antarctic ecosystem (Smale & Barnes, 2008). For the infaunal communities in glacial fjords, especially in the outer areas where the level of disturbance is similar, the climate change influence on benthic fauna is pronounced in a similar way. Our study was based on the datasets collected in both fjords at the same time, when climate change was already visible. According to Grange & Smith (2013), Admiralty Bay and other fjords at South Shetlands are in later stage of climate warming compared to the fjords located further south in the WAP region and are more similar to the Spitsbergen fjords. Our study further confirms this opinion. In consequence climate related changes, which will be observed in Ezcurra Inlet and also in Hornsund, can reflect future modifications of the more southern Antarctic basins. Those sites can function as natural laboratories where increasing dynamics of glacial processes will influence the marine biota, giving us the opportunity to predict future scenarios and discuss their applicability in different inlets distributed along the Antarctic Peninsula.

Conclusions

On the habitat level, in the areas characterized by similar environmental properties and similar level of disturbance (outer areas of the fjords), species richness of soft bottom polychaete communities is almost the same in the Arctic and in the Antarctic. That seems to be independent from the species pool of the whole fjords and probably not influenced by differences in evolutionary history of both polar regions. Those results demonstrate how important role local conditions play in shaping benthic diversity in polar regions. Our results suggest very complex interactions occurring even at relatively small scale which makes the reliable conclusions on the regional scale difficult and far from being completed. The discrepancy between the habitat scale (Hornsund inner and outer—88

species, Ezcurra Inlet inner and outer-90 species; this study) and the intermediate scale of whole fjords (Hornsund—93 species, Ezcurra Inlet—138 species) (Siciński, 1986; Hartmann-Schoder & Rosenfeldt, 1988, 1989; Siciński, 2004; Kędra et al., 2010a; Pabis & Siciński, 2010; Siciński et al., 2012) imply that further analysis should include studies of processes associated with migrations of species between the habitats, as well as the influence of pre- and postsettlement events, competition, development modes, and dispersal abilities. Further studies should also include comparisons of various types of benthic habitats. It will allow for more comprehensive evaluation of differences and similarities of benthic diversity in polar regions and make extrapolations from habitat to local and regional scale more adequate.

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