Community dynamics of nematodes after Larsen ice-shelf collapse in the eastern Antarctic Peninsula

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Abstract

Free-living marine nematode communities of the Larsen B embayment at the eastern Antarctic Peninsula were investigated to provide insights on their response and colonization rate after large-scale ice-shelf collapse. This study compares published data on the post-collapse situation from 2007 with new material from 2011, focusing on two locations in the embayment that showed highly divergent communities in 2007 and that are characterized by a difference in timing of ice-shelf breakup. Data from 2007 exposed a more diverse community at outer station B.South, dominated by the genus Microlaimus. On the contrary, station B.West in the inner part of Larsen B was poor in both numbers of individuals and genera, with dominance of a single Halomonhystera species. Re-assessment of the situation in 2011 showed that communities at both stations diverged even more, due to a drastic increase in Halomonhystera at B.West compared to relatively little change at B.South. On a broader geographical scale, it seems that B.South gradually starts resembling other Antarctic shelf communities, although the absence of the genus Sabatieria and the high abundance of Microlaimus still set it apart nine years after the main Larsen B collapse. In contrast, thriving of Halomonhystera at B.West further separates its community from other Antarctic shelf areas.

Introduction

The Antarctic Peninsula is one of the most affected areas worldwide by rapid regional warming (Vaughan et al. 2003), and this has led, among other things, to large-scale ice-shelf destabilization and disintegration. The Larsen area east of the Peninsula is one of the regions where ice-shelf collapse is evident: in 1995, the Larsen A ice shelf (LIS-A) disintegrated almost completely, and in February–March 2002, the Larsen B ice shelf (LIS-B) lost with roughly 3250 km² the largest proportion of its surface after a decade of several smaller disintegration events and millennia of stability (Rack and Rott 2004; Domack et al. 2003; Rebesco et al. 2014). The sudden collapse of LIS-B was mainly attributable to surface processes, rather than basal melting in response to oceanic warming (Gilbert and Domack 2003; Vaughan et al. 2003; Rack and Rott 2004; Scambos et al. 2004; Rebesco et al. 2014). Prior to the actual breakup, there had been an exceptionally warm summer and the surface net mass balance of the ice shelf had been decreasing for several years (Rack and Rott 2004). This eventually led to ice thinning and the formation of meltwater ponds and crevasses at the surface, further enhancing rapid disintegration (Gilbert and Domack 2003; Rack and Rott 2004). Currently, the remnant LIS-B (and its tributary glaciers; Rott et al. 2011; Berthier et al. 2012) continues to decrease, evidenced by an additional loss of 50% of the initial collapsed area over the period 2002–2009 (Shuman et al. 2011).

Sudden ice-shelf collapse results in profound changes for associated marine benthic ecosystems. In areas like Larsen (e.g., the western Antarctic Peninsula; Moline et al. 2004; Clarke et al. 2007), loss of permanent shelf ice enables phytoplankton to bloom in areas previously ice-locked for several millennia (Bertolin and Schloss 2009; Barnes and Clarke 2011). Furthermore, ice algae released...
upon seasonal ice melt may provide a valuable additional food source, especially in seasonally opened polynyas nearby the continent (Cape et al. 2014). Together, both processes enhance direct fresh food supply to seafloor-dwelling organisms, triggering colonization of previously ice-covered habitats from nearby sources. On the downside, sudden ice-shelf decay increases the risk of iceberg scouring as large icebergs break off and ground in areas further offshore (Gutt et al. 1996; Lee et al. 2001).

Despite all efforts in the study of benthic response to climate-induced events such as ice-shelf collapse and iceberg scouring, considerable uncertainty remains on how biodiversity is affected by, and what the resultant ecological responses are of these processes. To gain long-term information, several benthic faunal components of Larsen B were sampled during two expeditions onboard the German icebreaking RV Polarstern in austral summer of 2007 (ANT-XXIII/8) and 2011 (ANT-XXVII/3). Meiobenthos (32–1000 μm) of the first expedition was assessed by Raes et al. (2010) and Hauquier et al. (2011), focusing on the numerically most important Nematoda. Already then, 5 years after the main LIS-B collapse, significant differences were observed between Larsen stations for all faunal groups, driven by different response rates to the change from an oligotrophic sub-ice shelf to a more productive ecosystem (Gutt et al. 2011). Based on faunal abundance and diversity, stations B.South located at the original ice-shelf edge and B.West in the middle of the embayment contrasted most. For Nematoda, this observation was explained by a combination of the duration of the ice-free period and the connection with precollapse open Weddell Sea conditions (Raes et al. 2010). The main objective of expedition ANT-XXVII/3 in 2011 was to revisit 2007 locations and look at benthic ecosystem recovery and dynamics. This study re-analyses 2007 data for stations B.South and B.West and compares them with new (i.e., 2011) nematode community data to resolve nematode community response to ice-shelf collapse on a longer time scale. Given continued increase in vertical food supply and exchange with the open Weddell Sea, we hypothesize that:

1. Abundance and diversity at B.West will increase and nematode communities at both locations will converge in terms of numbers, diversity, and generic composition,

2. Communities within Larsen B will increasingly resemble other Antarctic shelf areas of similar water depth that do not necessarily share the same history of permanent ice shelter.

**Material and Methods**

**Sampling area and strategy**

Stations B.South and B.West of Polarstern expedition ANT-XXIII/8 (January 2007) were re-sampled during ANT-XXVII/3 (March 2011) using five random replicate multicorer deployments (MUC, inner diameter 57 mm; Barnett et al. 1984) per location, allowing for equivalent and comparable sample coverage (Table 1, Fig. 1). B.South was always located at the border of the original ice shelf in open connection to the Weddell Sea (hence referred to as “outer” station), whereas B.West (inner station) experienced permanent ice cover until after the 2002 collapse (evolution of ice-shelf extent is depicted in Fig. 1; see also Raes et al. 2010).

The top 0–5 cm of one core per replicate deployment was sliced at a 1-cm resolution and preserved in 4–8% formalin for meiofauna analysis. Table 1 gives the geographic position and depth of the 2011 and – for ease of comparison – the 2007 samples. Meiofauna was extracted from the sediment using 1 mm and 32 μm sieves and density gradient centrifugation with Ludox (specific density 1.18 g cm$^{-3}$; Heip et al. 1985; Vincx 1996), fixed

| Station | Replicate | Latitude | Longitude | Depth (m) | Replicate | Latitude | Longitude | Depth (m) |
|---------|-----------|----------|-----------|-----------|-----------|----------|-----------|-----------|
| B.South | 700-8     | 65° 54.98’ S | 60° 20.54’ W | 422       | 246-3     | 65° 54.95’ S | 60° 20.43’ W | 424       |
|         | 700-9     | 65° 54.95’ S | 60° 20.88’ W | 417       | 246-4     | 65° 54.95’ S | 60° 21.49’ W | 395       |
|         | 702-4     | 65° 55.12’ S | 60° 19.96’ W | 427       | 246-5     | 65° 54.99’ S | 60° 20.70’ W | 419       |
|         | 702-7     | 65° 54.49’ S | 60° 21.37’ W | 405       | 247-3     | 65° 55.12’ S | 60° 19.83’ W | 428       |
|         | 702-8     | 65° 54.95’ S | 60° 20.95’ W | 410       | 247-4     | 65° 55.15’ S | 60° 20.01’ W | 425       |
| B.West  | 710-2     | 65° 33.03’ S | 61° 36.98’ W | 277       | 233-4     | 65° 32.99’ S | 61° 36.94’ W | 277       |
|         | 710-3     | 65° 33.04’ S | 61° 37.18’ W | 281       | 233-5     | 65° 32.97’ S | 61° 36.94’ W | 278       |
|         | 710-7     | 65° 33.03’ S | 61° 37.01’ W | 275       | 235-4     | 65° 32.96’ S | 61° 36.88’ W | 276       |
|         | 710-8     | 65° 33.03’ S | 61° 37.00’ W | 283       | 235-5     | 65° 33.01’ S | 61° 36.96’ W | 280       |
|         | 710-9     | 65° 33.07’ S | 61° 37.06’ W | 288       | 235-6     | 65° 33.01’ S | 61° 37.00’ W | 279       |
Finally, sediment total organic matter (TOM) was multiplied by 14:12 to account for the difference in molar mass.

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in 4% formalin, and dyed with Rose Bengal (0.5 g L⁻¹).

All meiofauna was counted and identified at higher taxonomic level using a stereo microscope and the guide of Higgins and Thiel (1988).

From each layer, 150 nematodes were randomly picked (or all when the number of nematodes <150), transferred to anhydrous glycerol (Seinhorst 1959), and mounted on slides. Genus-level identification (using a Leica DMLS compound microscope, 1000× magnification) was based on the pictorial key of Warwick et al. (1998) and the NeMYS database (Vanaverbeke et al. 2014).

As for 2007, samples for faunal analysis were complemented with an additional sample set for the measurement of environmental variables. These were analyzed at a coarser vertical resolution, 0–3 cm and 3–5 cm. Sediment grain size distribution was determined by laser diffraction (Malvern Mastersizer 2000, size range 0.02–2000 μm) and classified following Wentworth (1922). Granulometric variables considered in this study were median grain size, silt (<63 μm) and sand (>63 μm) percentage. Pigments were extracted from lyophilized sediments by adding 10 mL 90% acetone, and chlorophyll a (chl a; μg g⁻¹) was measured with a fluorescence detector after HPLC (high-performance liquid chromatography) separation. Additionally, total organic carbon (TOC) and nitrogen (TN) fractions were measured on 2011 freeze-dried samples using a Flash 2000 organic elemental analyzer (protocol available through Interscience B.V., Breda, The Netherlands). Their ratio was calculated and multiplied by 14:12 to account for the difference in molar mass (C:Ng outf). Finally, sediment total organic matter (TOM) was determined after combustion at 550°C.

**Statistical analyses**

Nematode abundance and community composition in 2011 were analyzed both separately and in conjunction with 2007 data. Analyses were executed in PRIMER v6 (Clarke and Gorley 2006) with the PERMANOVA+ add-on (Anderson et al. 2008), unless mentioned otherwise. Nematode assembly data were standardized to individuals per 10 cm² (ind.10 cm⁻²) and square-root transformed to limit influence of dominant genera.

Differences in communities between areas and sediment depth layers in 2011 were assessed using a PERMANOVA (permutational ANOVA) design with two fixed factors (area, layer; Bray–Curtis similarity of genus ind. 10 cm⁻²; 9999 permutations) and visualized using PCO (principal coordinate analysis). SIMPER (similarity of percentages) identified which genera were responsible for (dis)similarities between samples. Community data were then summed for 0–3 and 3–5 cm depth for each replicate preceding correlation with environmental variables (as these were measured at a rougher scale) and averaged for both areas. TOM was log-transformed to reduce right-skewness, and sand content was omitted from the analysis owing to its high correlation (r > 0.9) with silt. All environmental variables were normalized. BEST analysis quantified the correlation between environmental setting and nematode assemblages.

Comparison of 2011 and 2007 data was made by PERMANOVA. Univariate analysis of nematode densities used a two-factor design (area, year = fixed; Euclidean distances of nematode ind.10 cm⁻²; 9999 permutations), multivariate nematode composition data a three-factor design (area, year, sediment depth = fixed; Bray–Curtis similarity of genus ind.10 cm⁻²; 9999 permutations). Pairwise tests were performed between all pairs of levels for significant factors. When the number of unique permutations exceeded 100, true permutational P-values were reliable. When this number was below 100, Monte Carlo P-values were interpreted. Results were accompanied by a PCO graph, combined with CLUSTER results, to gain visual insight in the data cloud.

Diversity indices (N₀ = number of genera; $H'$ = Shannon index (loge); $EG(200) =$ expected number of genera in a sample of 200 individuals; Hill’s $N_J$) and evenness (Hill’s $N_{Curtis}$; $J'$ = Pielou’s evenness) were calculated in accordance with Raes et al. (2010). The rarefaction index $EG(n)$ was based on 200 as the lowest number of identified specimens in one of the replicates was 215 (Clarke and Gorley 2006). After assumption testing in R (R Core Team, 2013), several indices did not fulfill requirements for two-way ANOVA; hence, differences in diversity between areas and years were assessed using PERMANOVA (design identical to that for abundance data).
Finally, the 2007 and 2011 Larsen data were included in a larger dataset on (sub)-Antarctic nematode shelf assemblages (0–1000 m) to examine relationships within a broader geographical context (Table 7). Data were grouped over larger geographical scales to simplify analysis. Groupings were chosen arbitrarily, disregarding geographical coordinates, and should not be interpreted as true biogeographical provinces. One-way ANOSIM (analysis of similarity) assessed differences between areas, which were visualized with nonmetric MDS (multidimensional scaling).

**Results**

**Nematode abundance and vertical distribution**

In all 2011 samples, regardless of their location or sediment depth, nematodes formed the most abundant meiofaunal taxon (relative contribution 93–95%). Whereas nematode total densities (i.e., summed over 0–5 cm) in both areas differed a lot in 2007, they were comparable in 2011 (and no longer significantly different; Tables 2 and 3: pairwise tests for factor area). This is the result of a clear increase in total nematode densities at B.West, and a slight (but insignificant) decrease at B.South (Tables 2 and 3: pairwise tests for factor year). Also nematode vertical distribution differed between stations and years (Fig. 2). Vertical profiles showed steep declines with depth in both years for B.South. Profiles were less steep at B.West, especially in 2011 when nematode density peaked at 1–2 cm and remained relatively high down to 4 cm depth.

**Nematode community composition**

Nematode community composition in 2011 differed significantly between stations and cm-layers (two-factor PERMANOVA, significant interaction “area × layer”; results not shown) with largest differences between surface layers, gradually declining when moving deeper into the sediment (all pairwise $P < 0.05$). This is visible in the PCO plot for both stations in 2011 (Fig. 3). The first PCO axis (37.1% variation) divides samples according to their location, while the second axis (20.2% variation) is related to sediment depth. The dominant genera were *Microlaimus* at B.South and (a single species of) *Halomonhystera* at B.West (Table 4), which together explained almost half of the dissimilarity between both stations (average dissimilarity = 85.58%; contribution *Halomonhystera + Microlaimus* = 46.92%; SIMPER).

These two genera were also responsible for the clear separation between stations in terms of years (Fig. 4), as they remained most abundant at their respective area (Table 4). As a result, genus composition at B.South was relatively similar in 2007 and 2011, apart from some smaller differences (e.g., no *Metadesmolaimus* in 2011, Table 4). Also diversity and evenness remained fairly similar over the years (Table 2), with few significant differences (only $N_1$ and $J'$; Table 3: pairwise tests for factor year). On the contrary, diversity and evenness at B.West were even lower than in 2007 (and differences were always significant; Table 3), due to a profound increase in *Halomonhystera*, mainly in the upper two centimeters of sediment (Table 4; Fig. 2). Hence, as was the case in 2007, genus diversity and evenness remained highest at B.South.

Three-factor PERMANOVA, including sediment depth, showed that the differences in nematode assemblages between areas and years further depended on depth in the sediment (significant three-way interaction, $P < 0.05$; Table 5). Communities of both areas differed mostly in surface layers and became more similar with depth (Fig. 5A: pairwise differences between stations for all levels of factors “year” and “layer”). This trend was more obvious in 2007 since communities in 2011 were more distinct in almost all depth layers. Alternatively, communities of both years became more similar at B.West with increasing depth, while the opposite occurred at B.South (Fig. 5B: pairwise dissimilarities across all levels of factors “area” and “layer”). This means that nematode assemblages in deeper layers of B.South diverged over the years.

### Table 2. Overview of total nematode density (ind. 10 cm$^{-2}$), diversity ($N_0$, *EG*(200), $H'$, $N_1$), and evenness ($N_{	ext{tot}}$, $J'$), averaged for five replicates per area × year combination. Values in brackets represent standard deviation.

|       | Density (ind. 10 cm$^{-2}$) | $N_0$   | *EG*(200) | $H'$      | $N_1$       | $N_{	ext{tot}}$ | $J'$       |
|-------|-----------------------------|---------|-----------|-----------|-------------|-----------------|-----------|
| **2011** |                             |         |           |           |             |                 |           |
|       | B.South                     | 2547.8 (472.38) | 35.60 (4.56) | 24.13 (3.44) | 2.29 (0.32) | 10.23 (3.10) | 2.38 (0.51) | 0.64 (0.07) |
|       | B.West                      | 4832.24 (1038.26) | 10.80 (2.39) | 6.24 (1.35) | 0.40 (0.16) | 1.51 (0.26) | 1.09 (0.06) | 0.17 (0.05) |
| **2007** |                             |         |           |           |             |                 |           |
|       | B.South                     | 3075.94 (235.34) | 29.40 (1.52) | 24.63 (0.82) | 2.53 (0.08) | 12.57 (1.01) | 3.16 (0.41) | 0.75 (0.02) |
|       | B.West                      | 604.71 (63.03) | 20.80 (4.97) | 16.90 (4.36) | 1.57 (0.32) | 4.99 (1.58) | 1.89 (0.32) | 0.52 (0.06) |
while they increasingly resembled each other at B.West (due to the large Halomonhystera contribution in all sediment layers in 2011).

**Environmental setting**

Averaged environmental variables for each layer in both stations and years (where available) are given in Table 6 (data grouped over 0–3 and 3–5 cm for 2007; n (2007) = 5; n(2011) = 2). Silt was the dominant grain size for all layers at both locations. B.South had a slightly higher sand content in 2007, but only for the upper centimeters. The biggest difference was a significant increase in chl a from 2007 to 2011, for both B.South and B.West (Fig. 2). Chl a content was higher in surface layers (0–3 cm) than deeper down (3–5 cm). Chl a values in 2011 alone did not differ much between stations, only between sediment layers. B.South samples had about twice as much TOC and TOM than B.West, leading to a higher C: Nmolar as well (2011 only). BEST routine attributed 64% of 2011 nematode community variation to a combination of chl a and TOC.

**Broader geographic comparison**

Plotting of Larsen communities within a larger geographical context showed that, despite large dissimilarities observed within the area, communities differed substantially from those in other Antarctic shelf regions (Table 7, Fig. 6). Significant differences were found between all regions (R = 0.633, P < 0.05; one-way ANOSIM), but they were largest between B.West and the other locations. Pairwise differences between the Larsen B stations and the other areas decreased from 2007 to 2011 for B.South, but increased for B.West (data not shown). Differences with other regions were (mostly) due to the low abundance of Sabattieria and high abundance of Microlaimus for B.South, while they were mainly attributable to high contributions of the Monhysteridae (including Halomonhystera) and the absence of Sabattieria in the case of B.West (SIMPER).
Discussion

Large-scale ice-shelf disintegration is one of the many consequences of the rapid warming trend observed along the Antarctic Peninsula. Although most of the LIS-A/B disintegration occurred over a rather short time period (1995–2002), its effects will persist over a longer time span. Therefore, the aim of both ANT-XXIII/8 and ANT-XXVII/3 was to collect information at different time intervals for several components of the marine food web to be able to anticipate to future responses, and relate changes and patterns to the situation observed before (Gutt et al. 2011).

Environmental setting and implications for benthic communities

Four years after the first sampling campaign, rapid regional warming in Antarctic Peninsula surroundings continues, evoking additional ice-mass loss in the Larsen area (Shuman et al. 2011; Berthier et al. 2012). Consequently, seasonal phytoplankton blooms emerge (Barnes and Clarke 2011), further modifying benthic habitats at former ice-shelf locations from an oligotrophic to a more productive state. New organic matter production in the Larsen area was demonstrated by remote sensing of net primary productivity in 1997–2011 (Cape et al. 2014), and diatom siliceous frustules found in the upper two centimeters of the sediment (i.e., the layer corresponding to post-ice-shelf deposition; Sañé et al. 2013). Productivity in Larsen A and B is now as high as that for other Antarctic shelf locations and tightly linked to seasonal polynya dynamics (Cape et al. 2014). Average chl \( a \) values reported in 2011 surface sediments (Table 6) are indeed comparable to those found elsewhere on the Antarctic shelf (e.g., Fabiano and Danovaro 1999: 0.25–0.38 \( \mu g g^{-1} \) at 430–590 m in the Ross Sea; Vanhove et al. 2004: 0.36–0.52 \( \mu g g^{-1} \) at 750 m in the Weddell Sea). The fivefold to tenfold increase in sediment chl \( a \) compared to 2007 conceivably demonstrates higher productivity in the area as more time passed since ice-shelf collapse. However, considering a time lag between production in surface photic layers and transport of phytodetritus through the water column, summer-bloom chlorophyll could have already reached the seafloor and benthic communities in 2011 (late-summer sampling), but not in 2007 (early-summer sampling). Furthermore, primary production in the Larsen area depends heavily on the sporadic breakup of seasonal sea ice, which makes food supply to the benthos hardly predictable in space and time, especially in terms of the high interannual variability (Gutt et al. 2013).

As meiofaunal assemblages are tightly linked to fresh food input (Lins et al. 2014, 2015), it is almost inevitable that the transition to a more productive (yet still highly
seasonal) state will influence nematode communities (cf. TOC and chl \(a\) main explanatory variables in BEST results). Organic matter in surface marine sediments lies usually within the range of 0.1–5 wt%, of which the lower extreme (0.1–0.2 wt%) typically occurs in fine-grained sediments of well-oxygenated bathyal and abyssal depths, while average TOC values of 0.5–3 wt% dominate in deltas and on upper continental margins (Hedges and Oades 1997). Surface TOC content at B.West was thus relatively low compared to global means, while values at B.South were clearly higher, situated within the intermediate range, and comparable to values reported in other Antarctic studies at similar depths (0.2–0.75 wt%; Domack and Ishman 1997; Giordano et al. 1999).

Not only the quantity, but also the quality and source of food can influence benthic community composition. Due to the cold temperatures of Antarctic waters, phytodetritus degradation is slow, allowing its accumulation in sediment “foodbanks” (Smith et al. 2006, 2008;
Mincks et al. 2008). These foodbanks can sustain a rich benthic community throughout the year (especially in long winters), even when fresh input is lacking. In addition, phytoplankton supply to the seafloor in sub-ice zones is considerably lower than in open water owing to lower sedimentation rates (Post et al. 2007). Combining both phenomena (i.e., low degradation and sedimentation rates) and taking into account the closer connection of B.South to open water and phytodetritus input, a substantial foodbank could have developed at this site; and nematode assemblages could be feeding on organic matter that accumulated over the course of many years (cf. higher TOC and TOM; Table 6). In contrast, longer persisting ice cover at B.West prevented the establishment of an extensive foodbank, rendering communities highly dependent upon short pulses of fresh material after ice-shelf collapse (demonstrated by higher chl<sub>a</sub> values).

### 2007–2011 Nematode community change

The original high dissimilarity in nematode community composition between B.South and B.West in 2007 (Raes et al. 2010) was still evident 4 years later, and temporal changes in nematode assemblages were quite different for both stations (Table 7).

Density, diversity, and generic composition at B.South remained fairly similar over the years (Figs 2 and 4; Tables 2 and 4) and changed with depth into the sediment. The community was still dominated by *Microlaimus*, an epistratum-feeder (Wieser 1953) that is generally widespread in shallow and deep-sea habitats (Tita et al. 2002; Gambi et al. 2003; Vanhove et al. 2004; Sebastian et al. 2007; Van Gaever et al. 2009a; Portnova et al. 2010; Vanreusel et al. 2010b). This opportunistic genus often attains elevated abundance in deeper areas that are more organically enriched (Van Gaever et al. 2004, 2006, 2009a; Sebastian et al. 2007) or recently disturbed (e.g., after iceberg scouring; Lee et al. 2001), in which case it is considered a pioneering colonizer. As there were no signs of disturbance related to iceberg scouring at the time of sampling, the first explanation seems more likely. To reach current numbers at B.South, *Microlaimus* could have benefited from lateral advective food input from the Weddell Sea during ice-shelf cover,
complemented by increased levels of phytodetritus accretion after ice-shelf collapse. Even so, in spite of continued seasonal ice-free periods and enhanced food conditions, changes between 2007 and 2011 nematode assemblages at B.South were not very prominent, suggesting a relatively steady community, comparable to other Antarctic shelf areas in terms of abundance and biodiversity (Raes et al. 2010). Nevertheless, generic composition at B.South was not entirely comparable to that of other Antarctic shelf areas, mainly attributable to the genera *Sabatieria* and *Microlaimus* (SIMPER; Fig. 6). Only a few individuals of *Sabatieria* were observed at B.South, while it is usually quite common in shelf samples, especially in muddy sediments (as was the case in the Larsen; Schratzberger et al. 2009; Van Gaever et al. 2009a). It tends to reside in deeper sediment layers (associated with the Redox Potential Discontinuity (RPD) layer; Vanreusel et al. 2010a; Guilini et al. 2011), where a substantial fraction of organic material becomes incorporated below the oxic zone. Perhaps, Larsen sediment conditions were not yet favorable for *Sabatieria*, as organic matter burial was rather limited in the millennia preceding ice-shelf collision; or, alternatively, *Sabatieria* could not reach the area or establish a stable population within the 4 years time. Either way, the nematode community at B.South did not converge with other Antarctic shelf fauna as we hypothesized, although differences with other areas did decline over the years (pairwise ANOSIM).

Nematode assemblages at B.West were even more distinct from other shelf communities than at B.South (Fig. 6), as >90% of total abundance consisted of *Halomonhystera* (and *Sabatieria* was virtually absent). *Halomonhystera* is classified as a nonselective deposit feeder sensu Wieser (1953) and a general opportunistic genus (Bongers et al. 1991). Compared to the 2007 situation, densities increased drastically while diversity decreased due to proliferation of *Halomonhystera*. According to Raes et al. (2010), low density and low genus richness in 2007 reflected precollapse oligotrophic conditions. At that time, *Halomonhystera* was mainly found in deeper

Table 7. Location and depth range of references included in the (sub)-Antarctic database.

| Reference          | Publication year | Region                  | Broader area         | Depth range (m) | Collection year |
|--------------------|------------------|-------------------------|----------------------|-----------------|-----------------|
| Chen et al.        | 1999             | Beagle Channel, Magellan Strait | Magellan            | 10–550 m        | 1994            |
| Hauquier et al.    | 2011             | Larsen B                | Larsen 2007          | ~820 m          | 2007            |
| Hauquier et al.    | Unpublished      | Drake Passage, NE Weddell Sea | Peninsular           | 470–520 m       | 2013            |
| Ingels et al.      | 2006             | Signy Island, South Georgia | Peninsular           | ~300 m          | 2002            |
| Lee et al.         | 2001             | Kapp Norvægia           | Eastern Weddell      | 200–300 m       | 1998            |
| Lee et al.         | Unpublished      | Bransfield Strait, Drake Passage | Peninsular           | 200–430 m       | 1998            |
| Lee et al.         | Unpublished      | Kapp Norvægia, Vestkapp | Eastern Weddell      | ~200 m          | 1996            |
| Luyten             | 1999             | Adelaide Island          | Peninsular           | ~5–30 m         | 1998            |
| Manachini          | 1997             | Kapp Norvægia, Ross Sea | Eastern Weddell/Ross Sea | 200–600 m     | 1994, 1996      |
| Raes et al.        | 2010             | Unpublished             | Elephant Island      | ~430 m          | 2006            |
| Raes et al.        | 2010             | Larsen A, Larsen B      | Larsen 2007          | 240–430 m       | 2007            |
| Vanhove et al.     | 1997             | Kapp Norvægia, Halley Bay | Eastern Weddell      | 200–800 m       | 1989            |
| Vanhove et al.     | 1998             | Signy Island            | Peninsular           | ~10 m           | 1994            |
| Vanhove et al.     | 2004             | South Sandwich Trench  | Peninsular           | ~750 m          | 2002            |

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sediment layers (upper cm dominated by *Thalassomonhystera*), which generally contain less food. The drastic increase in *Halomonhystera* densities at station B.West over the course of only a few years is thus at least remarkable. One possible explanation is that increased direct supply of fresh food to the seabed has triggered opportunistic feeding behavior of *Halomonhystera*. Earlier research on one species of *Halomonhystera*, *H. disjuncta*, has classified it as an efficient colonizer, capable of expressing priority effects (Derycke et al. 2007b; Van Gaever et al. 2009b), a situation where first colonizing individuals have such a strong population development that they inhibit the settlement of other species. This could explain why community composition at B.West was still very different from B.South, even after a longer time period: *Microlaimus* and other genera potentially able to profit from open-water conditions do not get a chance to settle in the *Halomonhystera*-dominated sediments (provided that they did reach the area though; see further). Alternatively, it is possible that *Halomonhystera* is responding to sedimentary features other than fresh phytodetritus input. In fact, the subsurface (1–2 cm) maximum in *Halomonhystera* abundance strongly resembles the vertical profile observed at station Larsen B.Seep reported by Hauquier et al. (2011), where a low-active cold seep was found (~800 m; Niemann et al. 2009). Also there, nematode assemblages were characterized by high densities, deeper density maxima, and high dominance of one *Halomonhystera* species. This prompted the question whether *Halomonhystera* depended upon chemosynthetically derived organic matter, as was the case with *Halomonhystera hermesi* (earlier identified as *H. disjuncta*) in sulfidic, microbial mat sediments at the Håkon Mosby Mud Volcano (~1300 m; Van Gaever et al. 2006). However, stable isotope data for B.Seep did not indicate such a relationship, leading to the conclusion that *Halomonhystera* thrives on phytoplanktonic rather than chemosynthetic resources (Hauquier et al. 2011). The fact that there were no signs of elevated sulfide levels, anoxia or seepage at the time of sampling at B.West further strengthens this conclusion.

Whatever the reason or mechanism behind it, the success of *Halomonhystera* at B.West in 2011 further isolated the community from B.South (and by extension any other Antarctic shelf region) compared to 2007. Instead of anticipated convergence of communities at both stations, they increasingly diverged from each other.

**Nematode colonization dynamics**

Besides food availability as a local, environmental driver for differences between areas, also more regional processes such as colonization ability of organisms can structure benthic communities. Marine nematode dispersal is dependent on body morphology, swimming ability, and feeding strategies (Thomas and Lana 2011), and as nematodes lack pelagic larvae or propagules, dispersal is in this case synonymous to gene flow (Derycke et al. 2013). It was already shown that nematode colonization is a slow process (Post et al. 2007), predominantly driven by passive transport via bottom currents (Boeckner et al. 2009); and not necessarily related to higher food input (e.g., Guilini et al. 2011). Furthermore, colonization dynamics depend on the distance (Derycke et al. 2007a), proximity of a source population, and the time needed for successful settlement and reproduction (Schratzberger et al. 2006; Raes et al. 2010). Closer connection of B.South to the open Weddell Sea as a source for new recruits may therefore partly explain observed differences with B.West. Raes et al. (2010) calculated a speed of recovery of 60.8 m year$^{-1}$, and hence, approximately 1000 years needed to cross the distance of 70.8 km between B.West and B.South. So far, too little time has passed for the nematodes to travel between both Larsen stations on one hand and between larger geographical areas on the other hand. With time, and if local conditions allow it, maybe *Microlaimus* will be able to reach the innermost part of the Larsen B, and maybe other genera, such as *Sabatieria*, will find their way into Larsen.

**Comparison with other benthic groups**

Nematodes are only one taxonomic player in the Antarctic marine benthic food web, and it can be valuable to compare their response with other food-web compartments, as changes at one trophic level may impact other faunal components (either bottom-up or top-down) or remineralization processes in the sediment (e.g., Moline et al. 2004; Montes-Hugo et al. 2009). As already shown in 2007 (Gutt et al. 2011), different benthic components react in different ways to the ice-shelf collapse, each at their own pace (some organisms are more sensitive to disturbance, especially long-lived taxa such as Porifera). Results on other trophic levels for the 2011 expedition remain scarce so far, but Gutt et al. (2013) found a drastic decrease in the aggregations of two ascidians between 2007 and 2011 but an increase in abundances of deposit-feeding ophiuroids. Although they could not relate their findings to particular environmental characteristics, it clearly shows the high dynamics of Antarctic benthos and the probability for both negative and positive effects to arise after large-scale alterations. Together with this study, their research highlights the difficulties to relate changes in faunal communities to environmental factors because benthic responses may take a long time and are highly variable.
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Conflict of Interest

None declared.

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