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Foraminiferal community response to seasonal anoxia in Lake Grevelingen (the Netherlands)

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Abstract. Over the last decades, hypoxia in marine coastal environments has become more and more widespread, prolonged and intense. Hypoxic events have large consequences for the functioning of benthic ecosystems. In severe cases, they may lead to complete anoxia and the presence of toxic sulfides in the sediment and bottom-water, thereby strongly affecting biological compartments of benthic marine ecosystems. Within these ecosystems, benthic foraminifera show a high diversity of ecological responses, with a wide range of adaptive life strategies. Some species are particularly resistant to hypoxia–anoxia, and consequently it is interesting to study the whole foraminiferal community as well as species-specific responses to such events. Here we investigated the temporal dynamics of living benthic foraminiferal communities (recognised by CellTracker™ Green) at two sites in the saltwater Lake Grevelingen in the Netherlands. These sites are subject to seasonal anoxia with different durations and are characterised by the presence of free sulfide (H₂S) in the uppermost part of the sediment. Our results indicate that foraminiferal communities are impacted by the presence of H₂S in their habitat, with a stronger response in the case of longer exposure times. At the deepest site (34 m), in summer 2012, 1 to 2 months of anoxia and free H₂S in the

surface sediment resulted in an almost complete disappearance of the foraminiferal community. Conversely, at the shallower site (23 m), where the duration of anoxia and free H₂S was shorter (1 month or less), a dense foraminiferal community was found throughout the year except for a short period after the stressful event. Interestingly, at both sites, the foraminiferal community showed a delayed response to the onset of anoxia and free H₂S, suggesting that the combination of anoxia and free H₂S does not lead to increased mortality, but rather to strongly decreased reproduction rates. At the deepest site, where highly stressful conditions prevailed for 1 to 2 months, the recovery time of the community takes about half a year. In Lake Grevelingen, *Elphidium selseyense* and *Elphidium magellanicum* are much less affected by anoxia and free H₂S than *Ammonia* sp. T6. We hypothesise that this is not due to a higher tolerance for H₂S, but rather related to the seasonal availability of food sources, which could have been less suitable for *Ammonia* sp. T6 than for the elphidiids.

1 Introduction

Hypoxia affects numerous marine environments, from the open ocean to coastal areas. Over the last decades, a general decline in oxygen concentration was observed in marine waters (Stramma et al., 2012), with an extent varying between the concerned regions. In coastal areas, oxygen concentrations have been estimated to decrease 10 times faster than in the open ocean, with indications of a recent acceleration, expressed by increasing frequency, intensity, extent and duration of hypoxic events (Diaz and Rosenberg, 2008; Gilbert et al., 2010). This is due to the combination of (1) global warming, which is strengthening seasonal stratification of the water column and decreasing oxygen solubility, and (2) eutrophication resulting from increased anthropogenic nutrient and/or organic matter input, which is enhancing benthic oxygen consumption in response to increased primary production (Diaz and Rosenberg, 2008). Bottom-water hypoxia has serious consequences for the functioning of all benthic ecosystem compartments (see Riedel et al., 2016, for a review). Benthic faunas are strongly impacted by these events (Diaz and Rosenberg, 1995), even though the meiofauna, especially foraminifera, appears to be less sensitive to low dissolved oxygen (DO) concentrations than the macrofauna (e.g. Josefson and Widbom, 1988). Many foraminiferal taxa are able to withstand seasonal hypoxia–anoxia (see Koho et al., 2012, for a review), and consequently they can play a major role in carbon cycling in ecosystems affected by seasonal low-oxygen concentrations (Woulds et al., 2007). Anoxia is often accompanied by free sulfide (H_2S) in pore and/or bottom waters (e.g. Jørgensen, 1982; Seitaj et al., 2015), which is considered very harmful for the benthic macrofauna (Wang and Chapman, 1999). Neutral molecular H_2S can diffuse through cellular membranes and inhibits the functioning of cytochrome *c* oxidase (a mitochondrial enzyme involved in ATP production), finally inhibiting aerobic respiration (Nicholls and Kim, 1982; Khan et al., 1990; Dorman et al., 2002).

Lake Grevelingen (southwestern Netherlands) is a former branch of the Rhine–Meuse–Scheldt estuary, which was closed in its eastern part (riverside) by the Grevelingen Dam in 1964 and in its western part (seaside) by the Brouwers Dam in 1971. The resulting saltwater lake, with a surface of 115 km², is one of the largest saline lakes in western Europe. Lake Grevelingen is characterised by a strongly reduced circulation (even after the construction of a small sluice in 1978) with a strong thermal stratification occurring in the main channels in summer, leading to seasonal bottom-water hypoxia–anoxia in late summer and early autumn (Bannink et al., 1984). This situation results in a rise of the H_2S front in the uppermost part of the sediment, sometimes up to the sediment–water interface.

These observations especially concern the Den Osse Basin (i.e. one of the deeper basins, maximum depth 34 m; Hagens et al., 2015), which has been intensively monitored over the

last decades, so that a large amount of environmental data are available (e.g. Wetsteijn, 2011; Donders et al., 2012). The annual net primary production in the Den Osse Basin (i.e. 225 g C m^{−2} yr^{−1}; Hagens et al., 2015) is comparable to other estuarine systems in Europe (Cloern et al., 2014). However, there is almost no nutrient input from external sources; thus primary production is largely based on autochthonous recycling (> 90 %; Hagens et al., 2015), both in the water column and in the sediment, with a very strong pelagic–benthic coupling (de Vries and Hopstaken, 1984). The benthic environment is characterised by the presence of two antagonistic groups of bacteria, with contrasting seasonal population dynamics (i.e. cable bacteria in winter–spring and *Beggiatoaceae* in autumn–winter), which have a profound impact on all biogeochemical cycles in the sediment column (Seitaj et al., 2015; Sulu-Gambari et al., 2016a, b). The combination of hypoxia–anoxia with sulfidic conditions, which is rather unusual in coastal systems without external nutrient input, and the activity of antagonistic bacterial communities makes Lake Grevelingen a very peculiar environment. In the Den Osse Basin, seasonal anoxia coupled with the presence of H_2S at or very close to the sediment–water interface occurs in summer (i.e. between July–September). However, euxinia (i.e. diffusion of free H_2S in the water column) does not occur, because of cable bacterial activity (Seitaj et al., 2015).

Although the tolerance of foraminifera towards low DO contents and long-term anoxia (from weeks to 10 months) has been well documented for many species from different types of environments in laboratory culture (e.g. Moodley and Hess, 1992; Alve and Bernhard, 1995; Bernhard and Alve, 1996; Moodley et al., 1997; Duijnsteet et al., 2003, 2005; Geslin et al., 2004, 2014; Ernst et al., 2005; Pucci et al., 2009; Koho et al., 2011) as well as in field studies (e.g. Piña-Ochoa et al., 2010b; Langlet et al., 2013, 2014), their tolerance of free H_2S is still debated. In the vast majority of previous studies, no decrease in the total abundances of living foraminifera (i.e. strongly increased mortality) was observed during anoxic events. Unfortunately, studies on foraminiferal response in systems affected by seasonal hypoxia–anoxia with sulfidic conditions are still very sparse. The few available observations are not conclusive, but they suggest that H_2S could be toxic for foraminifera even on fairly short timescales (Bernhard, 1993; Moodley et al., 1998b; Panieri and Sen Gupta, 2008; Langlet et al., 2014).

To our knowledge, all earlier studies show that the foraminiferal response to hypoxia–anoxia is species-specific (e.g. Bernhard and Alve, 1996; Ernst et al., 2005; Bouchet et al., 2007; Geslin et al., 2014; Langlet et al., 2014). However, this species-specific response generally follows the same scheme (usually decrease in density, reduction of growth and/or reproduction), with different response intensities. Duijnsteet et al. (2005) suggested that oxic stress leads to an increased mortality and inhibited growth and reproduction. The suggestion of inhibited growth is supported by LeKieffre et al. (2017), who observed that the morphos-

pecies *Ammonia tepida* (probably *Ammonia* sp. T6) showed minimal or no growth under anoxia. Conversely, Geslin et al. (2014) and Nardelli et al. (2014) suggested that, in the same morphospecies, reproduction was strongly reduced, but growth would not be affected by hypoxic and/or short anoxic events. Additionally, under low-oxygen conditions, some species are able to shift to anaerobic metabolism (i.e. denitrification; Risgaard-Petersen et al., 2006; Piña-Ochoa et al., 2010a), to sequester chloroplast (i.e. kleptoplastidy; Jauffrais et al., 2018), to associate with bacterial symbionts (Bernhard et al., 2010) or to enter into a state of dormancy (Ross and Hallock, 2016; LeKieffre et al., 2017).

The highly peculiar environmental context of Lake Grevelingen offers an excellent opportunity to study this still poorly known aspect of foraminiferal ecology.

The conventional method to discriminate between live and dead foraminifera uses Rose Bengal, a compound which stains proteins (i.e. organic matter). This method was proposed for foraminifera by Walton (1952) and is based on the assumption that “the presence of protoplasm is positive indication of a living or very recently dead organism”. The author already noted that this assumption implied that the rate of degradation of organic material should be relatively high. Previous studies of living benthic foraminifera in environments subjected to hypoxia–anoxia were almost all based on Rose Bengal-stained samples (e.g. Gustafsson and Nordberg, 1999, 2000; Duijnsteet et al., 2004; Panieri, 2006; Schönfeld and Numberger, 2007; Polovodova et al., 2009; Papaspyrou et al., 2013). However, foraminiferal protoplasm may remain stainable from several weeks to months after their death (Corliss and Emerson, 1990), especially under low dissolved oxygen concentrations where organic matter degradation may be very slow (Bernhard, 1988; Hannah and Rogers, 1997; Bernhard et al., 2006). The Rose Bengal staining method is therefore not suitable for studies in environments affected by hypoxia–anoxia. Consequently, the results of foraminiferal studies in low-oxygen environments based on this method have to be considered with reserve. In order to avoid this problem, we used CellTracker™ Green (CTG) to recognise living foraminifera. CTG is a fluorescent probe which marks only living individuals with cytoplasmic (i.e. enzymatic) metabolic activity (Bernhard et al., 2006). Since metabolic activity stops after the death of the organism, CTG should give a much more accurate assessment of the living assemblages at the various sampling times and thereby avoid overestimation of the live foraminiferal abundances.

In this study, samples were collected in August and November 2011 and then every month through the year 2012, at two different stations in the Den Osse Basin, with two replicates dedicated to foraminifera. The two stations were chosen in contrasted environments regarding water depth (34 and 23 m, respectively) and duration of seasonal hypoxia–anoxia and sulfidic conditions. Living foraminiferal assemblages were studied in the uppermost sediment and size distributions were determined in order to get insight

into the possible moment(s) of reproduction or accelerated growth in test size. The seasonal variability study of the foraminiferal community allows us (1) to better understand the foraminiferal tolerance of seasonal hypoxia–anoxia with the presence of free H_2S in their microhabitat and (2) to obtain information about the responses of the various species to adverse conditions. This knowledge will be useful for the development of indices assessing environmental quality (i.e. biomonitoring) and may also improve palaeoecological interpretations of coastal records (e.g. Murray, 1967; Gustafsson and Nordberg, 1999).

2 Material and methods

2.1 Studied area – environmental settings in the Den Osse Basin

Lake Grevelingen is a part of the former Rhine–Meuse–Scheldt estuary, in the southwestern Netherlands. This former estuarine branch was turned into an artificial saltwater lake during the Delta Works project. In Lake Grevelingen, the water circulation is strongly limited by the construction of dams (in the early 1970s) and only a small sluice allows water exchanges with open seawater (i.e. very weak hydrodynamics). In the lake, development of bottom-water hypoxia–anoxia occurs in the deepest part of the basin in summer (i.e. July–September) to early autumn (i.e. October–December; Bannink et al., 1984; Hagens et al., 2015). In the literature, the terminology and threshold values used to describe oxygen depletion are highly variable (e.g. oxic, dysoxic, hypoxic, suboxic, microxic, postoxic; see Jorissen et al., 2007; Altenbach et al., 2012). In this study we defined hypoxia as a concentration of oxygen $< 63 \mu\text{mol L}^{-1}$ (1.4 mL L^{-1} or 2 mg L^{-1}) whereas anoxia is defined as no detectable oxygen (following Rabalais et al., 2010).

In Den Osse Basin, the nutrient input from external sources is very low and pelagic–benthic coupling is essential, as already noted by de Vries and Hopstaken (1984). In 2012, phytoplankton blooms occurred in April–May and July (Hagens et al., 2015) in response to the increasing solar radiation and nutrient availability in the water column following organic matter recycling in winter. This led to an increased food availability in the benthic compartment in the same periods. In general, Chl *a* concentrations in Den Osse Basin are below $10 \mu\text{g L}^{-1}$, excluding very short peaks during blooms in April–May and July which did not exceed $30 \mu\text{g L}^{-1}$ in 2012 (Hagens et al., 2015). Thermal stratification of the water column and increased oxygen consumption due to organic matter input (i.e. from phytoplankton blooms) are both responsible for the development of seasonal bottom-water hypoxia–anoxia in summer (i.e. July–September). Although euxinia (i.e. the presence of free H_2S in the water column) does not occur in the Den Osse Basin due to cable bacterial activity in winter, free H_2S is present in the uppermost layer of the

sediment in summer (Seitaj et al., 2015). Summarising, in the benthic ecosystem, increased food availability in summer is counterbalanced by strongly decreasing oxygen contents, sometimes accompanied by the presence of free sulfides in the topmost sediment.

2.2 Field sampling

The two studied sites are located along a depth gradient in the Den Osse Basin of Lake Grevelingen. Both station 1 (51°44.834' N, 3°53.401' E) and station 2 (51°44.956' N, 3°53.826' E) are located in the main channel, at 34 and 23 m depth, respectively (Fig. 1).

Measurements of bottom-water oxygen (BWO) concentrations were performed at 2 m above the sediment–water interface and are from Donders et al. (2012), whereas the data for 2012 were published in Hagens et al. (2015). Sediment cores were collected monthly in 2012 using a single core gravity corer (UWITEC, Austria) using PVC core liners (6 cm inner diameter, 60 cm length). All cores were inspected upon retrieval and only visually undisturbed sediment cores were used for further analysis (Seitaj et al., 2017). Oxygen penetration depth (OPD) and depth of free H₂S detection were determined by Seitaj et al. (2015) using profiling microsensors for station 1. The data for station 2 (Supplement Table S1) were acquired similarly and during the same cruises but never published; for further details about the sampling method, see Seitaj et al. (2015).

Two replicate sediment cores dedicated to the foraminiferal study were sampled in August and November 2011 using the same gravity corer (UWITEC, Austria) and then monthly throughout the year 2012 at the same sampling time as for BWO concentration and OPD and H₂S measurements in the sediment (see Seitaj et al., 2015). Consequently, for 2012 at stations 1 and 2, OPD and H₂S were measured in the sediment column at the same time as foraminifera were sampled (Seitaj et al., 2015). For each replicate, the uppermost centimetre (0–1 cm) of the core was then transferred on board in a vial of 250 mL, and 30 mL of seawater (at the same temperature as in situ) was added to the vial. Then we labelled the samples with CellTracker™ Green CMFDA (CTG, 5-chloromethylfluorescein diacetate, final concentration of 1 µmol L⁻¹ following Bernhard et al., 2006) and slowly agitated manually to allow the CTG diffusion in the whole sample. Samples were then fixed in 5 % sodium-borate-buffered formalin after 24 h of incubation in the dark.

2.3 Sample treatment

All samples were sieved over 315, 150 and 125 µm meshes, and foraminiferal assemblages were studied in all three size fractions. Individuals were picked wet under an epifluorescence stereomicroscope (Olympus SZX12, light fluorescent source Olympus URFL-T, excitation/emission wavelengths:

Table 1. Sampling dates of the samples which were investigated for living foraminifera for stations 1 and 2. X: one core investigated; O: no core investigated.

Year	Month	Day	Station 1	Station 2
2011	Aug	22	X X	X X
2011	Nov	15	X X	X X
2012	Jan	23	X X	X X
2012	Mar	12	X X	X X
2012	May	30	X X	X X
2012	Jul	24	X X	X X
2012	Sep	20	X X	X X
2012	Oct	18	O	X X
2012	Nov	2	X X	X X
2012	Dec	3	O	X X

492 nm/517 nm) and placed on micropalaeontological slides. Only specimens that fluoresced brightly green were considered living and were identified to the (morpho)species level when possible. Since picking foraminifera under an epifluorescence stereomicroscope is particularly time-consuming, we decided to study samples only every 2 months for the year 2012. At a later stage, in view of the large differences in foraminiferal abundances between the samples of September and November 2012 at station 2, we decided to study the October and December 2012 samples as well for this station. The sampling dates investigated in this study are listed in Table 1.

Abundances were then standardised to a volume of 10 cm³. The abundances of living foraminifera for each sampling time and replicate are listed in Tables S2 and S3. The mean abundance and standard deviation ($\bar{x} \pm \text{SD}$) for the two replicates for each sampling date were calculated for both the total living assemblage and the individual species, as an indication of spatial patchiness.

2.4 Taxonomy of dominant species

Four dominant species (>1 % of the total assemblage) were present in our material: *Ammonia* sp. T6, *Elphidium magellanicum* (Heron-Allen and Earland, 1932), *Elphidium selseyense* (Heron-Allen and Earland, 1911) and *Trochammina inflata* (Montagu, 1808). As we identified these species on the basis of morphological criteria, we will use them as “morphospecies”.

Concerning the genus *Ammonia*, two living specimens collected at Grevelingen station 1 were molecularly identified (by DNA barcoding) as phylotype T6 by Bird et al. (2019). At the same site, we genotyped seven other living *Ammonia* specimens, which were all T6. Their sequences were deposited in GenBank (accession numbers MN190684 to MN190690), and Supplement Fig. S1 shows scanning electron microscope (SEM) images of the spiral side and of the penultimate chamber at 1000× magnification for four indi-

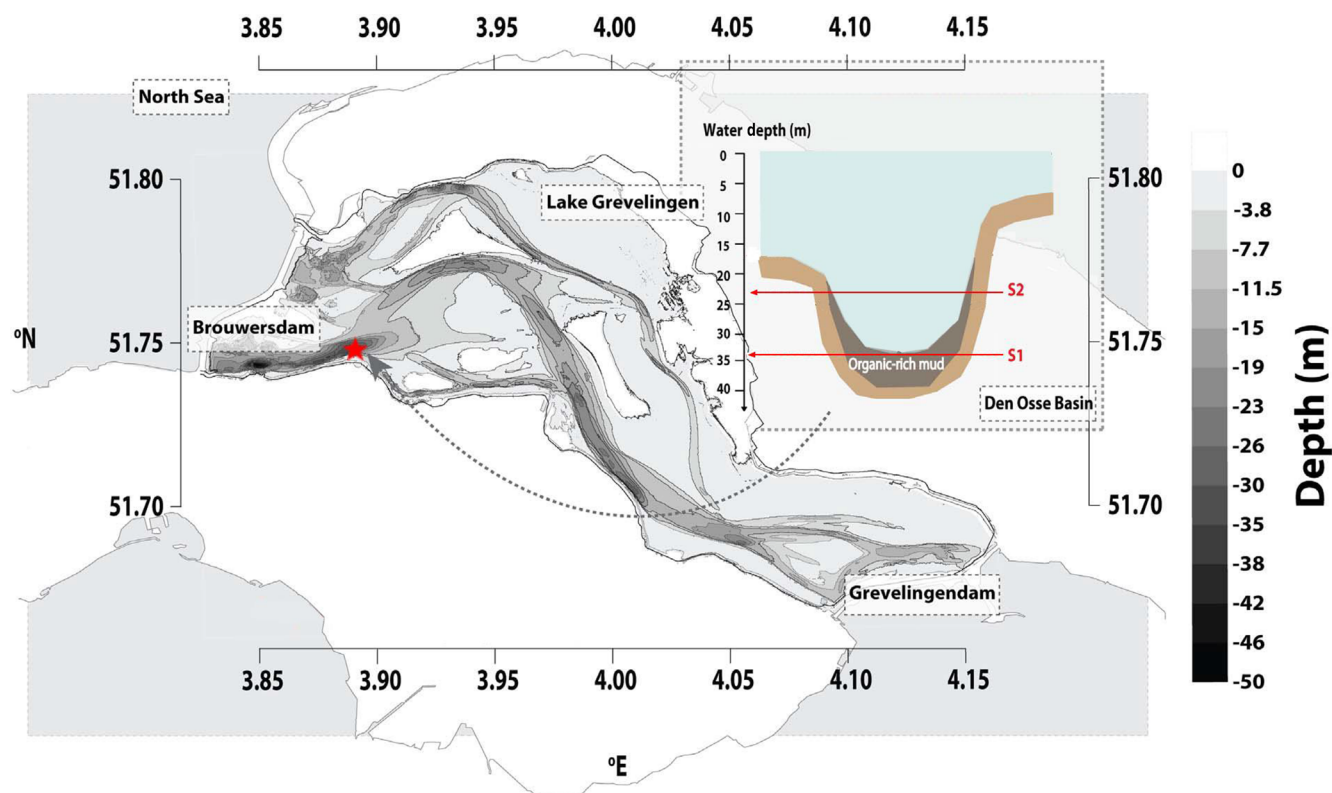


Figure 1. Map of Lake Grevelingen showing the location of the two sampled stations in the Den Osse Basin (red star). The transversal section of the Den Osse Basin (top right) shows the depth at which station 1 (S1) and station 2 (S2) were sampled (34 and 23 m depth, respectively). This figure was modified from Sulu-Gambari et al. (2016b).

viduals. A morphological screening based on the criteria proposed by Richirt et al. (2019) confirmed that T6 accounts for the vast majority (>98 %) of *Ammonia* individuals, whereas phylotypes T1, T2, T3 and T15 are only present in very small amounts (Table S3).

The specimens of *Elphidium magellanicum* were identified exclusively on the basis of morphological criteria, as there are no molecular data available yet. This morphospecies, although rare, is regularly recognised in Boreal and Lusitanian provinces of Europe (e.g. Gustafsson and Nordberg, 1999; Darling et al., 2016; Alve et al., 2016). However, as the type species was described from the Strait of Magellan (Southern Chile), the European specimens may represent a different species and further studies involving DNA sequencing of both populations are needed to confirm or disprove this taxonomic attribution (see Roberts et al., 2016).

Elphidium selseyense has often been considered an ecophenotype of *Elphidium excavatum* (Terquem, 1875) and has been identified as *E. excavatum* forma *selseyensis* (e.g. Feyling-Hanssen, 1972; Miller et al., 1982). Four living specimens were already sampled for DNA analysis at station 1 and were all identified as the species *E. selseyense* (phylo-type S5, Darling et al., 2016). We only observed minor morphological variations in our material, especially concerning

the number of small bosses in the umbilical region, which we considered to be intraspecific variability. Consequently, we identified all our specimens as *E. selseyense*.

The specimens attributed to *Trochammina inflata* were also identified exclusively on the basis of morphological criteria, as no molecular data are available yet.

2.5 Size distribution measurement

In order to detect periods of increased growth and/or reproduction, size measurements were performed on all samples of 2012. The measurements were made for all species (4176 individuals for station 1 and 19624 individuals for station 2), and trochospiral species were all orientated spiral side up prior to measurements. High-resolution images (3648 pixels × 2736 pixels) of all micropalaeontological slides were taken with a stereomicroscope (Leica S9i, 10× magnification) and individual measurements were processed using ImageJ software (Schneider et al., 2012, Fig. S2).

Each individual was isolated (Fig. S2) and its maximum diameter was measured (i.e. Feret's diameter). We represented all size distributions using histograms with 20 µm classes (the best compromise between the total number of individuals and the size range (Fig. S3)). As we only examined the size fractions >125 µm, our analysis mainly concerns

adult specimens and does not include juveniles. This limitation should be kept in mind when interpreting the results.

Assuming that the size distribution was a sum of Gaussian curves, each of them representing a cohort, we tried to identify the approximate mode for the Gaussian curves (i.e. cohorts) using the changes in slope (i.e. inflexion points) of the second-order derivative of the total size distribution (Gammón et al., 2017). Unfortunately, this tentative attempt to distinguish cohorts by using a deconvolution method was not conclusive. The main problem was the lack of information concerning individuals smaller than 125 µm, so that our size distributions were systematically skewed toward small individuals. Because the identification of individual cohorts was not successful, a study of population dynamics was not possible. For this reason, the data are only shown in Figs. S2 and S3. Nevertheless, the size distribution data give some clues concerning the possible moment(s) of reproduction or intensified test growth for the different species.

2.6 Encrusted forms of *E. magellanicum*

In our samples, we found abundant encrusted forms of *E. magellanicum* at station 1 (May 2012) and station 2 (May, July, September and December 2012, Fig. 2). Most individuals were totally encrusted (Fig. 2a), others only partly (Fig. 2b). These crusts were hard, firmly stuck to the shell (difficult to remove with a brush), thin (Fig. 2c–e) and rather coarse. In order to determine if the crust matrix is constituted of carbonate, we placed some specimens in microtubes and exposed them to 0.1 M of EDTA (ethylenediaminetetraacetic acid) diluted in 0.1 M cacodylate buffer (acting as a carbonate chelator). After an exposition of 24 h, we checked under a stereomicroscope if the crust was still cohesive (no carbonate in the crust) or was disaggregated (crust contains carbonate).

3 Results

3.1 Total abundances of foraminiferal assemblages

Averaged total abundances varied between 1.1 ± 1.5 and 449.9 ± 322.1 ind. 10 cm^{-3} for station 1 and between 91.1 ± 25.0 and 604.8 ± 3.5 ind. 10 cm^{-3} for station 2 (Fig. 3 and Table 2). For every studied month, the total density was higher at station 2 than at station 1. The seasonal succession is very different between the two sites (Fig. 3). Station 1 shows very low total foraminiferal abundances for most months, contrasting with much higher densities in May and July. Conversely, station 2 shows high total foraminiferal abundances throughout the year, with somewhat lower values in November 2011 and October and November 2012 (Fig. 3).

At station 1, almost no individuals were present in August ($\bar{x} = 3.4 \pm 1.3$) and November 2011 ($\bar{x} = 1.1 \pm 1.5$). In 2012, total abundances were very low in January ($\bar{x} = 11.5 \pm 9.3$), showed a slight increase in March ($\bar{x} = 62.1 \pm 19.3$) and reached a maximal abundance in May ($\bar{x} = 449.9 \pm 322.1$).

Total abundances then progressively decreased from May to September ($\bar{x} = 34.0 \pm 17.0$) and almost no foraminifera were present in November ($\bar{x} = 1.6 \pm 0.3$).

At station 2, total abundances were comparatively low in August and November 2011 ($\bar{x} = 174.0 \pm 48.0$ and $\bar{x} = 128.7 \pm 25$ ind. 10 cm^{-3} , respectively). In 2012, total abundances were relatively high and stable from January to September (between $\bar{x} = 523.6 \pm 30.7$ and $\bar{x} = 604.8 \pm 3.5$), then decreased in October ($\bar{x} = 211.5 \pm 8.0$) and November ($\bar{x} = 91.1 \pm 25.3$), and finally increased again in December ($\bar{x} = 377.9 \pm 38.8$).

3.2 Dominant species

At station 1, the major species were, in order of decreasing abundances, *Elphidium selseyense* (Fig. 4a–b), *Elphidium magellanicum* (Fig. 4c–d) and *Ammonia* sp. T6 (Fig. 4e–g). In Fig. 4, we added *Trochammina inflata* (Fig. 4h–j) to facilitate comparison with station 2, where this species is among the dominant ones. The “other species” account only for 2.2 % of the total assemblage at station 1. The fact that they are well represented in some months (e.g. 26.3 % of the assemblage in August 2011) is due to the extremely low number of individuals (see Fig. 3 and Table 2). At station 2, the dominant species, in order of decreasing abundances, were *E. selseyense*, *Ammonia* sp. T6, *E. magellanicum* and *T. inflata* (Table 2). Here, “other species” account only for 2.6 % of the total assemblage. Whereas *E. selseyense* and *E. magellanicum* were dominant species at both stations, both *Ammonia* sp. T6 and *T. inflata* were present in much higher abundances at station 2 compared to station 1, where the latter species was almost absent (Figs. 5–6).

At station 1, only some very scarce individuals of *E. selseyense* were observed in August and November 2011 (Fig. 5 and Table 2). In 2012, *E. selseyense* abundances were very low in January and started to increase in March ($\bar{x} = 23.9 \pm 6.8$), reaching maximal values in May ($\bar{x} = 336.5 \pm 275.8$). In July, values for *E. selseyense* were still high ($\bar{x} = 162.0 \pm 121.5$) and further decreased until an almost total absence in November 2012. No specimen of *E. magellanicum* was observed in 2011 (Fig. 5 and Table 2). The abundance of *E. magellanicum* was very low in January 2012, started to increase in March ($\bar{x} = 21.6 \pm 11.0$), reaching maximal values in May ($\bar{x} = 96.4 \pm 47.3$), and then strongly decreased in July ($\bar{x} = 3.7 \pm 0.3$). The species was absent from samples in September and November 2012. *Ammonia* sp. T6 was almost absent in August and November 2011 and present with very few specimens in January 2012 ($\bar{x} = 3.2 \pm 3.5$). Maximum abundances were reached between March and July 2012 (ranging between $\bar{x} = 9.2 \pm 6.5$ and $\bar{x} = 12.9 \pm 1.3$). Then abundances rapidly decreased until the species was almost absent in November. *Trochammina inflata* was absent in 2011 and was only present in very low abundances from January to May and in September 2012.

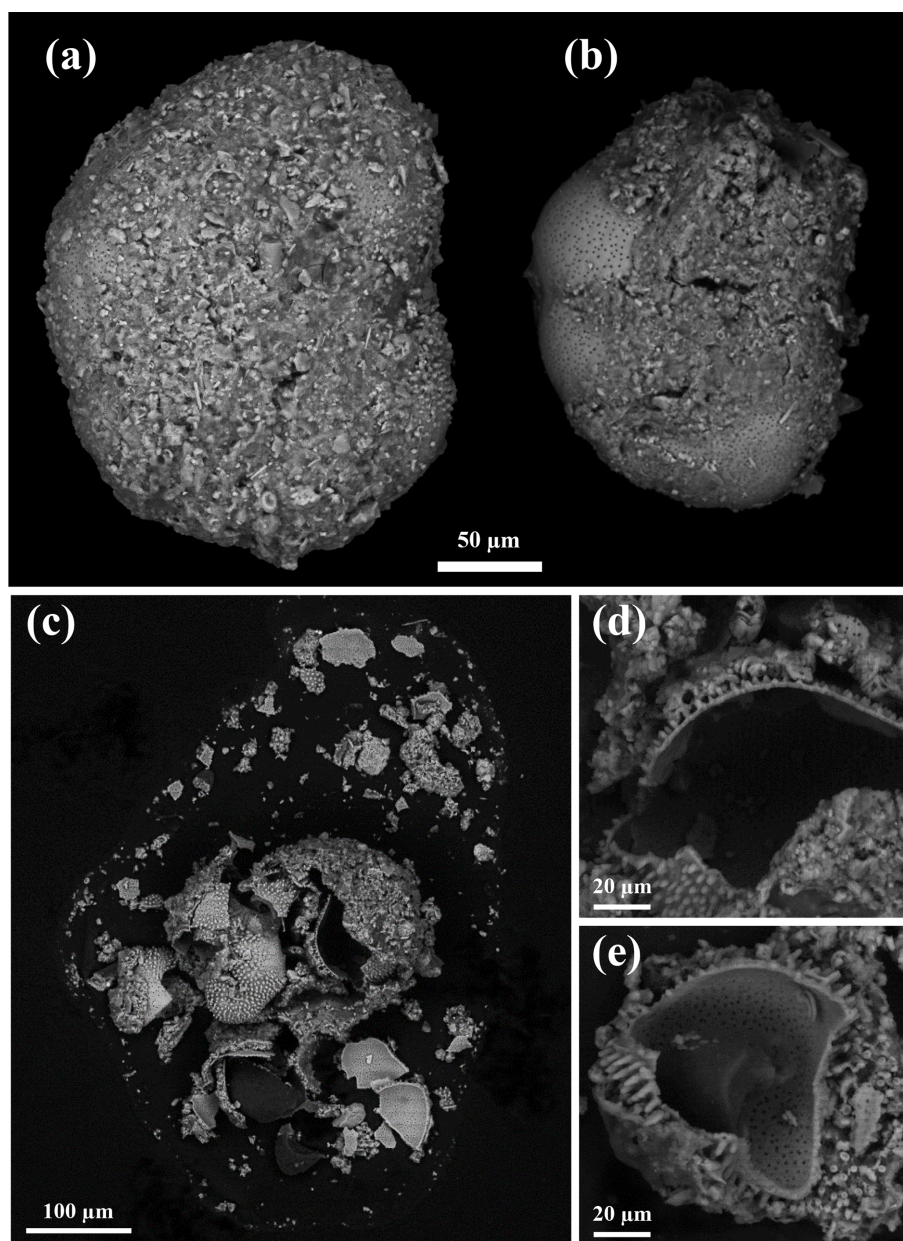


Figure 2. SEM images of fully encrusted specimen (a), partially encrusted specimen (b) and crushed encrusted specimen of *Elphidium magellanicum* (c). Note the thinness of the crust and the spinose structures in (d) and (e).

At station 2, the two dominant species were *E. selseyense* and *Ammonia* sp. T6, which together always represented at least 70 % of the total assemblage (Fig. 6 and Table 2). These two species showed a different seasonal pattern over the considered period. Abundances of *E. selseyense* were comparable in August ($\bar{x} = 74.8 \pm 29.8$) and November 2011 ($\bar{x} = 52.3 \pm 27.0$) and then showed a progressive increase until a maximum in September 2012 ($\bar{x} = 365.5 \pm 70.3$). Abundances then showed a sharp decrease in October and November (respectively $\bar{x} = 98.7 \pm 8.5$ and $\bar{x} = 30.9 \pm 2.3$) to increase again in December ($\bar{x} = 252.2 \pm 41.0$). For *Ammo-*

nia sp. T6, abundances strongly increased between November 2011 ($\bar{x} = 60.8 \pm 1.5$) and January 2012 ($\bar{x} = 226.2 \pm 52.3$) and then progressively decreased until the end of 2012 ($\bar{x} = 48.1 \pm 26.0$ in November 2012). *Trochammina inflata* showed an analogous pattern to *Ammonia* sp. T6. Abundances strongly increased between November 2011 ($\bar{x} = 11.8 \pm 1.8$) and January 2012 ($\bar{x} = 121.5 \pm 29.8$) and then progressively decreased until very low abundances in November ($\bar{x} = 3.7 \pm 3.0$). *E. magellanicum* was completely absent in August and November 2011, almost absent in January 2012 ($\bar{x} = 0.9 \pm 0.3$), and then suddenly increased until a max-

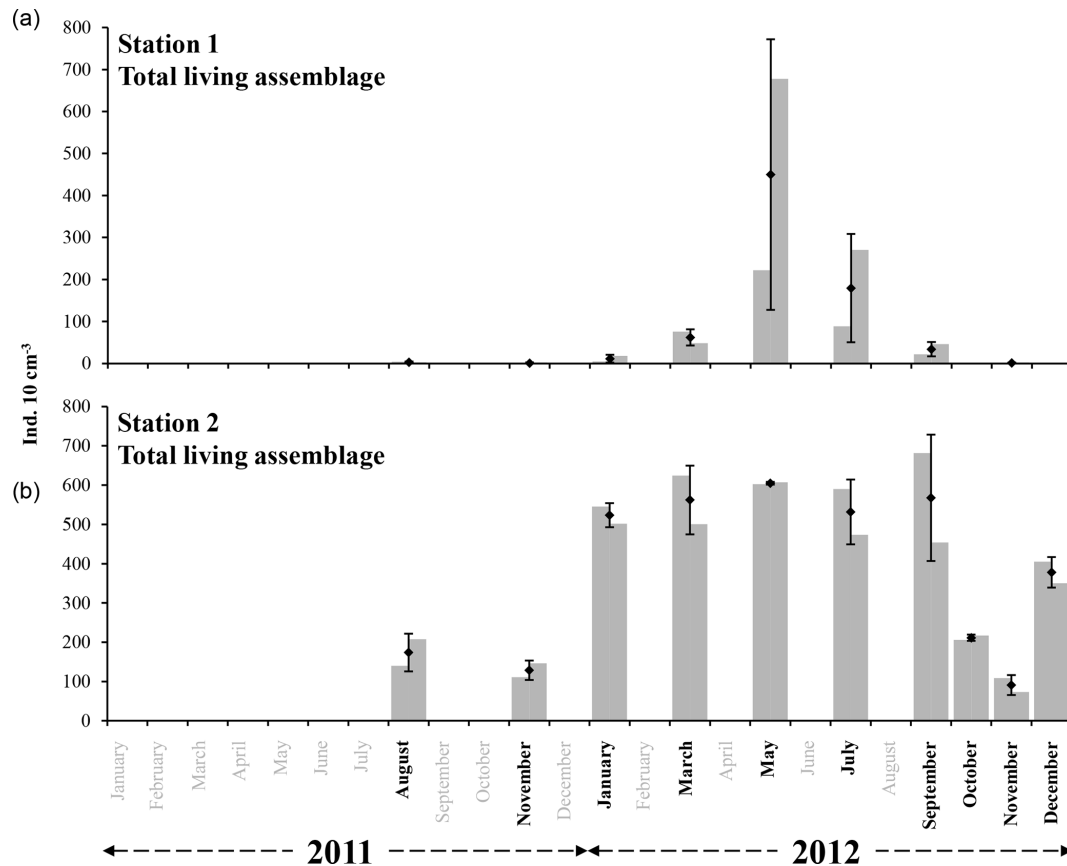


Figure 3. The grey bars represent the living foraminiferal abundances for the two replicates. The mean abundances (diamonds) and standard deviations (black error bars) were calculated for the two replicates for stations 1 (34 m depth, **a**) and 2 (23 m depth, **b**). All abundance values are for the 0–1 cm layer and were standardised to 10 cm³. Months where foraminiferal communities were investigated are indicated in bold (excluding October and December at station 1).

imum of $\bar{x} = 116.0 \pm 6.5$ in May. Abundances stayed relatively high in July ($\bar{x} = 37.8 \pm 2.5$) and September ($\bar{x} = 72.0 \pm 35.8$) and then drastically decreased until minimum numbers in October and November. Finally, like all other species, *E. magellanicum* abundances increased again in December ($\bar{x} = 25.5 \pm 13.0$).

3.3 Encrusted forms of *Elphidium magellanicum*

After exposition to 0.1 M of EDTA diluted in 0.1 M cacodylate buffer, the crusts remained cohesive, indicating that they do not consist of carbonate and suggesting that they are composed of sediment particles cemented by an organic matrix.

At station 1, encrusted forms of *E. magellanicum* were present in moderate proportions in May (26.8 % of the total *E. magellanicum* population, Fig. 7) and July (47.6 %); the species disappeared thereafter. At station 2, encrusted forms strongly dominated the *E. magellanicum* population from May (72.3 %) to December (88.0 %, Fig. 7).

4 Discussion

4.1 Tolerance of foraminiferal communities towards anoxia and free sulfide

At station 1, bottom waters were hypoxic in July 2012 and became anoxic in August (Fig. 8). Both in July and in August, oxygen penetration into the sediment was null, whereas it was 0.7 ± 0.1 mm depth in September. In all 3 months (July to September 2012), sulfidic conditions were observed very close to the sediment–water interface (1 mm or less, Fig. 8 and Table S1). In view of these results, the duration of anoxic and sulfidic conditions in the uppermost sediment layer can be estimated as 1 to 2 months (in July and August, Fig. 8).

After the strong increase in foraminiferal densities in May 2012, there was a decrease starting in July, leading to a near absence of foraminifera at station 1 in November (Fig. 8). The most probable cause of the strong decline of the foraminiferal community appears to be a prolonged presence of sulfides in the foraminiferal microhabitat. However, the fact that foraminiferal abundances reached almost

Table 2. Mean living foraminiferal absolute (ind. 10 cm⁻³) and relative abundances (percentage of the total fauna, in parentheses) of the dominant species. Last column: absolute abundance of the total fauna.

Year	Month	<i>Elphidium selseyense</i>	<i>Ammonia</i> sp. T6	<i>Elphidium magellanicum</i>	<i>Trochammina inflata</i>	Others	Total
Station 1							
2011	Aug	1.2 (36.8)	1.2 (36.8)	0.0 (0.0)	0.0 (0.0)	0.9 (26.3)	3.4
2011	Nov	0.5 (50.0)	0.4 (33.3)	0.0 (0.0)	0.0 (0.0)	0.2 (16.7)	1.1
2012	Jan	5.1 (44.6)	3.2 (27.7)	0.2 (1.5)	1.2 (10.8)	1.8 (15.4)	11.5
2012	Mar	23.9 (38.5)	12.9 (20.8)	21.6 (34.8)	1.4 (2.3)	2.3 (3.7)	62.1
2012	May	336.5 (74.8)	9.2 (2.0)	96.4 (21.4)	1.8 (0.4)	6.0 (1.3)	449.9
2012	Jul	162.0 (90.2)	10.3 (5.7)	3.7 (2.1)	0.0 (0.0)	3.5 (2.0)	179.5
2012	Sep	29.7 (87.5)	2.3 (6.8)	0.0 (0.0)	0.4 (1.0)	1.6 (4.7)	34.0
2012	Nov	1.1 (66.7)	0.4 (22.2)	0.0 (0.0)	0.0 (0.0)	0.2 (11.1)	1.6
Sum		560.0 (75.4)	39.8 (5.4)	121.8 (16.4)	4.8 (0.6)	16.4 (2.2)	742.9
Station 2							
2011	Aug	74.8 (43.0)	82.1 (47.2)	0.0 (0.0)	14.7 (8.4)	2.5 (1.4)	174.0
2011	Nov	52.3 (40.7)	60.8 (47.3)	0.0 (0.0)	11.8 (9.2)	3.7 (2.9)	128.7
2012	Jan	161.8 (30.9)	226.2 (43.2)	0.9 (0.2)	121.5 (23.2)	13.3 (2.5)	523.6
2012	Mar	214.7 (38.2)	214.0 (38.1)	48.8 (8.7)	75.0 (13.3)	9.9 (1.8)	562.3
2012	May	288.2 (47.7)	147.1 (24.3)	116.0 (19.2)	36.1 (6.0)	17.3 (2.9)	604.8
2012	Jul	282.6 (53.2)	158.4 (29.8)	37.8 (7.1)	31.5 (5.9)	21.2 (4.0)	531.6
2012	Sep	365.5 (64.4)	102.4 (18.0)	72.0 (12.7)	16.1 (2.8)	11.5 (2.0)	567.5
2012	Oct	98.7 (46.7)	99.0 (46.8)	1.8 (0.8)	7.4 (3.5)	4.6 (2.2)	211.5
2012	Nov	30.9 (34.0)	48.1 (52.8)	4.1 (4.5)	3.7 (4.1)	4.2 (4.7)	91.1
2012	Dec	252.2 (66.7)	78.0 (20.6)	25.5 (6.7)	12.7 (3.4)	9.5 (2.5)	377.9
Sum		1821.8 (48.3)	1216.1 (32.2)	306.8 (8.1)	330.5 (8.8)	97.7 (2.6)	3773.0

zero only in September (about 2 months after the first occurrence of anoxic and sulfidic conditions in the upper sediment, in July) suggests that the presence of H₂S did not cause instantaneous mortality, but that the disappearance of the foraminiferal community was a delayed response, probably caused by inhibited reproduction and, eventually, increased mortality. Inhibited reproduction has previously been suggested as a response to hypoxic–short anoxic (Geslin et al., 2014) and sulfidic conditions (Moodley et al., 1998b).

Such a time lag between a change in foraminiferal abundances and changes in environmental parameters affecting reproduction and/or growth of foraminifera has been suggested previously by Duijnste et al. (2004). These authors highlighted that the density patterns of some foraminiferal species showed a higher correlation with measured environmental parameters (e.g. oxygenation or temperature) when a time lag of about 3 months was applied.

For 2011, at station 1, no pore-water O₂ and H₂S measurements are available. However, severe hypoxia was observed in the bottom waters from May to August, with anoxia in June 2011 (Fig. 8). We therefore assume that, like in 2012, anoxic and probably co-occurring sulfidic conditions were responsible for the very low standing stocks in August and November 2011 and January 2012.

Our observations confirm the suggestion in previous studies that the foraminiferal community is severely affected by a long-term presence of H₂S in its habitat but does not show instant mortality. In fact, after a 66 d incubation in euxinic conditions (a maximum of $11.9 \pm 0.4 \mu\text{mol L}^{-1}$ of H₂S in the overlying water) of foraminiferal assemblages collected at a 19 m deep site in the Adriatic Sea, Moodley et al. (1998a) found a strong decrease in the total density of Rose Bengal-stained foraminifera. After 21 d, living specimens were still observed, whereas after 42 and 66 d, the live checks (based on protoplasm movement) gave only negative results. Langlet et al. (2013, 2014) performed an in situ experiment with closed benthic chambers at a 24 m deep site in the Gulf of Trieste, in the Adriatic Sea. They observed a decrease in living foraminiferal density (labelled with CTG), but they also found that almost all species survived after 10 months of anoxia and periodically co-occurring H₂S in the sediment and overlying water. However, the duration of sulfidic conditions, which was estimated to be several weeks, could not be assessed precisely (Metzger et al., 2014). The suggestion that short exposure to euxinic conditions is not directly lethal for foraminifera is confirmed by the experimental results of Bernhard (1993), who found that foraminiferal activity (as determined by ATP content) was not significantly affected after 30 d exposure to euxinia ($32.6 \pm 8.6 \%$ of active indi-

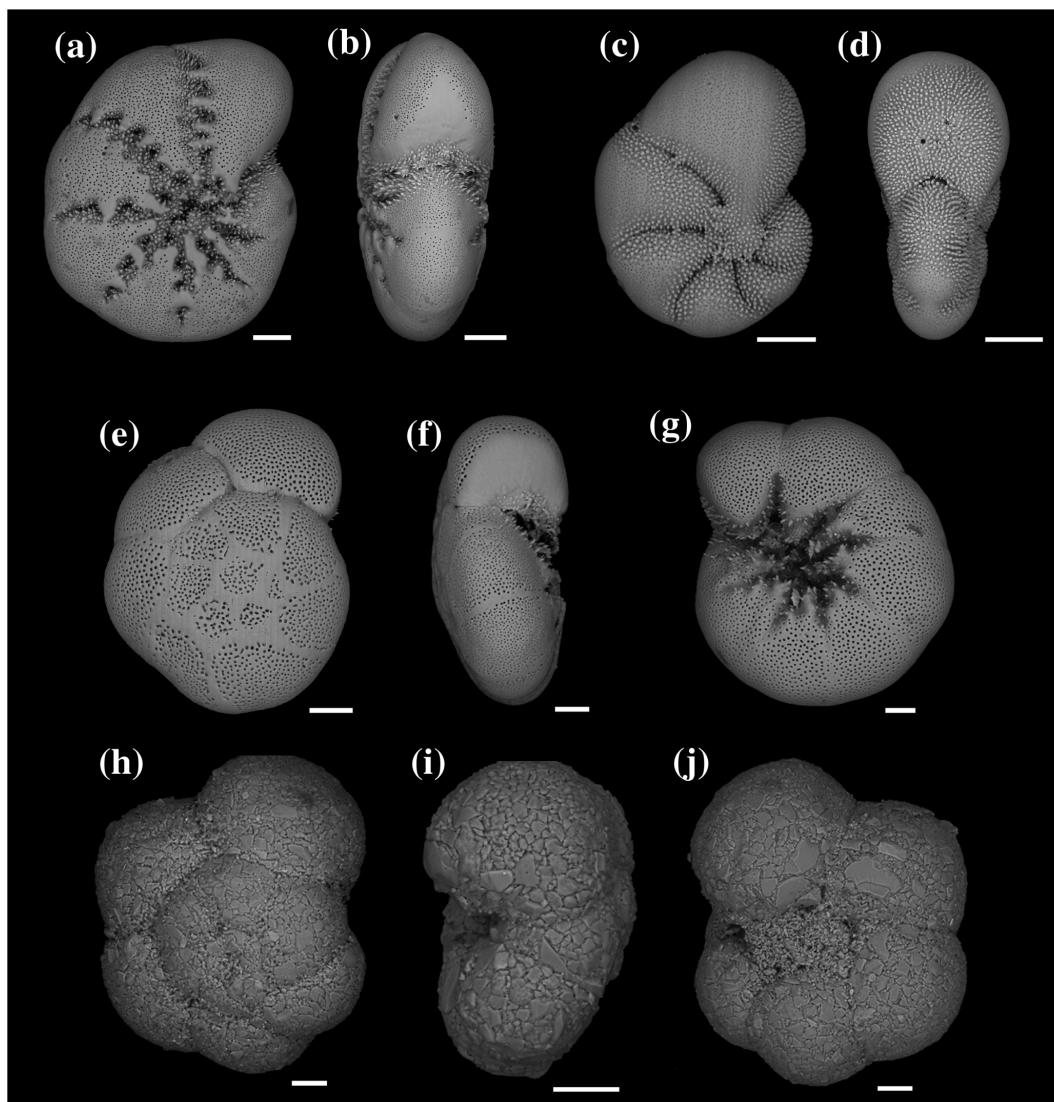


Figure 4. SEM images of *Elphidium selseyense* in lateral (a) and peripheral (b) views; *Elphidium magellanicum* in lateral (c) and peripheral (d) views; *Ammonia* sp. T6 in spiral (e), peripheral (f) and umbilical (g) views; and *Trochammina inflata* in spiral (h), peripheral (i) and umbilical (j) views. All scale bars are 50 µm.

viduals, $n = 174$ in control conditions versus $29.5 \pm 6.2\%$, $n = 173$ in sulfidic conditions).

After the 2011 hypoxia–anoxia, standing stocks at station 1 only started to increase in March 2012, indicating a very long recovery time (about 6 months) of the foraminiferal faunas after a temporary near-extinction due to anoxic and sulfidic conditions. This confirms observations of relatively long recovery times in the literature (e.g. Alve, 1995, 1999; Gustafsson and Nordberg, 2000; Hess et al., 2005). For instance, Gustafsson and Nordberg (1999) showed that in the Koljö Fjord, at comparable water depths, foraminiferal populations responded with increased densities only 3 months after a renewal of sea-floor oxygenation following hypoxic conditions in the bottom waters. However, in that case, the

disappearance of the foraminiferal population was only partial and not nearly as complete as in our study.

At station 2, in 2012, hypoxia was only observed in August, when the OPD was zero, and sulfidic conditions were observed in the superficial sediment (i.e. from 0.4 ± 0.2 mm downwards, Fig. 9, Table S1). Both in July and in September, oxygen penetrated more than 1 mm into the sediment (1.3 ± 0.4 and 1.2 ± 0.2 mm, respectively). However, free H_2S was still detected at about 1 mm depth in the sediment (1.1 ± 0.8 mm in July and 0.8 ± 0.2 mm in September). Although the sampling plan does not allow us to be very precise about the duration of anoxic and sulfidic conditions, we can estimate their duration to be 1 month or less (Fig. 9).

Station 1

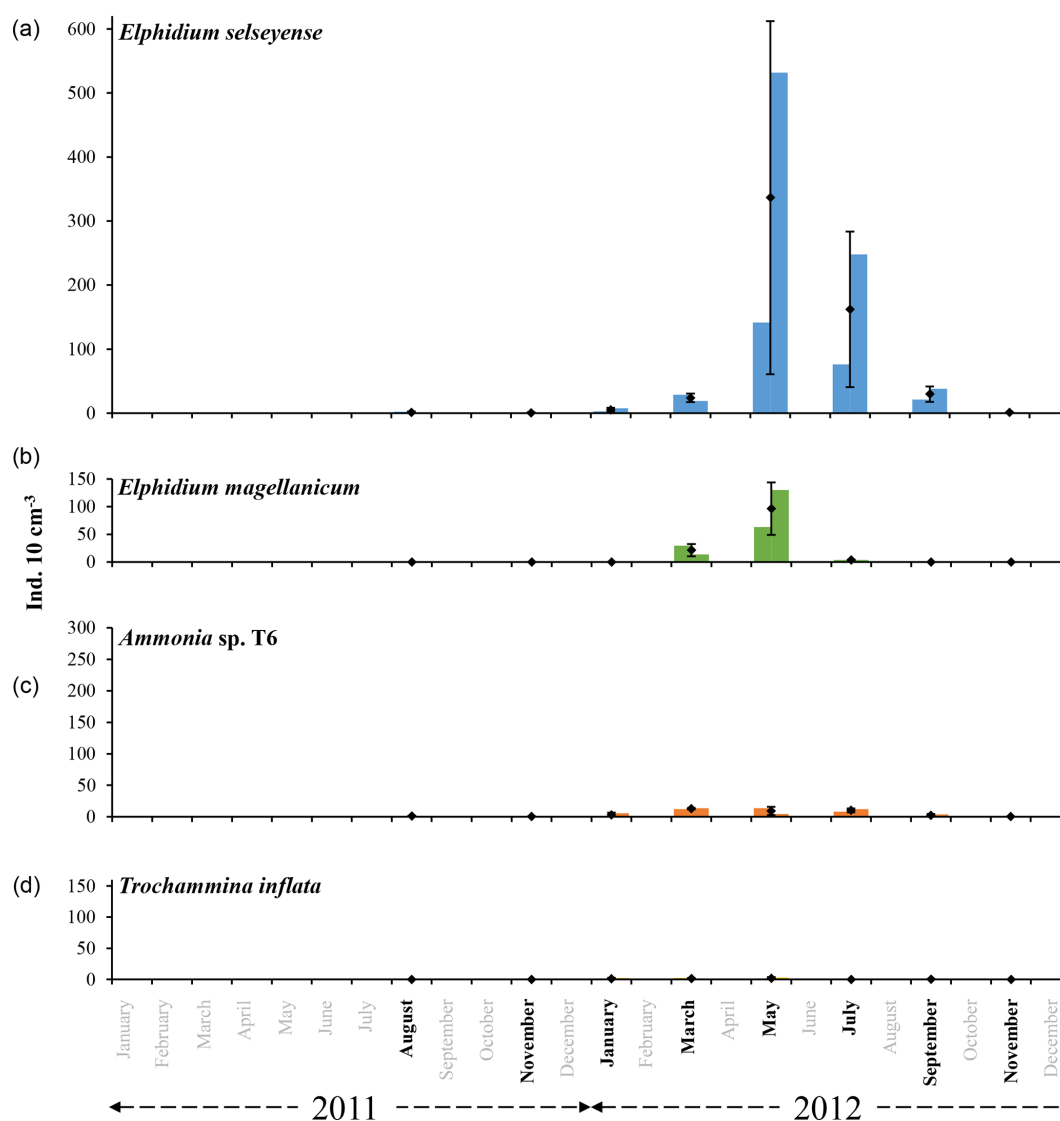


Figure 5. The bars represent the living foraminiferal abundances for the two replicates for *Elphidium selseyense* (a), *Elphidium magellanicum* (b), *Ammonia* sp. T6 (c) and *Trochammina inflata* (d) at station 1 in 2011 and 2012. The mean abundances (diamonds) and standard deviations (black error bars) were calculated for the two replicates. All abundance values are for the 0–1 cm layer and were standardised to 10 cm³. Months when foraminiferal communities were investigated are indicated in bold. Scales were chosen in order to facilitate comparison with station 2.

Foraminiferal abundances showed a strong decrease in October and November 2012, about 2 months after the presence of anoxic and sulfidic conditions in the topmost part of the sediment (Fig. 9). Like at station 1, this temporal offset between the presence of anoxia–sulfidic conditions at station 2 (in August) and the strong decrease in faunal densities may be explained as a delayed response, mainly due to inhibited reproduction during the anoxic–sulfidic event. If true, the mortality of adults did not strongly increase in the months following the H₂S production in the uppermost sediment.

Nevertheless, there was no replacement in the > 125 µm fraction by growing juveniles, probably because reproduction was interrupted when H₂S was present in the foraminiferal microhabitat. A renewed recruitment after the last stage of sulfidic conditions somewhere in September would then explain why the faunal density in the > 125 µm fraction increased again in December 2012 (Fig. 9).

In 2011, at station 2, bottom waters oscillated between hypoxic and oxic conditions between May and August (Fig. 9). Although we have no measurements of H₂S in the pore

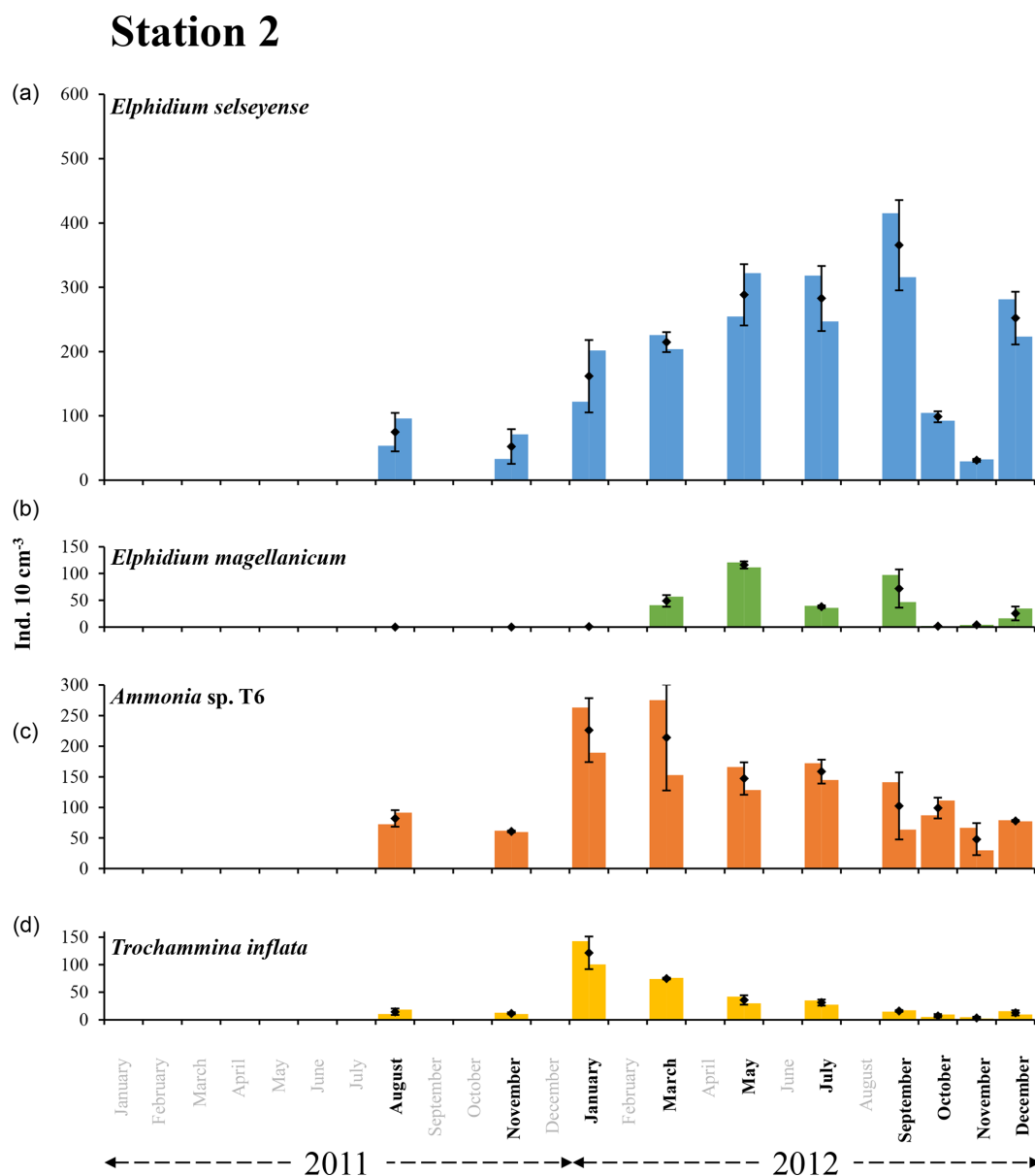


Figure 6. The bars represent the living foraminiferal abundances for the two replicates for *Elphidium selseyense* (a), *Elphidium magellanicum* (b), *Ammonia* sp. T6 (c) and *Trochammina inflata* (d) at station 2 in 2011 and 2012. The mean abundances (diamonds) and standard deviations (black error bars) were calculated for the two replicates. All abundances values are for the 0–1 cm layer and were standardised to 10 cm³. Months when foraminiferal communities were investigated are indicated in bold. Scales were chosen in order to facilitate comparison with station 1.

waters for this year, it seems probable that bottom-water hypoxia was accompanied by the presence of free H₂S very close to the sediment surface, strongly affecting the foraminiferal communities. If we assume that, like in 2012, rich foraminiferal fauna was present in May–July 2011 at both stations, the low faunal densities observed in August and November 2011 could suggest that foraminifera may have also shown a delayed response to sulfidic conditions in 2011.

It is interesting to note that the foraminiferal densities observed at station 2 were lower in August 2011 than in July or September 2012. This may be a consequence of the repetition of short hypoxic events in the bottom water between May and August 2011 (probably associated with anoxia and maybe H₂S in the uppermost part of the sediment), which possibly affected the foraminiferal community more substantially in 2011 than in 2012, when a hypoxic event was recorded in August only.

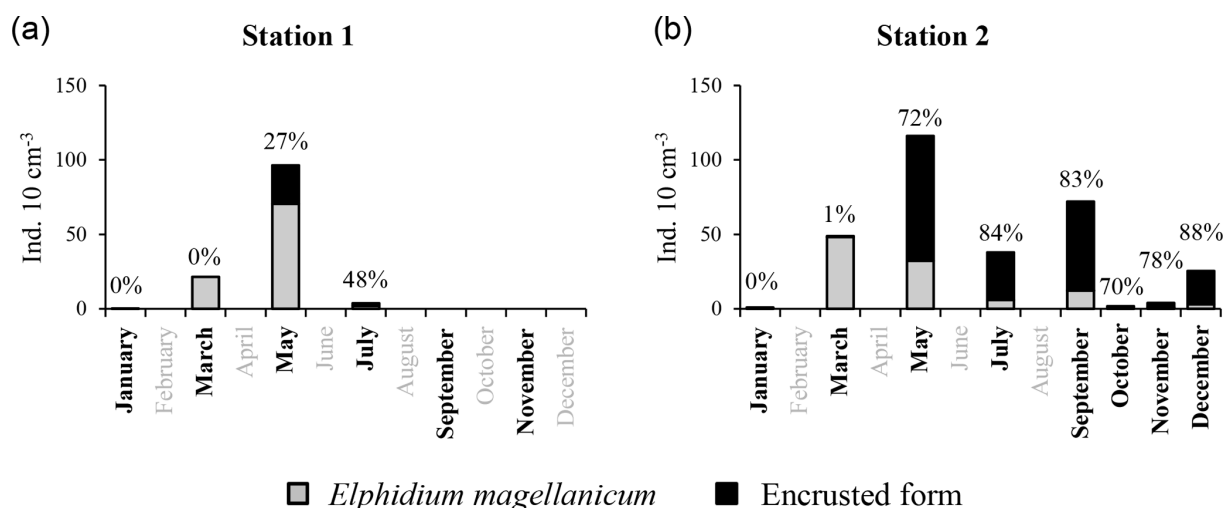


Figure 7. Mean abundances (ind. 10 cm⁻³) of non-encrusted (grey) and encrusted forms (black) of *Elphidium magellanicum* in 2012, at stations 1 (a) and 2 (b), with proportion of encrusted forms above each bar (%). Investigated months are indicated in bold.

The important decrease in total standing stocks at station 2 in October and November 2012 (Fig. 9) suggests that, in spite of the shorter duration of anoxia and sulfide conditions (compared to station 1; 1 month or less compared to 1 to 2 months), the foraminiferal faunas were still strongly affected. However, at station 2, foraminiferal abundances increased again in December 2012, suggesting a recovery time of about 2 months, which is likely much shorter than at station 1, where standing stocks in the >125 µm fraction only increased 6 months after the presence of anoxia and free sulfides.

Summarising, the foraminiferal communities of both stations 1 and 2 seem strongly impacted by the anoxic and sulfidic conditions developing in the uppermost part of the sediment in summer (i.e. July–September). However, at station 1, where anoxic and sulfidic conditions lasted for 1 to 2 months, the response is much stronger, leading ultimately (in November) to almost complete disappearance of the foraminiferal fauna. The delayed response at both stations shows that instantaneous mortality was limited and suggests that the decreasing standing stocks might rather be the result of inhibited reproduction and, eventually, increased mortality. Recovery is much faster at station 2 (about 2 months) than at station 1 (about 6 months), probably because at station 1 (in contrast to station 2) the foraminiferal extinction was nearly complete, and the site had to be recolonised (e.g. possibly by nearby sites or by the remaining few individuals) after reoxygenation of the sediment. At station 2, a reduced but significant foraminiferal community remained present, explaining the faster recovery.

4.2 Species-specific response to anoxia, sulfide and food availability in Lake Grevelingen

The comparison of the different seasonal patterns of the major species at the two investigated stations allows us to draw some conclusions about interspecific differences in the response to seasonal anoxic and sulfidic conditions.

First, there is a clear faunal difference between the two stations. Station 1 is dominated by *E. selseyense* and *E. magellanicum* while at station 2 these two taxa are accompanied by *Ammonia* sp. T6 and *T. inflata*. The latter species is almost absent at station 1, where *Ammonia* sp. T6 is present with low densities. At first glance, the dominance of the two *Elphidium* species at station 1 would suggest that they have a greater tolerance of the seasonal anoxic and sulfidic conditions, which lasted much longer there. It is interesting to note that the temporal evolution of standing stocks at station 1 is different for the two *Elphidium* species. *Elphidium magellanicum* shows a strong drop in absolute density in July 2012, at the onset of H₂S presence in the uppermost part of the sediment, whereas the diminution of *E. selseyense* is more progressive and the species disappears almost completely only in November (Fig. 5). This strongly suggests that *E. magellanicum* is more affected by increased mortality than *E. selseyense* in response to the combined effects of anoxic and sulfidic conditions. This hypothesis is confirmed by the patterns observed at station 2, where the drop in standing stocks in October–November is also more drastic in *E. magellanicum* than in *E. selseyense* (Fig. 6).

As mentioned earlier, certain species of foraminifera can use an anaerobic metabolism (i.e. denitrification; Risgaard-Petersen et al., 2006; Piña-Ochoa et al., 2010a), sequester chloroplasts (i.e. kleptoplastidy; Jauffrais et al., 2018), host bacterial symbionts (Bernhard et al., 2010) or enter dor-

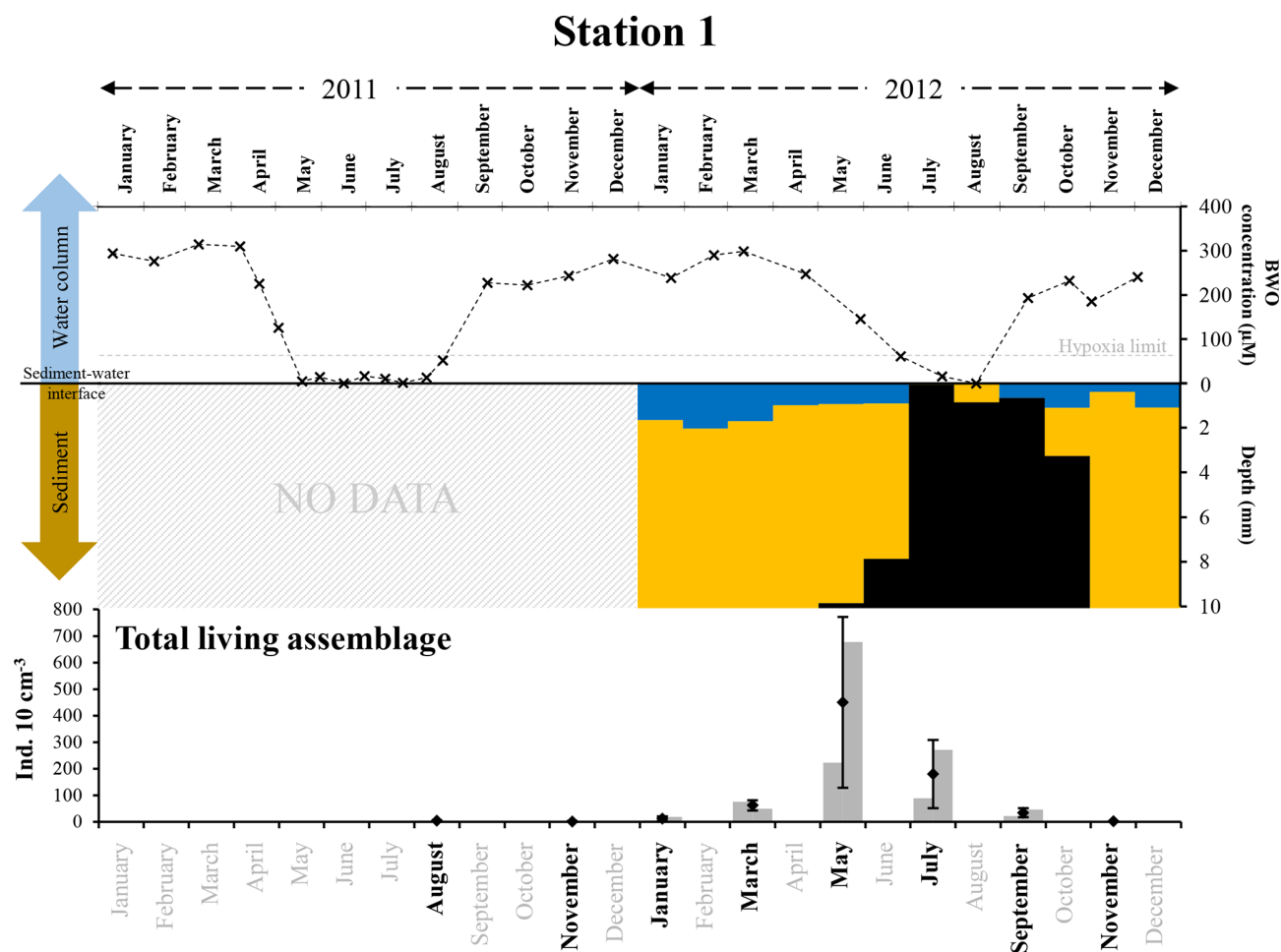


Figure 8. The top panel represents bottom-water oxygen concentrations ($\mu\text{mol L}^{-1}$) in 2011 and 2012 at station 1, from Donders et al. (2012) and Seitaj et al. (2017). The grey horizontal dotted line indicates the hypoxia limit ($63 \mu\text{mol L}^{-1}$). The middle panel represents the depth (mm) distribution of the oxic zone (blue), absence of oxygen and sulfides (orange), and sulfidic zone (black) within the sediment in 2012, from Seitaj et al. (2015). The bottom panel shows the total living foraminiferal abundances for both replicates (grey bars), mean abundances (diamonds) and standard deviations (black error bars) calculated for the two replicates, for all investigated months (in bold) in 2011 and 2012.

mancy (Ross and Hallock, 2016; LeKieffre et al., 2017) to deal with low-oxygen conditions. Concerning the species found in this study, although the presence of intracellular nitrate was shown for *Ammonia*, denitrification tests yielded negative results (Piña-Ochoa et al., 2010a; Nomaki et al., 2014). Similarly, the presence of active symbionts was previously suggested for *Ammonia* but never confirmed (Nomaki et al., 2016; Bernhard et al., 2018). To our knowledge, denitrification or the presence of bacterial symbionts was never shown for *Elphidium* either. In conclusion, a shift to an alternative anaerobic metabolism or an association with bacterial symbionts has never been shown conclusively for the dominant foraminiferal species found in Lake Grevelingen.

The greater tolerance of *E. selseyense* towards low-oxygen conditions could be explained by the fact that it is able to sequester chloroplasts from ingested diatoms and keep

them active for several days to weeks, in contrast to *Ammonia* sp. T6 (Jauffrais et al., 2018). These active chloroplasts could serve as an alternative source of oxygen and/or food through photosynthesis (Bernhard and Alve, 1996) or another metabolic pathway (Jauffrais et al., 2019) and thereby increase the capability of this species to survive anoxic events. Although sequestration of chloroplasts was never investigated for *E. magellanicum*, its abundant spinose ornamentation in the umbilical region and in the vicinity of the aperture (Fig. 4c–d) suggests that this species is capable of crushing diatom frustules like some kleptoplastic species (Bernhard and Bowser, 1999; Austin et al., 2005). Hagens et al. (2015) observed that the light penetration depth in the Den Osse Basin never exceeded 15 m in 2012, and therefore photosynthesis by kleptoplasts (Bernhard and Alve, 1996) appears unlikely for both our aphotic stations (34 and 23 m

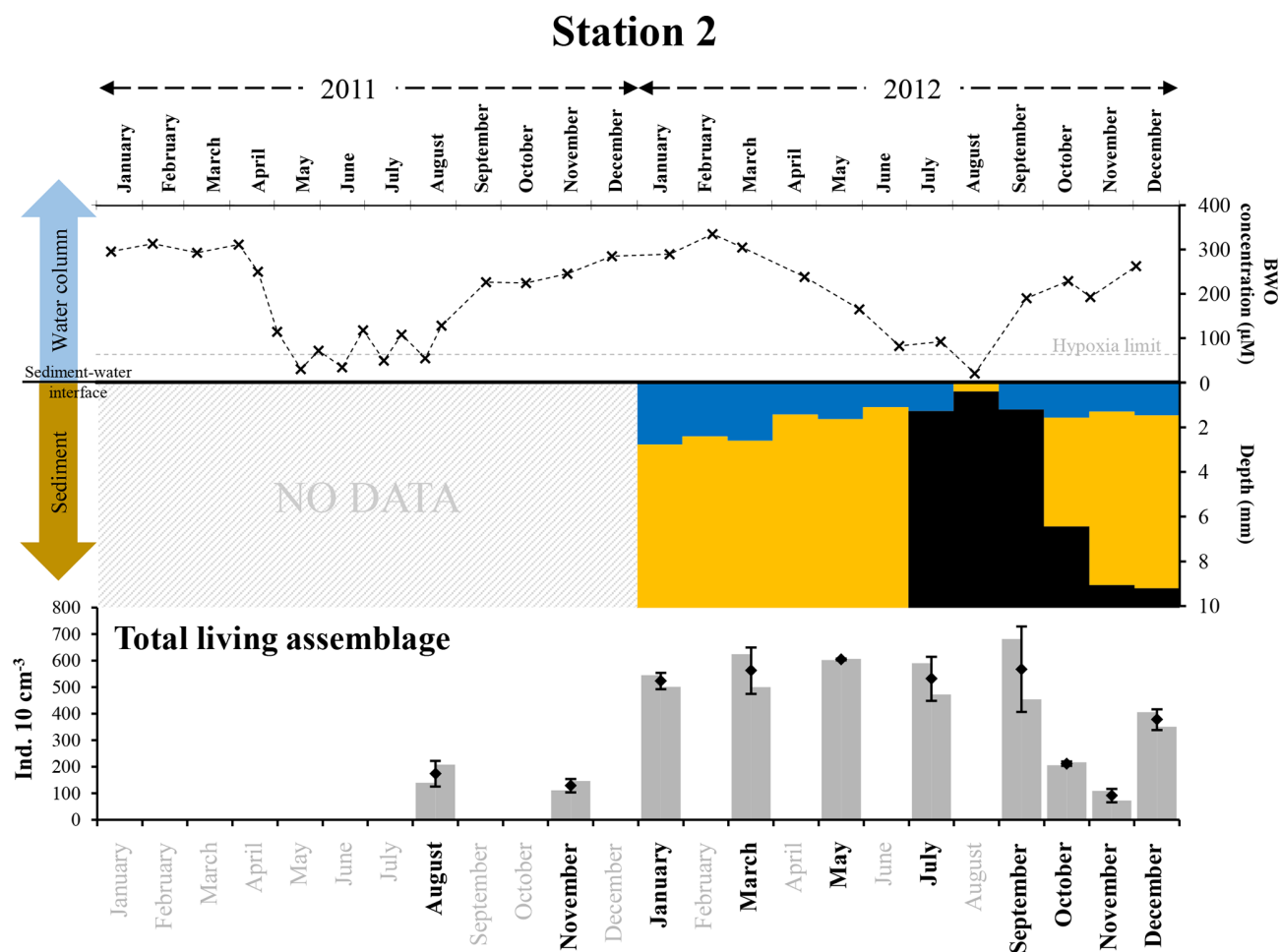


Figure 9. The top panel represents bottom-water oxygen concentrations ($\mu\text{mol L}^{-1}$) in 2011 and 2012 at station 2, from Donders et al. (2012) and Seitaj et al. (2017). The grey horizontal dotted line indicates the hypoxia limit ($63 \mu\text{mol L}^{-1}$). The middle panel represents the depth (mm) distribution of the oxic (blue), suboxic (orange, absence of oxygen and sulfides) and sulfidic (black) zones within the sediment in 2012. The bottom panel shows the total living foraminiferal abundances for both replicates (grey bars), mean abundances (diamonds) and standard deviations (black error bars) calculated for the two replicates, for all investigated months (in bold) in 2011 and 2012.

depth). However, other foraminifera from aphotic and anoxic environments such as deep fjords are kleptoplastic and use these kleptoplasts for a yet unknown purpose (Jauffrais et al., 2019).

Rather surprisingly, the drop in foraminiferal densities at station 2 in October–November, which we interpreted as a delayed response to sulfidic conditions, is less strong for *Ammonia* sp. T6 than for the two *Elphidium* species, suggesting that this species is less affected. However, this does not agree with our previous suggestion that the two *Elphidium* species would be more tolerant to anoxic and sulfidic conditions. As already proposed by LeKieffre et al. (2017), *Ammonia* seems to be able to deal with anoxia (up to 28 d, but with no sulfide) by reducing its metabolic activity, but this ability was never shown for *Elphidium* species. If *E. selseyense* and *E. magellanicum* are indeed unable to resist anoxia by reducing their metabolism or by entering a dormancy state, this could ex-

plain their stronger decrease in density at station 2 compared to *Ammonia* sp. T6. Nevertheless, further studies about the ability and mechanisms of the two *Elphidium* species to resist anoxic–sulfidic conditions are necessary.

Another remarkable observation is that *Ammonia* sp. T6 (and *T. inflata*) shows maximum densities in January–March, contrasting with the two *Elphidium* species, which have their density maxima later in the year (May–September). This temporal offset could possibly be explained by a difference in preferential food source, with food particles available in winter (January–March) being more suitable for *Ammonia* sp. T6 (and *T. inflata*) and food particles available later in the year, resulting from phytoplankton blooms, being more favourable for *E. selseyense* and *E. magellanicum*.

In our study, for *E. selseyense* (and *E. magellanicum*), the continuous presence of a high proportion of small-sized specimens and progressively increasing densities between

January and September 2012 strongly suggest ongoing and continuous reproduction (Fig. S3a). Continuous reproduction during the year has been described earlier for different foraminiferal genera, such as *Elphidium*, *Ammonia*, *Haynesina*, *Nonion* and *Trochammina* (e.g. Jones and Ross, 1979; Murray, 1983, 1992; Cearreta, 1988; Basson and Murray, 1995; Gustafsson and Nordberg, 1999; Murray and Alve, 2000). Conversely, for *Ammonia* sp. T6, a decrease in densities coupled with a rapid increase in overall test size between March and May 2012 (small sized specimens remain present but in smaller proportions) could be indicative of a period of reduced recruitment (Fig. S3b).

In fact, foraminifera exhibit a large range of feeding strategies, with several species showing selective feeding with specific food particles (Muller, 1975; Suhr et al., 2003; Chronopoulou et al., 2019). Hagens et al. (2015) reported that in Lake Grevelingen the phytoplankton composition was different between April–May and July 2012. In April–May, the phytoplankton bloom was mainly composed of the haptophyte *Phaeocystis globosa* (Scherffel, 1899), whereas it was dominated by the dinoflagellate *Prorocentrum micans* (Ehrenberg, 1834) in July. *Elphidium* was reported to be able to feed on various food sources (e.g. diatoms, dinoflagellates, green algae; Correia and Lee, 2002; Pillet et al., 2011). However, diatoms are a major food source for kleptoplastic species (Bernhard and Bowser, 1999), such as *E. selseyense* (Jauffrais et al., 2018; Chronopoulou et al., 2019). *Ammonia* spp. seem able to feed on very diverse food sources including microalgae, diatoms, bacteria or even metazoans (Lee et al., 1969; Moodley et al., 2000; Dupuy et al., 2010; Jauffrais et al., 2016; Chronopoulou et al., 2019). Recently, Chronopoulou et al. (2019) showed different feeding preferences for *Ammonia* sp. T6 and *E. selseyense* in intertidal environments in the Dutch Wadden Sea. Although diatoms are ingested by both species (but much more by *E. selseyense*), dinoflagellates were consumed by *E. selseyense* but not by *Ammonia* sp. T6. The latter species is also capable of feeding on metazoans by active predation (Dupuy et al., 2010).

These observations suggest that at station 2 the different seasonal density patterns of *Ammonia* sp. T6 and the two *Elphidium* species are not the consequence of a large difference in tolerance of anoxia–sulfides, but rather a different adjustment to the seasonal cycle of food availability. At station 1, the very low densities of *Ammonia* sp. T6 could possibly be explained by a recolonisation starting in January, when food conditions were favourable for this taxon (as testified by the strong density increase in January 2012 at station 2). However, once a more abundant pioneer population had developed (in March–May), food conditions may have been no longer favourable for *Ammonia* sp. T6, explaining why its density did not show a further increase. Conversely, the food conditions may have become optimal for the two *Elphidium* species, explaining their strong density increase between March and May 2012. If true, this would mean that the lower densities of *Ammonia* sp. T6 would not be due

to a lower resistance to anoxia and free sulfides, but rather due to an unfavourable seasonal succession of food availability. Previous studies already suggested that hypoxic–anoxic conditions coupled with increased food input from autumnal phytoplankton blooms (composed of diatoms and dinoflagellates) would favour the development of *E. magellanicum* (Gustafsson and Nordberg, 1999). The fact that also at station 2 this species was mainly observed between March and September 2012 corroborates our conclusion of its dependence on a specific food regime.

Finally, encrusted forms of *E. magellanicum* were observed at both stations from May until the end of the year but were absent in the samples of March 2012. In view of the fact that the crusts consist mainly of organic matter, the encrusted individuals appear to be specimens with preserved feeding cysts. The precise functions of cysts observed around foraminifera are not clear and include feeding, reproduction, chamber formation, protection or resting (Cedhagen, 1996; Heinz et al., 2005). Concerning the cysts of *E. magellanicum* described here, very similar observations have been made for *Elphidium incertum* at different locations (Norwegian Greenland Sea and Baltic Sea in Linke and Lutze, 1993; Koljö Fjord in Gustafsson and Nordberg, 1999; Kiel Bight in Polovodova et al., 2009). If we assume that encrusted specimens indeed present the remains of feeding cysts, the observation of abundant encrusted specimens corroborates our conclusion that the surface water phytoplankton bloom in May 2012 (i.e. probably mainly *Phaeocystis globosa*) provided a food source particularly well suited to the nutritional preferences of this species.

5 Conclusions

In this study we examined the foraminiferal community response to different durations of seasonal anoxia coupled with the presence of sulfide in the uppermost layer of sediment at two stations in Lake Grevelingen. In both stations investigated, foraminiferal communities are highly impacted by the combination of anoxia and H₂S in their habitat. The foraminiferal response varied depending on the duration of adverse conditions and led to a near extinction at station 1, where anoxic and sulfidic conditions were present for 1 to 2 months, compared to a drop in standing stocks at station 2, where these conditions lasted for 1 month or less. At both sites, foraminiferal communities showed a 2-month delay in the response to anoxic and sulfidic conditions, suggesting that the presence of H₂S inhibited reproduction, whereas mortality was not necessarily increased. The duration of the subsequent recovery depended on whether the foraminiferal community was almost extinct (station 1) or remained present with reduced numbers (station 2). In the former case, 6 months were needed for faunal recovery, whereas in the latter case, it took only 2 months. We hypothesise that the dominance of *E. selseyense* and *E. magellanicum* at sta-

tion 1 is not due to a lower tolerance of *Ammonia* sp. T6 towards anoxic and sulfidic conditions, but is rather the consequence of a different adjustment between the two *Elphidium* species and *Ammonia* sp. T6 with respect to the seasonal cycle of food availability.

Data availability. Raw data are available in the Supplement.

Supplement. The supplement related to this article is available online at: <https://doi.org/10.5194/bg-17-1415-2020-supplement>.

Author contributions. BR, DL and JR produced the foraminiferal data. DS, AM and CPS provided and interpreted geochemical data. MS provided, verified and integrated all available genetic information concerning the foraminiferal taxa of Lake Grevelingen. FJRM and CPS coordinated a much larger research project concerning Lake Grevelingen, of which this foraminiferal study is a part. They were also responsible for the foraminiferal sampling and provided environmental data. FJJ designed the foraminiferal study and directed the postdoctoral research of BR and, together with AM and MS, the PhD thesis of JR. All authors contributed actively to the several successive versions of the manuscript.

Competing interests. The authors declare that they have no conflict of interest.

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