Scale dependence of temporal biodiversity change in modern and fossil marine plankton

Aleksandra M. Lewandowska1,2 | Lukas Jonkers3 | Holger Auel4 | Jan A. Freund1 | Wilhelm Hagen4 | Michal Kucera3 | Helmut Hillebrand1,5

1Institute for Chemistry and Biology of the Marine Environment, Carl von Ossietzky University of Oldenburg, Oldenburg, Germany
2Tvärminne Zoological Station, University of Helsinki, Hanko, Finland
3MARUM – Center for Marine Environmental Sciences, University of Bremen, Bremen, Germany
4BreMarE – Bremen Marine Ecology, Marine Zoology, University of Bremen, Bremen, Germany
5Helmholtz-Institute for Functional Marine Biodiversity at the University of Oldenburg (HIFMB), Oldenburg, Germany

Correspondence
Aleksandra M. Lewandowska, Tvärminne Zoological Station, University of Helsinki, J.A. Palmenin tie 260, 10900 Hanko, Finland
Email: aleksandra.lewandowska@helsinki.fi

Funding information
Lower Saxony Ministry of Science and Culture (MWK), Grant/Award Number: MarBAS - Marine Biodiversity Across Scales

Editor: Adam Tomasovych

Abstract

Aim: Biodiversity dynamics comprise evolutionary and ecological changes on multiple temporal scales from millions of years to decades, but they are often interpreted within a single time frame. Planktonic foraminifera communities offer a unique opportunity for analysing the dynamics of marine biodiversity over different temporal scales. Our study aims to provide a baseline for assessments of biodiversity patterns over multiple time-scales, which is urgently needed to interpret biodiversity responses to increasing anthropogenic pressure.

Location: Global (26 sites).

Time period: Five time-scales: multi-million-year (0–7 Myr), million-year (0–0.5 Myr), multi-millennial (0–15 thousand years), millennial (0–1,100 years) and decadal (0–32 years).

Major taxa studied: Planktonic foraminifera.

Methods: We analysed community composition of planktonic foraminifera at five time-scales, combining measures of standing diversity (richness and effective number of species, ENS) with measures of temporal community turnover (presence-absence-based, dominance-based). Observed biodiversity patterns were compared with the outcome of a neutral model to separate the effects of sampling resolution (the highest in the shortest time series) from biological responses.

Results: Richness and ENS decreased from multi-million-year to millennial time-scales, but higher standing diversity was observed on the decadal scale. As predicted by the neutral model, turnover in species identity and dominance was strongest at the multi-million-year time-scale and decreased towards the millennial scale. However, contrary to the model predictions, modern time series show rapid decadal variation in the dominance structure of foraminifera communities, which is of comparable magnitude as over much longer time periods. Community turnover was significantly correlated with global temperature change, but not on the shortest time-scale.

Main conclusions: Biodiversity patterns can be to some degree predicted from the scaling effects related to different durations of time series, but changes in the dominance structure observed over the last few decades reach higher magnitude,
probably forced by anthropogenic effects, than those observed over much longer durations.

**KEYWORDS**

biodiversity, community turnover, global change, planktonic foraminifera

# 1 | INTRODUCTION

Biodiversity change has been prevalent through the geological past (Fenton, Pearson, Dunkley Jones, Farnsworth, et al., 2016; Wade & Pearson, 2008; Yasuhara, Tittensor, Hillebrand, & Worm, 2017) and over recent decades (Dornelas et al., 2014; Elahi et al., 2015; Hillebrand, Blasius, et al., 2018). Even though ongoing biodiversity change appears consistent with forcing by anthropogenic global change (Hillebrand, Brey, et al., 2018; Newbold et al., 2015; Poloczanska et al., 2013), it remains unclear if the observed species turnover exceeds the natural range of variability. This is because most observational time series are short and lack historical baselines (Cardinale, Gonzalez, Allington, & Loreau, 2018; Gonzalez et al., 2016; Yasuhara, Hunt, Breitburg, Tsujimoto, & Katsuki, 2012). Clearly, a proper baseline is needed to make accurate predictions on how climate change and anthropogenic activities will influence biodiversity in the future (Cardinale et al., 2018; Jackson, 2008). Defining a clear baseline for biodiversity before industrialization is difficult, because most monitoring programs post-date the onset of the industrial revolution (Dornelas et al., 2014; Elahi et al., 2015; Lotze et al., 2006). As a result, we lack information on the variation in species numbers and community composition (as one aspect of biodiversity) before human impact. However, for some taxonomic groups, the fossil record provides the necessary temporal context encompassing the range of ecological variation on time-scales from decades to millions of years and, therefore, there is increasing interest in using palaeontological records to answer ecological questions (Fritz et al., 2013; Harnik et al., 2012; Nogués-Bravo et al., 2018; Yasuhara et al., 2017). Recent findings by Jonkers, Hillebrand, and Kucera (2019) confirmed that composition of modern foraminifera assemblages differ from those in pre-industrial times as a result of temperature change, highlighting the importance of historical baselines for the assessment of ecological communities.

Another critical aspect of studies reporting anthropogenic impact on biodiversity is the focus on local species richness (Chase & Knight, 2013; Hillebrand, Blasius, et al., 2018). Recent analyses reveal that some communities are undergoing major compositional change without change in species number (Hillebrand, Blasius, et al., 2018; Magurran et al., 2018). This is because colonization by new species sometimes equals or exceeds rates of local extinctions (Sax & Gaines, 2003), so that local richness fluctuates around a steady state, despite changes in community composition and species abundance. Reordering of species dominance patterns can occur along with changes in species identity (species appearance and disappearance) or can be independent of identity change. If there is a change in dominance, but no change in species presence/absence, biodiversity change is driven by reordering the relative abundances of species already present in the local community (Hillebrand, Blasius, et al., 2018). Therefore, an analysis of baseline biodiversity change that is relevant for biodiversity conservation has to consider both richness (presence/absence) and dominance (relative abundance) (Barnosky et al., 2017; Cardinale et al., 2018). Accordingly, we here focus on two metrics of biodiversity change: (a) change in species presence/absence (species turnover measured by Jaccard’s dissimilarity), and (b) reordering of species dominance patterns (Wishart’s dissimilarity) as described by Hillebrand, Blasius, et al. (2018). For completeness, we also discuss changes in species richness (as it is the most common measure of biodiversity), and the effective number of species (ENS), which has been suggested to be a more robust diversity metric than richness regarding different sampling schemes (Chase & Knight, 2013). By comparing the identity-based metrics (richness, Jaccard) with dominance-based metrics (ENS, Wishart), we provide a comprehensive assessment of the magnitude of biodiversity change across timescales.

We use planktonic foraminifera as a model system to study scale dependence of temporal biodiversity patterns in the marine realm. Planktonic foraminifera are open-ocean protists that have the most complete taxonomic record of any clade, which allows for direct comparisons between recent and geologically distant communities (Fenton, Pearson, Dunkley Jones, Farnsworth, et al., 2016). They are widely sampled both in the recent past and throughout deep-time, so that rich datasets on foraminiferal assemblages are available across large spatial and temporal scales (Fenton, Pearson, Dunkley Jones, & Purvis, 2016; Yasuhara et al., 2017). Planktonic foraminifera are sensitive to environmental change, such as temperature (Ezard, Quental, & Benton, 2016; Jonkers et al., 2019), and are therefore widely used for palaeoclimate reconstructions (Kucera, 2007). In this study, we analysed data on community composition of planktonic foraminifera over the past 7 million years to track dynamics of biodiversity change in the ocean. Rather than defining the drivers of biodiversity change, of which there are many, we assessed the magnitude of biodiversity fluctuations across five distinct temporal scales from decades to millions of years. Benefitting from access to modern monitoring time series as well as to fossil data, we analysed community turnover in planktonic foraminifera at these time-scales, specifically asking how the magnitude of change detected in modern time series of species turnover compares with observations in the fossil record. In particular, we tested the following hypotheses:

Hypothesis 1: In the multi-million-year time series, an increasing number of speciation and extinction events increases temporal turnover based on presence-absence, and might increase the overall
number of species observed at locations where the speciation rates exceed the extinction rates. For dominance-based metrics (ENS, Wishart’s dissimilarity), we expect similar patterns but even higher turnover as – in addition to speciation and extinction – changes in relative abundance will accumulate over time.

Hypothesis 2: In the more recent records (million-year to decadal scales), where speciation and extinction do not affect temporal biodiversity patterns (Yasuhara, Hunt, Dowsett, Robinson, & Stoll, 2012; Yasuhara et al., 2017), the number of species will increase and turnover will decrease in longer time series that are intrinsically characterized by higher averaging than shorter time series.

In addition, we compared the observed changes in biodiversity to changes in temperature over the same time, testing the Hypothesis 3 that global temperature cycles are associated with higher rates of turnover during warming events. We expect that these effects are most pronounced during ice ages and after the last glaciation (million-year and multi-millennial scales) characterized by the strongest temperature fluctuations.

2 | METHODS

2.1 | Data sources

We compiled data on foraminifera community composition (species’ relative abundances) from 20 sediment cores and 6 sediment traps with a minimum length of 5 years distributed over the world’s oceans (Figure 1). Only cores that contained the full taxonomic record were included in the analysis. Cores were selected to explore different scales of temporal variability in foraminifera community composition, thus datasets that had too low sampling resolution to cover this variability (e.g. too short records) were omitted. The datasets (Supporting Information Appendix S1) were grouped into five categories depending on how far back in time they extend: multi-million-year scale (0 to 7 Myr, eight cores), million-year scale (0 to 0.5 Myr, four cores), multi-millennial scale (0 to 15 thousand years, three cores), millennial scale (0 to 1,100 years, five cores) and decadal scale (0 to 32 years, six sediment trap records). The longest datasets were characterized by the highest sample integration (highest time averaging, Supporting Information Appendix S1: Table S1). All data from sediment traps were analysed within the decadal scale category. From the sediment traps we extracted yearly averages of the foraminifera fluxes to eliminate seasonal dynamics (Jonkers & Kučera, 2015). The year 2016 was set as ‘present day’ for all datasets (cores and traps) to facilitate a direct comparison of the results.

We also compiled estimates of global temperature for each time frame (Supporting Information Appendix S1), except for the 0–7 Myr period for which no global compilation is available. Instead, we used a tropical (almost exclusively Pacific) temperature stack (Herbert et al., 2016), which is appropriate as all data that span this period are from this basin.

FIGURE 1 Geographical distribution of time series analysed in this study. Names of the sites and their description are listed in the Supporting Information Appendix S1: Table S1 [Colour figure can be viewed at wileyonlinelibrary.com]
2.2 | Scaling in a neutral community model

To create reference time series for different scales, we designed a stochastic process simulating a neutral community model (for more detailed description, please see Supporting Information Appendix S2). In this model stochastic speciation events are introduced through a constant speciation rate that is balanced by the extinction rate in order to generate a species richness that fluctuates around the level of 20. When a speciation event occurs, a new species is released at the detection limit (zero). When species abundance declines crossing the detection limit, it is recorded with the value zero, but left in the simulation with a chance of reoccurrence. Eventually, when it hits the extinction threshold (at \(-1\)) it goes extinct with no possibility of returning. This model assumes that all species follow the same population dynamics and there are no community interactions between species (neutral community model). Time series of the community evolution are simulated based on equidistant sampling at a sampling rate \(f_s\), where \(f_s = 1/\Delta t\). Over continuous time the stochastic population dynamics of each species is red noise generated by an Ornstein-Uhlenbeck (OU) process with a mean reversion value \(\mu\) and a correlation time of \(\tau_c\) in combination with the absorbing extinction threshold. In discrete time the OU process is connected with a vector autoregressive process of order one:

\[ x_n = \mu + A(x_{n-1} - \mu) + \epsilon_n \]

where \(x_n\) is a vector that comprises concentrations of all \(N\) species at given time \(t = n\Delta t\), \(\mu\) is the mean reversion value, which is set to 1 for all species (neutral community model), \(A\) is a constant \(N \times N\) matrix with non-zero entries only along its diagonal according to the assumption of non-interacting species and diagonal elements are all identical (neutral community model), \(\epsilon_n\) is a vector of \(N\) statistically independent Gaussian white noise fluctuations (with zero mean and appropriately scaled variance, more details in Supporting Information Appendix S2).

We used this model to simulate time series and infer frequency distributions of diversity metrics at different time-scales (time-series lengths 0.1, 1, 10, 1,000 and 10,000 kyr, corresponding to decadal, millennial, multi-millennial, million-year and multi-million-year time-scale categories, respectively). We used six, five, three, four and eight simulation runs for decadal, millennial, multi-millennial, million-year and multi-million-year time-scale categories, respectively. In contrast to the empirical data, time resolution was adapted to render 100 equidistant time points per simulation run with a sampling interval of 0.001, 0.01, 0.1, 10 and 100 kyr, respectively. This means that going from the shortest to the longest scale the simulated time series were uniformly downsampled by a factor of \(10^5\) and time-series length is positively correlated with sampling interval. The initial foraminifera community contained 20 species, except for the decadal scale records, for which we started with 6, 6, 11, 11, 12 and 13 species in the related six runs.

2.3 | Statistical analysis

We calculated species richness as the number of morphospecies (fossil species can only be defined using the morphological species concept) identified in each sample. The ENS was calculated as described by Chase and Knight (2013) (see also Supporting Information Appendix S2). The advantage of using ENS over richness is that ENS is less sensitive to sampling effort and does not overestimate the impact of rare species on the overall diversity.

Jaccard’s dissimilarity and Wishart’s dissimilarity were calculated as described by Hillebrand, Blasius, et al. (2018). Jaccard’s dissimilarity is a presence–absence-based species exchange ratio between two samples and represents changes in community composition in terms of species identities, whereas Wishart’s dissimilarity is a measure of dominance change (turnover in species proportional abundances). The latter is closely related to the Simpson index and thus related to robust measures of diversity, such as the ENS (Chase & Knight, 2013). For each core and trap series we calculated both dissimilarity metrics comparing foraminifera assemblages between each unique pair of samples (the diagonal of the dissimilarity matrix was always omitted to exclude zero dissimilarities between each sample and itself). Then, we calculated a median of pairwise dissimilarities between a sample of a given age and all other samples in the core or the sediment trap (Figure 2). The advantage of this procedure is that the temporal trend does not depend on a single reference community at the beginning or at the end of the time series. Finally, for each site we calculated a trend using generalized additive models (GAMs) with cubic regression splines. The error covariance was assumed to follow a time-dependent autoregressive process to account for temporal correlation.

General trends within each five categories were derived by applying GAMs to the fitted values for individual series and site ID was included as a random effect (random intercept) to account for differences between sampling sites (different location and sample integration). We used the GAM approach, because it enables us to account for statistical irregularities in the data, such as non-uniform sampling frequency and interpolating over missing observations. GAMs also allow the response to be estimated as a linear or nonlinear function of the predictors, giving a better chance of detecting non-monotonic patterns, for example, glacial-interglacial fluctuations. Model assumptions, including normality, independence and homogeneity of variance, were verified by visual analysis of residuals. All analyses were performed in R (R Development Core Team, 2017, version 3.3.3) using package ‘mgcv’ for GAMs (Wood, 2017). More details on statistical procedures are included in Supporting Information Appendix S2.

To further explore the effects of temporal scaling (Tomašových & Kidwell, 2010) we made a simple comparison of the mean diversity values (measured using our four indices) with time-series length, sampling frequency (time separation between the samples) and sample integration (inverse of the sedimentation rate measured as a
first derivative of the depth-age model for each analysed sediment core).

To address whether observed biodiversity patterns were associated with temperature change, we performed cross-correlation analyses for each time category. Results of these analyses are presented in detail in Supporting Information Appendix S2.

3 | RESULTS

3.1 | Frequency distribution of diversity metrics

The frequency distribution of diversity measures shows substantial changes in standing diversity, identity and dominance change over the different time-scales (Figure 3). All diversity measures decrease with decreasing duration (and increasing resolution) of time series, with the exception of the shortest decadal records. Richness tends to decrease over pre-modern time categories (from multi-million-year to millennial time-scales), but also diverges from a rather normally distributed sample at scales of millions of years to a multimodal distribution at millennial time-scales (Figure 3a). For decadal samples, the richness is higher again.

In terms of species turnover recorded as presence–absence (Jaccard), decadal time series show less turnover than time series at millennial to multi-million-year time-scales (Figure 3b), which is consistent with the outcome of our neutral scaling model (Figure 4b) revealing the effects of time-series resolution (higher turnover values in longer cores). The duration of the decadal time series (5–32 years) is too short to record consistent extirpation or invasion, thus suggesting lower turnover. A distinctly higher turnover is seen on time-scales up to $10^6$ years (Figures 3b and 4b), where we also see the strongest response to climatic forcing (Supporting Information Appendix S2: Figure S5).

These conclusions change when taking dominance into account. The effective number of species declines from multi-million-year to millennial time-scales (Figure 3c), which is not a time averaging effect as confirmed by the model simulation (Figure 4c). This indicates that not only richness but also evenness declines (dominance increases) along these time-scales. In terms of compositional turnover (Figure 3d), we observe a similar trend towards a leftward skew of the frequency distribution from multi-million-year to millennial time series. Thus, the combination of species identity exchange and reordering of dominance of persistent species shows a large magnitude and range of changes across time-scales. The range is still large for multi-millennial datasets, whereas millennial time series show much less turnover, potentially due to the absence of large climatic forcing over the last millennium compared to the transition out of the last ice age that is covered in the multi-millennial time series. Interestingly, decadal time series show a similar range of dominance-based turnover as in glacial/interglacial time series (see both million-year and multi-millennial datasets) that cannot be attributed to the effects of temporal scaling (Figure 4d). Thus, at the smallest temporal scales we see dominance shifts that are not observed at the scales of 100–1,000 years and rarely visible in longer, but lower resolution, time series.
3.2 Dominance-based biodiversity change over time

Based on the observed changes in the mode and range of dominance-based diversity metrics across time-scales, we focus on temporal trends of ENS and Wishart’s dissimilarity here, but show similar graphs for richness, and Jaccard turnover in Supporting Information Appendix S3. Analysis of foraminifera ENS over the past 7 million years shows nonlinear pattern in single cores, but highlights a consistent diversity maximum around 3 million years ago (Figure 5a), close to the time of major intensification of the Northern Hemisphere glaciation (Figure 5c) (Raymo, 1994), corresponding to the aftermath of the last diversification of planktonic foraminifera after which only extinctions occurred (Fritz et al., 2013). Dominance-based turnover is generally high over these time-scales (Figure 5b, see above), but shows a decline towards the time of the Northern Hemisphere glaciation 3 Ma (Figure 5b). After the glaciation started (Figure 5c), ENS decreased and Wishart’s dissimilarity did not change, but remained at a high overall level.

Over glacial–interglacial cycles (million-year time-scale), ENS and Wishart’s dissimilarity show cycles with a frequency corresponding to the glacial cycles and peaks during interglacial periods (Figure 5d–f and Supporting Information Appendix S2: Figure S5). Details differ between cores, but maximum ENS and turnover in species composition is often associated with warming periods. The overall magnitude of turnover is much smaller than on the multi-million-year time-scale (cf. Figure 5b and e, respectively).

In the multi-millennial datasets, temporal turnover dynamics are related to climate instability at the end of the last glacial period c. 12,000 years ago (Figure 5g–i). However, ENS peaked significantly later (c. 10 kyr ago) than the turnover metric (c. 13.5 kyr ago), indicating that reordering of community composition can take place without gaining or losing effective species. In fact, turnover (Figure 5h) is high during the warming phase, when it is in the same range as at...
the multi-million-year time-scale, and decreases when temperature is constant.

In contrast to the significant changes in ENS and Wishart’s dissimilarity over longer time-scales, both remained constant over the last hundreds to thousand years (Figure 5j–l). Although small sample size should be acknowledged, differences between sites are obvious, as both metrics are higher and more variable at lower latitudes (sites C90, SO90 and SBB2 compared to MD99_3 and MD99_5). However, the overall trend of ENS and Wishart’s dissimilarity remains neutral, likely reflecting the stable climate during this period (Figure 5l and Supporting Information Appendix S2: Figure S2) and no effect of extinction and speciation on this temporal scale.

In the decadal records, ENS increased with time and the shortest trap series showed the highest ENS (Figure 5m). Notably, however, Wishart’s dissimilarity remained very high (comparable to glacial-interglacial turnover; Figure 5n).

### 3.3 Effects of temporal resolution

The duration of the time series had no effect on the mean ENS in the empirical data (Figure 6b). Highest species richness and turnover values (Jaccard’s and Wishart’s dissimilarity) were observed in the longest records (multi-million-year time series), because of a high number of speciation and extinction events that could be captured on this time-scale. As the longest time series had also significantly longer temporal separation between sampling events, the same pattern was observed for the relationships between biodiversity measures and sampling frequency (Figure 6e–h), that is, time series with extremely low sampling frequency were characterized by highest richness and turnover values.

Fossil assemblages are averaged over a certain amount of time (sample coarsening), which can influence estimates of species turnover (Tomašových & Kidwell, 2010), but such sample integration does not affect biodiversity patterns in our study (Figure 6i–l). This
can be probably explained by counteracting effects of duration and sample integration time (Tomašových & Kidwell, 2010) with the duration effects overriding the effects of sample resolution.

4 | DISCUSSION

Our analysis provides a first evaluation of marine biodiversity patterns across temporal scales from millions of years to decades. It indicates that trends in community turnover are key to understanding these patterns. The magnitude of temporal turnover observed in the decadal time series is smaller in terms of richness, but compositionally larger than what is seen on the millennial time-scale and the range of the compositional turnover is consistent with the climatically driven pattern of glacial and post-glacial time series (Figure 5 and Supporting Information Appendix S2: Figure S5). When considering the records from millennial to multi-million-year time-scales, we find a significant correlation between climatic changes and the turnover in composition and standing diversity of foraminifera in presence–absence-based estimates (richness, Jaccard), as well as in dominance-affected measures (ENS, Wishart) supporting Hypothesis 3 that global warming events are associated with higher rates of turnover. This is visible in Figure 5 and Supporting Information Figure S1 (Appendix S3), and supported by the results of cross-correlation analyses between global temperature records and observed biodiversity patterns (Supporting Information Appendix S2: Figure S5). Here, we focus our discussion on ENS and Wishart’s dissimilarity, as these are more likely to be informative on the short time-scales we are interested in for predicting future diversity responses to climate change.

On the geological time-scale of millions of years, ENS peaked at the onset of the Northern Hemisphere glaciation that occurred 3.5 million years ago, and turnover is decreasing in a cooling world. These changes are smaller in magnitude, but consistent at the million-year time-scale: turnover and ENS are higher in warming periods (Figure 5 and Supporting Information Appendix S2: Figure S5). The magnitude of change decreases when considering multi-millennial time frames and is very small to absent at millennial time-scales. On long time-scales (millions of years), changes in community turnover are in part driven by slowly occurring species extinctions (global or geographically selective) and speciation (evolution), whereas on shorter time-scales climate change plays the dominant role and the turnover dynamics reflect geographical redistribution of species, rather than extinction (Jackson, 2008; Yasuhara, Hunt, Dowsett, et al., 2012). Geographical differences between different sites are most pronounced in millennial records that are characterized by higher values of ENS and Wishart’s dissimilarity at low latitudes (Figure 5, sites C90, SBB2, SO90).
compared to high latitudes (Figure 5, sites MD99_3 and MD99_5), which is in line with the spatial distribution of fossil foraminifera diversity (high tropical and low polar biodiversity, Yasuhara, Hunt, Dowsett, et al., 2012). Climate-driven changes in species community structure have been described for many organism types in single locations or regions (small mammals: McGill, Hadly, & Maurer, 2005; dinoflagellates: de Vernal et al., 2005; corals: Pandolfi & Jackson, 2006; ostracodes: Yasuhara, Cronin, Hunt, & Hodell, 2009; benthic foraminifera: Reymond, Bode, Renema, & Pandolfi, 2011; planktonic foraminifera: Flower & Kennett, 1995; diatoms: Ren, Gersonde, Esper, & Sancetta, 2014; molluscs: Tomášových, Dominici, Zuschin, & Merle, 2014) and recently also for planktonic foraminifera before and after industrialization (Jonkers et al., 2019). However, most studies take a reverse approach and use changes in species assemblages to reconstruct past climate, rather than focusing on patterns of species turnover associated with climate change (Yasuhara, Hunt, Breitburg, et al., 2012). Our study goes beyond earlier work by analysing dominance-driven biodiversity change worldwide on different time-scales.

Consistent with the lack of substantial climatic change over the last millennium, ENS remained stable and turnover was small (Figure 5). This observation contrasts with the analysis of decadal records. Here, we see strong fluctuations of the dominance structure (Wishart’s dissimilarity) independent of changes in ENS, with the magnitude of change comparable to turnover observed over millions of years. This is in line with recent literature showing that community composition is rapidly changing, whereas the number of species remains stable over time (Dornelas et al., 2014; Hillebrand, Blasius, et al., 2018; Magurran et al., 2018). This high modern turnover of assemblages may reflect anthropogenic forcing and the comparison with our neutral scaling model indicates that stochastic processes cannot explain observed variability in foraminifera biodiversity on decadal time-scales (Figure 4). Thus, the lower turnover on the millennial time-scale is not a result of higher temporal averaging, but reflects ecological processes (Supporting Information Appendix S2: Figure S5). Since the decadal time series were annually averaged, seasonal variation in modern time series does not explain these differences as they are not reflected in our turnover estimates. These data thus suggest that the high turnover in modern assemblages is a response to fast environmental changes, such as increasing mean and variance in temperature.

Our analysis of foraminifera biodiversity patterns may be influenced by the effects of temporal resolution (time-series length,
sampling frequency and sample integration) on species turnover (Tomasových & Kidwell, 2010). In line with Hypothesis 1, we observed the highest number of species and highest values of turnover in the multi-million-year time series, but these results are partly due to scaling (Figure 6). However, scaling effects do not explain either the patterns in ENS (Figure 6) or the high variability in Wishart’s dissimilarity in the decadal records (Figure 3). Thus, Hypothesis 2 (decreasing turnover in modern times as a result of time averaging) can be supported for the presence–absence-based turnover, but not for the dominance-based turnover.

In conclusion, our analysis shows how biodiversity patterns are shaped at different time-scales, which is an important step towards understanding the processes driving diversity dynamics across scales. On the multi-million-year time-scale, slow processes, such as speciation and global extinction, can be addressed. On the shorter time-scales (thousands to hundreds of years), climatic forcing most likely prevails, whereas biodiversity patterns observed on the decadal scale are possibly a result of fast occurring processes, such as species dispersal. Temporal scaling effects (Tomasových & Kidwell, 2010) can explain higher richness and turnover values on the multi-million-year time-scale (Figure 6), but not the high variation in the dominance structure in modern times. We reinforce the importance of historical baselines of biodiversity change in interpretations of current dynamics and reveal that restructuring of communities occurs at all times, but changes in the modern dominance structure may reach higher magnitudes than expected from Earth’s history. Given the rapid change in climate, the critical question in ecology is whether species will be able to adapt or move, and thus communities to persist. Analysis of species richness alone is inconclusive and does not allow predictions on diversity dynamics (Cardinale et al., 2018), whereas turnover estimates prove to be more sensitive and consistent in their response to past climate change, with higher turnover during warming periods.

ACKNOWLEDGMENTS
We thank Ulrike Feudel and Marina C. Rillo (Carl von Ossietzky University of Oldenburg) for helpful discussions and comments on the earlier version of the manuscript. Julian Merder is acknowledged for his help with model simulations. This work was supported by the MarBAS (Marine Biodiversity Across Scales) project funded by the Lower Saxony Ministry of Science and Culture (MWK).

DATA AVAILABILITY STATEMENT
Data are available in the Dryad data repository (https://doi.org/10.5061/dryad.2z34tpmhf) and the sources are listed in the Appendix.

ORCID
Aleksandra M. Lewandowska https://orcid.org/0000-0003-2925-4509
Lukas Jonkers https://orcid.org/0000-0002-0253-2639
Jan A. Freund https://orcid.org/0000-0002-2076-3131
Wilhelm Hagen https://orcid.org/0000-0002-7462-9931
Michal Kucera https://orcid.org/0000-0002-7817-9018
Helmut Hillebrand https://orcid.org/0000-0001-7449-1613

REFERENCES
Barnosky, A. D., Hadly, E. A., Gonzalez, P., Head, J., Polly, P. D., Lawing, A. M., … Zhang, Z. (2017). Merging paleobiology with conservation biology to guide the future of terrestrial ecosystems. Science, 355, eaa4787. https://doi.org/10.1126/science.aah4787
Cardinale, B. J., Gonzalez, A., Allington, G. R. H., & Loreau, M. (2018). Is local biodiversity declining or not? A summary of the debate over analysis of species richness time trends. Biological Conservation, 219, 175–183. https://doi.org/10.1016/j.biocon.2017.12.021
Chase, J. M., & Knight, T. M. (2013). Scale-dependent effect sizes of ecological drivers on biodiversity: Why standardised sampling is not enough. Ecology Letters, 16, 17–26. https://doi.org/10.1111/ele.12112
de Vernal, A., Eynaud, F., Henry, M., Hillaire-Marcel, C., Londeix, L., Mangin, S., … Turon, J.-L. (2005). Reconstruction of sea-surface conditions at middle to high latitudes of the Northern Hemisphere during the Last Glacial Maximum (LGM) based on dinoflagellate cyst assemblages. Quaternary Science Reviews, 24, 897–924. https://doi.org/10.1016/j.quascirev.2004.06.014
Dornelas, M., Gotelli, N. J., McGill, B., Shimadzu, H., Moyes, F., Sievers, C., & Magurran, A. E. (2014). Assemblage time series reveal biodiversity change but not systematic loss. Science, 344, 296–299. https://doi.org/10.1126/science.1248484
Elahi, R., O’Connor, M. I., Byrnes, J. E. K., Dunic, J., Eriksson, B. K., Hensel, M. J. S., & Kearns, P. J. (2015). Recent trends in local-scale marine biodiversity reflect community structure and human impacts. Current Biology, 25, 1938–1943. https://doi.org/10.1016/j.cub.2015.05.030
Ezard, T. H. G., Quental, T. B., & Benton, M. J. (2016). The challenges to inferring the regulators of biodiversity in deep time. Philosophical Transactions of the Royal Society B: Biological Sciences, 371, 20150216. https://doi.org/10.1098/rstb.2015.0216
Fenton, I. S., Pearson, P. N., Dunkley Jones, T., Farnsworth, A., Lunt, D. J., Markwick, P., & Purvis, A. (2016). The impact of Cenozoic cooling on assemblage diversity in planktonic foraminifera. Philosophical Transactions of the Royal Society B: Biological Sciences, 371, 20150224. https://doi.org/10.1098/rstb.2015.0224
Fenton, I. S., Pearson, P. N., Dunkley Jones, T., & Purvis, A. (2016). Environmental predictors of diversity in recent planktonic foraminifera as recorded in marine sediments. PLoS ONE, 11, 1–22. https://doi.org/10.1371/journal.pone.0165522
Flower, B. P., & Kennett, J. P. (1995) Biotic responses to temperature and salinity changes during last deglaciation, Gulf of Mexico. In Effects of past global change on life (pp. 209–220). Washington, DC: National Research Council (US).
Fritz, S. A., Schnitzler, J., Eronen, J. T., Hof, C., Bönhning-Gaese, K., & Graham, C. H. (2013). Diversity in time and space: Wanted dead and alive. Trends in Ecology and Evolution, 28, 509–516. https://doi.org/10.1016/j.tree.2013.05.004
Gonzalez, A., Cardinale, B. J., Allington, G. R. H., Byrnes, J., Endsky, K. A., Brown, D. G., … Loreau, M. (2016). Estimating local biodiversity change but not systematic loss. Science, 355, 296–299. https://doi.org/10.1126/science.aah4787
Harnik, P. G., Lotze, H. K., Anderson, S. C., Finkel, Z. V., Finnegan, S., Lindberg, D. R., … Tittensor, D. P. (2012). Extinctions in ancient and modern seas. Trends in Ecology and Evolution, 27, 608–617. https://doi.org/10.1016/j.tree.2012.07.010
Herbert, T. D., Lawrence, K. T., Tzanova, A., Peterson, L. C., Caballero-Gill, R., & Kelly, C. S. (2016). Late Miocene global cooling and the rise of modern ecosystems. Nature Geoscience, 9(11), 843–847.
Hillebrand, H., Blasius, B., Borer, E. T., Chase, J. M., Downing, J. A., Eriksson, B. K., … Ryabov, A. B. (2018). Biodiversity change is uncoupled from species richness trends: Consequences for conservation and monitoring. Journal of Applied Ecology, 55, 169–184. https://doi.org/10.1111/1365-2664.12959
Hillebrand, H., Brey, T., Gutt, J., Hagen, W., Metfies, K., Meyer, B., & Lewandowska, A. (2018). Climate change: Warming impacts on...
marine biodiversity. In M. Salomon & T. Markus (Eds.), *Handbook on marine environment protection* (pp. 353–373). Heidelberg, Germany: Springer-Verlag GmbH.

Jackson, J. B. C. (2008). Ecological extinction and evolution in the brave new ocean. *Proceedings of the National Academy of Sciences USA*, 105, 11458–11465. https://doi.org/10.1073/pnas.0802812105

Jonkers, L., Hillebrand, H., & Kucera, M. (2019). Global change drives modern plankton communities away from the pre-industrial state. *Nature*, 570, 372–375. https://doi.org/10.1038/s41586-019-1220-3

Jonkers, L., & Kucera, M. (2015). Global analysis of seasonality in the shell flux of extant planktonic Foraminifera. *Biogeosciences*, 12, 2207–2226. https://doi.org/10.5194/bg-12-2207-2015

Kucera, M. (2007). Planktonic foraminifera as tracers of past oceanic environments. In C. Hillaire-Marcel & A. de Vernal (Eds.), *Proxies in late cenozoic paleoceanography*, (Developments in Marine Geology, Vol. 1, pp. 213–262). Elsevier.

Lotze, H. K., Lenihan, H. S., Bourque, B. J., Bradbury, R. H., Cooke, R. G., Kay, M. C., ... Jackson, J. B. C. (2006). Depletion, degradation, and recovery potential of estuaries and coastal seas. *Science*, 312, 1806–1809.

Maguran, A. E., Deacon, A. E., Moyes, F., Shimadzu, H., Dornelas, M., Phillip, D. A. T., & Rammarine, I. W. (2018) Divergent biodiversity change within ecosystems. *Proceedings of the National Academy of Sciences USA*, 115(8), 1843–1847.

McGill, B. J., Hadly, E. A., & Maurer, B. A. (2005). Community inertia of Quaternary small mammal assemblages in North America. *Proceedings of the National Academy of Sciences USA*, 102, 16701–16706. https://doi.org/10.1073/pnas.0504225102

Newbold, T., Hudson, L. N., Hill, S. L. L., Contu, S., Lysenko, I., Senior, R. A., ... Purvis, A. (2015). Global effects of land use on local terrestrial biodiversity. *Nature*, 520, 45–50. https://doi.org/10.1038/natur e14324

Nogués-Bravo, D., Rodríguez-Sánchez, F., Orsini, L., de Boer, E., Janssens, R., Morlon, H., ... Jackson, S. T. (2018). Cracking the code of biodiversity responses to past climate change. *Trends in Ecology and Evolution*, 33, 765–776. https://doi.org/10.1016/j.tree.2018.07.005

Pandolfi, J. M., & Jackson, J. B. (2006). Ecological persistence interrupted in Caribbean coral reefs. *Ecology Letters*, 9, 818–826. https://doi.org/10.1111/j.1461-0248.2006.00933.x

Poloczanska, E. S., Brown, C. J., Sydeman, W. J., Kiessling, W., Schoeman, D. S., Moore, P. J., ... Richardson, A. J. (2013). Global imprint of climate change on marine life. *Nature Climate Change*, 3, 919–925. https://doi.org/10.1038/nclimate1958

Raymo, M. (1994). The initiation of Northern Hemispher glaciation. *Annual Review of Earth and Planetary Sciences*, 22, 353–383. https://doi.org/10.1146/annurev.ea.22.050194.002033

Ren, J., Gersonde, R., Esper, O., & Sancetta, C. (2014). Diatom distributions in northern North Pacific surface sediments and their relationship to modern environmental variables. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 402, 81–103. https://doi.org/10.1016/j.palaeo.2014.03.008

Reymond, C. E., Bode, M., Renema, W., & Pandolfi, J. M. (2011). Ecological incumbency impedes stochastic community assembly in Holocene foraminifera from the Huon Peninsula, Papua New Guinea. *Paleobiology*, 37, 670–685. https://doi.org/10.1666/09087.1

Sax, D. F., & Gaines, S. D. (2003). Species diversity: From global decreases to local increases. *Trends in Ecology and Evolution*, 18, 561–566. https://doi.org/10.1016/S0169-5347(03)00224-6

Tomasových, A., Dominici, S., Zuschin, M., & Merle, D. (2014). Onshore-offshore gradient in metacommunity turnover emerges only over macroevolutionary time-scales. *Proceedings of the Royal Society B: Biological Sciences*, 281, 20141533.

Tomašových, A., & Kidwell, S. M. (2010). The effects of temporal resolution on species turnover and on testing metacommunity models. *The American Naturalist*, 175, 587–606. https://doi.org/10.1086/651661

Wade, B. S., & Pearson, P. N. (2008). Planktonic foraminiferal turnover, diversity fluctuations and geochemical signals across the Eocene/Oligocene boundary in Tanzania. *Marine Micropaleontology*, 68, 244–255. https://doi.org/10.1016/j.marmico.2008.04.002

Wood, S. N. (2017). Generalized additive models: An introduction with R (2nd ed.). Boca Raton, FL: CRC Press Taylor & Francis Group.

Yasuohara, M., Cronin, T. M., Hunt, G., & Hodell, D. A. (2009). Deep-sea ostracods from the South Atlantic sector of the Southern Ocean during the last 370,000 years. *Journal of Paleontology*, 83, 914–930. https://doi.org/10.1666/08-149.1

Yasuohara, M., Hunt, G., Breitburg, D., Tsujimoto, A., & Katsuki, K. (2012). Human-induced marine ecological degradation: Micropaleontological perspectives. *Ecology and Evolution*, 2, 3242–3268. https://doi.org/10.1002/ece3.425

Yasuohara, M., Hunt, G., Dowsett, H. J., Robinson, M. M., & Stoll, D. K. (2012). Latitudinal species diversity gradient of marine zooplankton for the last three million years. *Ecology Letters*, 15, 1174–1179. https://doi.org/10.1111/j.1461-0248.2012.01828.x

Yasuohara, M., Tittensor, D. P., Hillebrand, H., & Worm, B. (2017). Combining marine macroecology and palaeoecology in understanding biodiversity: Microfossil as a model. *Biological Reviews*, 92, 199–215. https://doi.org/10.1111/brv.12223

**BIOSKETCH**

Aleksandra M. Lewandowska is currently an assistant professor at the Tvärminne Zoological Station, University of Helsinki. Her research focuses on the impact of climate change on biodiversity in marine plankton. The highly interdisciplinary team includes experts on micropalaeontology and plankton ecology from the University of Bremen, and experts on biodiversity research and statistical data analysis from the University of Oldenburg, where the first author was previously affiliated.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

**How to cite this article:** Lewandowska AM, Jonkers L, Auel H, et al. Scale dependence of temporal biodiversity change in modern and fossil marine plankton. *Global Ecol Biogeogr*. 2020;29:1008–1019. https://doi.org/10.1111/geb.13078

**APPENDIX**

Andersson, C., et al. (2003). Late Holocene surface ocean conditions of the Norwegian Sea (Vøring Plateau). *Paleoceanography*, 18, 1044.
Asahi, H., & Takahashi, K. (2007). A 9-year time-series of planktonic foraminifer fluxes and environmental change in the Bering sea and the central subarctic Pacific Ocean, 1990–1999. Progress in Oceanography, 72, 343–363.

De Abreu, L., et al. (2003). Millennial-scale oceanic climate variability off the Western Iberian margin during the last two glacial periods. Marine Geology, 196, 1–20.

Deuser, W. G. (1987). Seasonal variations in isotopic composition and deep-water fluxes of the tests of perennially abundant planktonic foraminifera of the Sargasso sea: Results from sediment-trap collections and their paleoceanographic significance. Journal of Foraminiferal Research, 17, 14–27.

Deuser, W. G., & Ross, E. H. (1989). Seasonal abundance of planktonic foraminifera of the Sargasso Sea: Succession, deep-water fluxes, isotopic compositions, and paleoceanographic implications. Journal of Foraminiferal Research, 19, 268–293.

Deuser, W. G., et al. (1981). Seasonal changes in species composition, numbers, mass, size, and isotopic composition of planktonic foraminifera settling into the deep Sargasso Sea. Palaeogeography, Palaeoclimatology, Palaeoecology, 33, 103–127.

Field, D. B., et al. (2006). Planktonic foraminifera of the California Current reflect 20th-century warming. Science, 311, 63–66.

Herbert, et al. (2016). Late Miocene global cooling and the rise of modern ecosystems. Nature Geoscience, 9, 843.

Hüls, M., & Zahn, R. (2000). Millennial-scale sea surface temperature variability in the western tropical North Atlantic from planktonic foraminiferal census counts. Paleoceanography and Paleoclimatology, 15, 659–678.

Knudsen, K. L., Eiriksson, J., & Barteles-Jonsdottir, H. B. (2012). Oceanographic changes through the last millennium off North Iceland: temperature and salinity reconstructions based on foraminifera and stable isotopes. Marine Micropaleontology, 84–85, 54–73.

Marcott, S. A., et al. (2013). A reconstruction of regional and global temperature for the past 11,300 years. Science, 339, 1198–1201.

Martinez, J. I., Mora, G., & Barrows, T. T. (2007). Paleoceanographic conditions in the western Caribbean Sea for the last 560 kyr as inferred from planktonic foraminifera. Marine Micropaleontology, 64, 177–188.

Mojtahid, M., et al. (2015). Thirteen thousand years of southeastern Mediterranean climate variability inferred from an integrative planktic foraminiferal-based approach. Paleoceanography, 30, 402–422.

Morice, C. P., et al. (2012). Quantifying uncertainties in global and regional temperature change using an ensemble of observational estimates: The HadCRUT4 data set. Journal of Geophysical Research: Atmospheres, 117, D08101.

Munz, P. M., et al. (2017). The Indian winter monsoon and its response to external forcing over the last two and a half centuries. Climate Dynamics, 49, 1801–1812.

Peeters, F. J. C., et al. (2004). Vigorous exchange between the Indian and Atlantic oceans at the end of the past five glacial periods. Nature, 430, 661–665.

Poore, R. Z., Tedesco, K. A., & Spear, J. W. (2013). Seasonal flux and assemblage composition of planktic foraminifers from a sediment-trap study in the northern Gulf of Mexico. Journal of Coastal Research, 63, 6–19.

Rigual-Hernandez, A. S. (2012). Seasonal and interannual changes of planktic foraminifer fluxes in the Gulf of Lions (NW Mediterranean) and their implications for paleoceanographic studies: Two 12-year sediment trap records. Deep Sea Research Part I: Oceanographic Research Papers, 66, 26–40.

Salmon, K. H., et al. (2015). Upper ocean mixing controls the seasonality of planktonic foraminifer fluxes and associated strength of the carbonate pump in the oligotrophic North Atlantic. Biogeosciences, 12, 223–235.

Shakun, J. D., et al. (2012). Global warming preceded by increasing carbon dioxide concentrations during the last deglaciation. Nature, 484, 49–54.

Shakun, J. D., et al. (2015). An 800-kyr record of global surface ocean and implications for ice volume-temperature coupling. Earth and Planetary Science Letters, 426, 58–68.

Valletfuoco, M., et al. (2012). Climatic variability and anthropogenic signatures in the Gulf of Salerno (southern-eastern Tyrrhenian Sea) during the last half millennium. Rendiconti Lincei. Scienze Fisiche e Naturali, 23, 12–23.

Wang, L. (1994). Sea surface temperature history of the low latitude western Pacific during the last 5.3 million years. Palaeogeography, Palaeoclimatology, Palaeoecology, 108, 379–436.