The Dynamics of Diachronous Extinction Associated with Climatic Deterioration near the Neogene/Quaternary Boundary

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Key Points:

* We report size and abundance data of 16,091 individual planktonic foraminifer *Globoconella puncticulata* specimens preceding extinction
* Population extinctions are diachronous and do not follow a North-to-South sequence as glaciation of the Northern Hemisphere intensified
* Population dynamics in core populations differ from populations at the edge of their range

**Key words**

Pliocene, Northern Hemisphere glaciation, extinction, biogeographic range contraction, pre-extinction dwarfing, planktonic foraminifera

Abstract

To predict extinction we must understand the processes leading to terminal population decline. Once a critical threshold of population size is reached, small environmental perturbations can push a species over the cliff-edge to extinction, so the main drivers of extinction are the factors that cause the initial reduction in population size. Most studies of population decline leading up to extinction focus on modern species in a human-dominated world. The drivers of population decline leading to non-human mediated extinctions are less well known but changes in climate are arguably the most widely invoked mechanism. Here, we report data on >16,000 individuals of the planktonic foraminifer *Globoconella puncticulata* from six sites in the Atlantic Ocean along a 83 degree-long latitudinal transect, over a 600,000-year interval leading up to the species’ global extinction during the late Pliocene-earliest Pleistocene intensification of Northern Hemisphere glaciation. We show changes in geographic range, abundance and body size. We find that populations do not follow a North-to-South sequence in extinction as Earth cooled and developed large ice sheets in the high latitudes of the Northern Hemisphere. Instead, our results suggest that (1) populations are differentially adapted to local environmental conditions such as nutrient availability, (2) population dynamics in core populations differ from those at the edge of their range, and (3) individual population responses to external pressures are essential to understanding the drivers of global extinction. Our study demonstrates the potential to transform our understanding of extinction dynamics through spatially replicated sampling of the highly-resolved marine microfossil record.

# Introduction

To predict extinction it is crucial to understand the processes leading to population decline. Decreasing abundance, range contraction and habitat fragmentation lead to smaller population size, which increases vulnerability to environmental perturbations, demographic noise (i.e., population-level random variation) and reduced fitness due to inbreeding (Hedrick et al., 1996). As populations decline past a critical threshold they end up in a downward spiral of positive feedbacks called an extinction vortex (Gilpin and Soulé, 1986) to the point where small external perturbations can push a species over the cliff-edge to extinction (Gilpin and Soulé, 1986; Hedrick et al., 1996; Raup, 1981). Because these modest perturbations would likely have been more readibly absorbed by healthy populations, the main drivers of extinction are the factors that reduce population size initially and thus enhance extinction risk (Hedrick et al., 1996; Raup, 1981). Therefore, to understand extinction, long-term records of biotic and abiotic change preceding extinction are needed.

Geographic range size reduction is one of the main factors that increase extinction risk (Purvis et al., 2000; Saupe et al., 2015). Range contractions reduce the total population size, decreasing genetic variation. For example, a study of alpine chipmunks showed declining genetic diversity with reducing range size over the last century (Rubidge et al., 2012). Another study showed decreased fitness due to reduced genetic diversity and inbreeding in a declining population of southern dunlins (Blomqvist et al., 2010). These effects can be exacerbated by population fragmentation, which reduces gene flow even further (Gilpin, 1987). Geographic range contractions are suggested to fall into one of two models: the demographic hypothesis, which predicts that the range implodes towards the core of the historical range, and the contagion hypothesis, which expects the range to shrink away from local negative impacts (Channell and Lomolino, 2000) (Figure 1). The demographic hypothesis assumes that the intensity of environmental change is uniform across the range, whereas the contagion hypothesis assumes a single, local extinction factor, with chances of survival increasing further away from the source of the perturbation (Channell and Lomolino, 2000). It is not clear which of these hypotheses applies more commonly in nature. Range contractions preceding modern extinctions are often best described by the contagion hypothesis (e.g. Boakes et al., 2018; Lomolino and Channell, 1995; Rando et al., 2011; Steadman et al., 2005) but in the majority of these cases human influences were the cause of extinction. To understand the drivers of extinctions where humans are not the main agent of forcing, we need to study range size dynamics further back in time.

Declining populations often experience changes in morphology, either as a result of reduced genetic and/or morphological variation, through inbreeding or otherwise, or as a direct response to external biotic or abiotic pressures. Body size has been particularly well studied in this regard, with reductions in average size reported at both the community and species levels. For example, average size in macroinvertebrate assemblages has been found to decrease immediately prior to the end-Permian mass extinction (Kiessling et al., 2018; Wu et al., 2018; Zhang et al., 2016; Zhang et al., 2017) and up to several hundred thousand years before the early Toarcian extinction event (García Joral et al., 2018; Piazza et al., 2019). Pre-extinction dwarfing of individual species has been noted in planktonic foraminifera (Wade and Olsson, 2009) and Pleistocene megafauna (Guthrie, 2003). Shifts in body size can precede population decline, which has recently led several studies to conclude that a combination of population abundance and body size is a more reliable early warning signal of population collapse than population size alone (Baruah et al., 2019; Clements and Ozgul, 2016). Baruah et al. (2019) found that this is most likely when environmental change was slow relative to generation time. Trait changes can buffer the negative effects of environmental change for a while, until a critical threshold is crossed and adaptation is no longer possible. Thus, when studied quantitatively, traits such as body size provide a promising means to study extinction dynamics in deeper time, especially because environmental change in the geological past tends to be slow relative to generation time (the immediate environmental consequences of meteorite impacts are the most obvious exception).

Studies of geographic range, population abundance and morphology through time and space require highly resolved fossil records. Most fossil records lack this resolution, but marine microfossils are exceptional. Their large population size, global distribution and excellent preservation potential make it possible to generate abundance records and morphological reconstructions at resolutions exceeding environmental change over millions of years. Additionally, environmental reconstructions from the same sample medium allow for direct comparisons of biotic and abiotic change. Here, we study range size, abundance and morphology of the planktonic foraminifer *Globoconella puncticulata* preceding its extinction. This globally distributed Pliocene species became extinct around 2.4 Ma (Wei, 1994), shortly after Earth’s last major shift in global climate state: the intensification of Northern Hemisphere glaciation (iNHG, ~3.5 to 2.5 Ma) (Bailey et al., 2013; Westerhold et al., 2020). During this interval, Earth transitioned from a unipolar glacial state, with large ice sheets only on Antarctica (Spray et al., 2019), to one that was sufficiently cool by ~2.5 Ma to induce growth of large ice sheets in the Northern Hemisphere (Shakun et al., 2016) and associated rafting of icebergs across the subpolar North Atlantic Ocean (Shackleton et al., 1984). Previous work on *G. puncticulata* has demonstrated diachronous population extinctions around the world (Scott et al., 2007), a decrease in test size prior to extinction in a mid-latitude population (Brombacher et al., 2017a) and cyclical variations in abundance corresponding to glacial-interglacial cycles in a sub-polar population (Chapman et al., 1998). We studied 16,091 specimens of *G. puncticulata* from six Atlantic populations, over a 600,000-year time interval preceding its extinction at ~2.4 Ma to investigate the consistency of these results along a latitudinal transect (Figure2a). In particular, we address the following questions:

*1. Does the geographic range contract before extinction?*

We investigate extinction timing across populations. Where extinction is diachronous, we test the competing predictions of the demographic hypothesis (ranges implode towards the core of the historical range) and the contagion hypothesis (range shrinks away from local negative impacts).

*2. Does shell size decrease before extinction?*

One mid-latitude population of *G. puncticulta* showed a decrease in size prior to extinction (Brombacher et al., 2017a). Here, we investigate whether size changed consistantly across populations to test whether size changes are a reliable early warning signal of extinction in other/all populations.

*3. Do populations respond to the same environmental parameters similarly?*

The demographic hypothesis assumes that response to environmental change is uniform across the species’ range. Here, we compare local climate reconstructions to individual populations’ abundance and size to test if this was the case, and to determine whether any differences among populations can be explained by differential response to local environmental perturbations.

# Materials and Methods

## Study species

The planktonic foraminifer species *Globoconella puncticulata* (Deshayes, 1832) first appeared in the early Pliocene around 4.6 Ma and became extinct at 2.41 Ma (Wei, 1994), shortly after the Neogene/Quaternary boundary (~2.58 Ma) and the culmination of the late Pliocene-earliest Pleistocene intensification of Northern Hemisphere glaciation at ~2.5 Ma (Bailey et al., 2013). The species was globally distributed with highest abundance in thermocline waters at middle and low latitude sites (Aze et al., 2011; Kennett and Srinivasan, 1983). A review by Scott et al. (2007) of first and last occurrences in Atlantic, Mediterranean, Pacific and Southern Ocean sites shows asynchronous population extinctions of *G. puncticulata*, suggesting local environmental controls on population survival. Chapman et al. (1998) reported cyclical variations in relative abundance of a subpolar population in response to glacial/interglacial cycles, and a mid-latitude Atlantic population showed a decrease in test size ~300,000 years prior to extinction (Brombacher et al., 2017a), possibly indicating pre-extinction dwarfing (Wade and Olsson, 2009).

## Study sites

We studied specimens of *G. puncticulata* from six Atlantic Ocean sites (Figure 2) spanning ~83˚latitude:

* Ocean Drilling Program (ODP) Site 981 (Jansen et al., 1996): a sub-polar Northern Hemisphere site (55° N) currently situated under the North Atlantic Current.
* Integrated Ocean Drilling Program (IODP) Site U1313 (Channell et al., 2006): a mid-latitude Northern Hemisphere site (41° N) situated at the northern edge of the oligotrophic North Atlantic Subtropical Gyre.
* Deep Sea Drilling Program (DSDP) Site 606 (Ruddiman et al., 1987): a mid-latitude Northern Hemisphere site located just south of Site U1313 within the North Atlantic Subtropical Gyre (37° N). Site U1313 and Site 606 are situated closest to one another geographically and are the most similar of the sites studied oceanographically.
* ODP Site 659 (Ruddiman et al., 1988): a subtropical Northern Hemisphere site (18° N) situated in the Canary Current off the Northwest African margin.
* ODP Site 925 (Curry et al., 1995): a western equatorial site (3° N) located in the Southern Equatorial Current transporting South Atlantic surface waters fed from the Benguela Current into the North Atlantic. This site experiences high export production due to nutrient-rich Amazon river water runoff.
* ODP Site 1264 (Zachos et al., 2004): a subtropical Southern Hemisphere site (28° S) located at the eastern edge of the South Atlantic Subtropical Gyre in the Benguela Current. Of all study sites, this site is least directly affected by climate feedbacks associated the intensification of Northern Hemisphere glaciation(Zachos et al., 2004).

For age control, we used previously published site-specific astronomically resolved benthic foraminiferal δ18O stratigraphies (Bell et al., 2014; Bickert et al., 1997; Bolton et al., 2010; Franz and Tiedemann, 2002; Mc Intyre et al., 1999; Sikes et al., 1991; Tiedemann et al., 1994) which have been included in (Sites 606, 659, 981 and 925), or tuned to (Sites 1264 and U1313) the LR04 δ18O stack (Lisiecki and Raymo, 2005). Samples from ODP Sites 606, 659, 981 and 925 were acquired from the IODP Bremen Core Repository in Germany. The samples from IODP Sites 1264 and U1313 were originally used in Bell et al. (2014) and Bolton et al. (2010), respectively. To avoid time averaging of foraminifera among samples due to bioturbation, samples were spaced at least 15 cm apart corresponding to temporal resolutions of 5-20 kyr (see Table 1 for site-specific details). Calcareous microfossil preservation in sediments of late Pliocene age is good to excellent at all sites (Channell et al., 2006; Curry et al., 1995; Jansen et al., 1996; Ruddiman et al., 1987; Ruddiman et al., 1988; Zachos et al., 2004).

Table 1 Sampling resolution and number of samples analysed per site

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Site | Depth resolution (cm) | Temporal resolution (kyr) | Number of samples | Number of specimens |
| 981 | 45 | 10 | 37 | 1661 |
| U1313 | 30 | 5 | 75 | 6575 |
| 606 | 60 | 20 | 36 | 2398 |
| 659 | 60 | 10 | 57 | 3968 |
| 925 | 60 | 20 | 19 | 138 |
| 1264 | 15 | 20 | 29 | 1351 |

## Analysis

### Abundance and geographic range size

All samples were dry-sieved over a >150 μm mesh sieve and divided using a Green Geological aluminium microsplitter until a single split contained 50-100 adult individuals of *G. puncticulata*, all of which were picked to avoid size biases. 16,091 specimens of *G. puncticulata* were analysed. To enable direct comparisons of abundance among sites, abundance per time slice was calculated as flux rather than relative abundance. The estimated total number of specimens in a sample (the number of individuals found in an analysed split multiplied by the split fraction) was divided by sample weight, dry bulk density and the total time contained in the sample with final numbers expressed as individuals/cm2/kyr.

### Morphology

To measure test size, foraminiferal shells were mounted on glass slides in groups of 20 individuals using transparent double-sided adhesive tape with the apertures facing upwards. Groups were imaged using an Infinity 3 Lumenera camera mounted on an Olympus SZX10 light microscope, and test size was extracted from the images using an automated image analysis macro in the Image Pro Premier software. We measured size as test area. Previous error analyses of a subset of the data presented here have shown that test area is a repeatable measure (Brombacher et al., 2017b) and a more accurate representation of three-dimensional body size than test diameter (Brombacher et al., 2018a). All data are privately accessible through this link: <https://figshare.com/s/29f093ebba6abee22f18>, and will be made publicly available upon acceptance of the manuscript.

### Response to local environmental change

Abundance and morphology of one mid-latitude population of *G. puncticulata* have been shown to respond to a combination of environmental factors, rather than single variables (Brombacher et al., 2018b). Therefore, we study biotic response to multiple abiotic drivers. Planktonic foraminifera are known to have preferred temperature ranges (Lombard et al., 2011; Lombard et al., 2009) and growth rates are influenced by food availability (Bé et al., 1981; Takagi et al., 2018). Additionally, ocean pH influences selection for larger shell size and thickness with decreasing pH (Hemleben et al., 1989).

To avoid offsets among local environmental records it is important to compare population response to environmental change reconstructed using the same proxies. Two of our study sites met these criteria: Site U1313 has high-resolution reconstructions of sea surface temperature (SST) and export productivity available for our study interval, and SST and export productivity reconstructions from Site 982 were used as indicators of local environment at Site 981 given the close proximity of these two sites (Jansen et al., 1996). Reconstructions of sea surface temperature are based on the alkenone unsaturation index Uk’37 (982: Lawrence et al., 2009; U1313: Naafs et al., 2010) and productivity records are reconstructed from C­37 alkenone mass accumulation rates (C37 MAR) (982: Bolton et al., 2011; U1313: Naafs et al., 2010). Additionally, we used a record of atmospheric CO2 (Martínez-Botí et al., 2015) to compare biotic response to global as well as local environmental change. Where climate data point ages were offset relative to our new biotic data, we followed the approach of Brombacher et al. (2018b) using Generalised Additive Models (GAMs) to interpolate the climate records to the foraminiferal sample ages by applying a smooth function to the climate data (R package ‘mgcv’, version 1.8.33 (R Core Team, 2013; Wood, 2006). The response variables in a GAM depend linearly on the additive combination of polynomial functions multiplied by statistically estimated coefficients (Wood, 2006) making them more suitable to interpolate environmental data (which often do not vary linearly with time) than simple linear interpolations. This approach is capable of distinguishing the glacial-interglacial dynamics of our study interval (Brombacher et al., 2018b). Although including only two of our six study sites in this analysis will not allow us to quantify the full range of responses of *G. puncticulata* to climate change across its geographical range, this is not necessary to distinguish between the two competing hypotheses of the demographic and contagion hypotheses. The two sub-Arctic and mid-latitude sites experienced the largest environmental change across the species’ range due to their proximity to the Northern Hemisphere ice sheets (Herbert et al., 2010; Lawrence et al., 2009; Naafs et al., 2010), providing an opportunity to study biotic response to the maximum environmental change experienced by both a core- and edge- of range population. The multiproxy climate records at these two sites allow investigation of synergistic climate effects (Brombacher et al., 2018b) to provide a first indication of possible variation among populations.

Environmental conditions at sites U1313 and 982 were influenced by both global and local factors. SST records at both sites show increasing glacial-integlacial variation through our study interval but, whereas Site 982 also experienced an overall decreasing trend in SST, only average glacial temperatures decreased at Site U1313 (Lawrence et al., 2009; Naafs et al., 2010). Changes in ocean circulation and global atmospheric CO2 decline were responsible for SST fluctuations at both sites (Friedrich et al., 2013; Lawrence et al., 2009; Naafs et al., 2010), and glacial-interglacial SST differences at Site 982 were further amplified by the site’s close proximity to the Northern Hemisphere ice sheets (Lawrence et al., 2009). Transient peaks in export productivity are caused by increased wind-driven mixing of deeper, nutrient-rich waters at Site 982 (Bolton et al., 2011) whereas southward deflections of the subarctic front and increased glacial dust delivery enhanced productivity at Site U1313 (Bolton et al., 2018; Naafs et al., 2012; Naafs et al., 2010).

Our study sites are situated in different palaeoceanographic settings that respond differently to environmental change (Figure 2a). To test the similarities and differences in population response to the same underlying environmental driver, the intensification of Northern Hemisphere Glaciation (iNHG), and its local expression we used a Linear Mixed Effect model using the R package ‘nlme’ (version 3.1.152) (Pinheiro et al., 2021; R Core Team, 2013). This model looks for linear trends among subsets of the data (here: populations) by comparing support for six different scenarios:

1. no ocean-wide climate trends, random effects or variance differences between sites (model input: trait ~ 1),
2. no ocean-wide climate trends or random effects but variance differences between sites (model input: trait ~ 1 | site)
3. no ocean-wide climate trends, but site-specific intercept differences and variance differences (model input: trait ~ 1 + (1 | site) )
4. no ocean-wide climate trends, but site-specific intercept differences, temperature relationships and variance differences (trait ~ 1 + (sst | site) )
5. ocean-wide climate trends, site-specific intercept differences and variance differences (trait ~ climate + (1 | site) );
6. ocean-wide climate trends, site-specific intercept differences, temperature relationships and variance differences (trait ~ climate + (sst | site) )(see also Figure S1).

Models without random effects were fitted using generalized least squares models. Variance differences among sites were assessed using the varIdent function in nlme (Pinheiro and Bates, 2000). The model with the lowest Akaike Information Criterion (AIC) value (a measure for model parsimony) and the highest Akaike weight (a measure for relative model likelihood) has the most statistical support among those models fitted. ‘Climate’ was taken as the combination of local SST and export productivity and global atmospheric CO2, and their two-way interactions (model input: climate = (SST + productivity + CO­2)2, with the squared term calculating two-way interactions between the environmental parameters). A visual inspection of the residuals shows that they are approximately normally distributed, satisfying the model prerequisites (Supplementary Figures S2-S5).

# Results

Abundance varies among sites (Figure 3a-d). Abundance is high and stable at mid-latitude sites U1313 and 606 as well as subtropical sites 659 and 1264, but has a lower mean with greater variability at subpolar Site 981 and equatorial Site 925. Based on these observations, sites U1313, 606, 659 and 1264 are identified as core populations, whereas sites 981 and 925 are inferred to sit at the edge of the species’ preferred environmental conditions. Populations at sites 981 and 925 experienced several extirpations (temporary local extinctions) before their final extinction (Figure 3a,d). At the subpolar site (981), several brief extirpations occurred before final population extinction at 2.44 Ma. In contrast, *G. puncticulata* disappeared from tropical Site 925 for much longer intervals of up to ~100 kyr.

Final population extinctions are spread over a ~100 kyr-long interval starting at 2.44 Ma with subpolar Site 981 and ending with the subtropical population at Site 659 at 2.34 Ma (Figure 4, Figure S6). First, the species’ northern range contracts to lower latitudes following extinctions at Sites 981 and U1313 (2.42 Ma). 10 kyr later the tropical population at Site 925 becomes extinct, resulting in reduced connectivity between Northern and Southern Hemisphere populations. The populations at Sites 606 and 1264 disappear soon afterwards (2.39 and 2.37 Ma respectively), leaving an isolated North Atlantic subtropical population at Site 659 that persists for another ~30,000 years until its extinction at 2.34 Ma.

Size also varies across space. The largest specimens are found at mid-latitude sites U1313 and 606 before 2.7 Ma, intermediate-sized specimens occur at Sites 659, 1264 and 981 and the smallest specimens come from Site 925 (Figure 3e-h). Only sites U1313 and 606 experience a sudden change in size similar to pre-extinction dwarfing around 2.7 Ma (Figure 3f). No dwarfing is observed in any of the other four sites.

The two populations at sites with environmental reconstructions available (sites U1313 and 981) show different responses to local abiotic change (Figure 5). The final extinction as well as two temporary extirpations at Site 981 occur during SST minima associated with MIS 96 (extinction), 100 and G6 (extirpations), whereas abundance at Site U1313 shows no response to local temperature or productivity. In contrast, mean test size at Site U1313 decreases during the SST minimum and productivity maximum associated with MIS G6 but no direct size response to temperature or productivity is observed at Site 981. The Linear Mixed Effects model results indicate no trend in the pooled abundance data of sites U1313 and 981, but response to local environment differs between the sub-polar and mid-latitude populations (Table 2). The biggest improvement in model fit comes from including abundance variance differences between sites (Model 2) as well changes to the mean abundance – the individuals at Site 981 are both smaller and, on a log10 scale, more variable than at U1313.

Size shows both a detectable trend in the pooled data, as well as site-specific trends of different magnitude (Scenario 6, Table 3). Unlike the abundance models, there is a smaller but still statistically significant impact of variance differences between the sites (cf Models 1 and 2). The impact of the variance differences is however much smaller than including global trends in the climate variables and their interactions (cf Models 3, 5 and 6) – that the biggest jump in AIC scores is through the climate variables as the fixed rather than random effects implies much stronger global controls on size than abundance (cf, Tables 2 and 3). There are clear site-specific differences in how these planktonic foraminifera respond to sea-surface temperature changes (cf the improvement from Model 5 to 6 in Table 3), supporting conclusions that the organismal response to local environmental settings is not constant through space and time.

Table 2 Linear Mixed Effect model results of *Globoconella puncticulata* abundance response to local environmental change. ΔAIC represents the difference between AIC and the set’s minimum AIC. The best performing model based on lowest AIC and highest Akaike weight is indicated in bold. See section 2.3.3 for model descriptions, and Figures S2 and S3 for residual distributions of the best performing model.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Model | Log Likelihood | Degrees of freedom | AIC | ΔAIC | Akaike Weight |
| 1: flux ~ 1 | -20.85 | 2 | 45.70 | 55.07 | 0 |
| 2: flux ~ 1 | site | -0.93 | 3 | 7.87 | 17.24 | 0 |
| **3: flux ~ 1 + (1 | site)** | **8.68** | **4** | **-9.37** | **0** | **0.861** |
| 4: flux ~ 1 + (sst | site) | 8.86 | 6 | -5.71 | 3.66 | 0.138 |
| 5: flux ~ climate2 + (1 | site) | -5.09 | 10 | 30.18 | 39.55 | 0 |
| 6: flux ~ climate2 + (sst | site) | -5.03 | 12 | 32.18 | 41.55 | 0 |

**Table 3** Linear Mixed Effect model results of *Globoconella puncticulata* size response to local environmental change. ΔAIC represents the difference between AIC and the set’s minimum AIC. The best performing model based on lowest AIC and highest Akaike weight is indicated in bold. See Figures S4 and S5 for residual distributions.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Model | Log Likelihood | Degrees of freedom | AIC | ΔAIC | Akaike Weight |
| 1: area ~ 1 | 8001.85 | 2 | -15999.71 | 331 | 0 |
| 2: area ~ 1 | site | 8007.76 | 3 | -16009.51 | 321.2 | 0 |
| 3: area ~ 1 + (1 | site) | 8008.97 | 4 | -16009.93 | 320.78 | 0 |
| 4: area ~ 1 + (sst | site) | 8043.40 | 6 | -16074.81 | 255.9 | 0 |
| 5: area ~ climate2 + (1 | site) | 8168.41 | 10 | -16316.81 | 13.9 | 0.001 |
| **6: area ~ climate2 + (sst | site)** | **8177.35** | **12** | **-16330.71** | **0** | **0.999** |

# Discussion

## Range contraction

The range contraction of *G. puncticulata* preceding extinction does not clearly follow predictions consistent with either the demographic or the contagion hypothesis. The demographic hypothesis predicts a range contraction to the core (e.g. edge sites 981 and 925 first, followed by core populations U1313, 6060, 659, 1264), and while one core population (Site 659) is indeed the last to become extinct, the core populations at Sites U1313 and 606 disappear much earlier (Figure 4, Figure S6). Similarly, the contagion hypothesis predicts extinctions away from the presumed extinction driver, in this case the environmental response to the expansion of Northern Hemisphere ice sheets (e.g. sub-Arctic Site 981 first, followed by mid-latitudes sites U1313 and 606, subtropical sites 659 and 1264 and finally tropical Site 925), and while the first two extinctions do follow this pattern it breaks down with the subsequent extinction of the equatorial population at Site 925. A combination of the two hypotheses better explains the data: the sub-polar (Site 981) and equatorial (Site 925) edge populations disappear early on as predicted by the demographic hypothesis, but the extinction of Northern Hemisphere core populations U1313, 606 and 659 follows a North-South pattern resembling the contagion hypothesis. These results suggest that population dynamics in core populations differ from those at the edge, and that biogeographic variation can obscure the detection of extinction driving mechanisms.

Our results differ from many studies of population range contractions. A review by Channell and Lomolino (2000) found that most modern ranges contract according to the contagion hypothesis. However, in most of these cases the extinction pressure came from humans. For example, Steadman et al. (2005) showed that, for late Quaternary sloth populations, extinction matched human arrival rather than global climate change and extinction of modern Galliform populations occurred close to the core in a human-dominated landscape (Boakes et al., 2018). Boakes et al. (2018) also showed that, in the absence of humans, extinction was equally likely to occur near the core or the edge of the geographical range, offering support for both the demographic and extinction hypotheses. Our data strongly suggest that contagion and demographics both played a role in extinction, but data from more extensive geological intervals and different types of species are needed to test the generality of this result.

A combination of the demographic and contagion hypothesis, rather than either of the theoretical ‘endmember’ extinction hypotheses on their own, might offer a more plausible explanation of extinction in general. The demographic hypothesis is based on the observation that populations are larger and more variable in the centre of their range than at the edge, which makes edge populations the most vulnerable to extinction (Channell and Lomolino, 2000). To test if an extinction follows the demographic hypothesis, the extinction factor must be synchronously and uniformly distributed across space (Channell and Lomolino, 2000). However, except in rare cases of species with an extremely small range, environmental change is never homogenous across the range. If certain areas are affected more than others, these populations would be more likely to follow the contagion hypothesis, calling into question the extent to which extinctions will ever solely resemble the demographic hypothesis. Similarly, the contagion hypothesis predicts extinction closest to the external stressor, but core populations might be more resilient to external perturbations than edge populations, decreasing detectability while also introducing elements of the demographic hypothesis.

## Abundance

In contrast to Chapman et al. (1998), we did not find clear cyclical variations in abundance as a response to glacial-interglacial cycles. This could be explained by different measures of abundance: we report abundance as absolute abundance or flux (individuals per area per kyr) whereas Chapman et al. (1998) measured abundance of *G. puncticulata* relative to other species in the assemblage. Relative abundance is a measure susceptible to closed sum problems. In the subpolar North Atlantic, decreases in the relative abundance of *G. puncticulata* during the early Northern Hemisphere glacials may well have been driven by increases in the abundance of cold-adapted species such as *Neogloboquadrina pachyderma*, *Neogloboquadrina atlantica* and *Neogloboquadrina incompta*. Flux on the other hand is more susceptible to age model uncertainty than relative abundance. Based on a maximum uncertainty of ~6 kyr in the LR04 age model in our time interval (Lisiecki and Raymo, 2005), and assuming one tie point per glacial-interglacial transition (i.e. one tie point per 20 kyr, see (Bolton et al., 2010)), the time contained in a sample and thus the flux could vary by a factor of ~0.25. However, as our flux patterns vary by 2-3 orders of magnitude (Figure 3a-d) the uncertainty introduced by the age model is unlikely to explain the absence of glacial-interglacial cyclicity in our data.

The use of relative abundance could also explain the different extinction timing we found for Site 659 as compared to Chapman et al. (1998). Chapman et al. (1998) reported extinction of the subtropical population at 2.44 Ma, similar to other populations, whereas we found *G. puncticulata* there for another 100,000 years until 2.34 Ma despite similar sampling resolutions. Assemblage counts typically analyse 300-500 specimens per sample, whereas here we included 50-100 specimens of the study species alone, or all specimens in a sample if there were less than 50. This approach is more likely to find (temporarily) rare species. Abundance of *G. puncticulata* decreased by a factor 10 on average in the last 100,000 years before extinction at Site 659, so it is possible that an assemblage study would have been less likely to find the species on the approach to extinction.

## Pre-extinction dwarfing

Two of the six studied populations (sites U1313 and 606) show a sudden size decrease consistent with pre-extinction dwarfing as described by Wade and Olsson (2009) (Figure 3f). Specimens showing pre-extinction dwarfing in that study mostly came from core populations, so it could be that dwarfing does not occur uniformly across space. Another possibility is that individuals at edge sites are smaller than those at core sites, resulting in size changes that are more difficult to detect due to the relative importance of different climate drivers (Brombacher et al., 2018b). Our Linear Mixed Effect model results support this explanation, showing different responses of size to local environmental change between populations at Sites U1313 and 981. More data on synchronicity of size changes across populations are needed before firm conclusions can be drawn, but our results suggest that the expression of early warning signals may vary across populations. In particular, a lack of change in one population does not necessarily imply low extinction risk or an absence of early warning signals for the species as a whole given the importance of the local palaeoceanographic context.

## Causes of extinction

The differences in population dynamics through space and time that we document suggest that the extinction of *G. puncticulata* cannot readily be explained by a single driver. If temperature was the main driver we would expect to see cyclical abundance variations in all populations with local extirpations during severe glacial stages. The subpolar population at Site 981 followed this pattern with temporary extirpations during glacials MIS G6 (2.72 Ma) and 100 (2.54 Ma, see Figure 5), but no other populations show similar responses. As predicted by both the demographic and contagion hypotheses, the population at Site 981 disappears first, but after that there is also no clear chain of extinction from North to South. Core sites U1313, 606 and 659 disappear in latitudinal sequence, but the equatorial edge population (Site 925), which under the contagion hypothesis should thrive under cooler conditions resembling the core environment, became extinct shortly after the first mid-latitude population disappeared. These results suggest that individual populations develop local adaptations that do not translate simply across space.

Local adaptation is further supported by the results of our Linear Mixed Effects models, which show significant differences in population response to local environment. As we analysed response to climate change in only two populations the species’ full range of response is not quantified, but the different responses observed at two relatively close sites suggests that differential response among populations to the same environmental parameters is not uncommon. These results agree with previous studies that found individualistic population response to climate change (Levin, 2019; Stewart, 2009; Stewart et al., 2010). Stewart (2009) found that Quaternary populations respond individualistically to biotic interactions through time and space. Stewart et al. (2010) also showed that the location of refugia following range contraction depends on population-specific adaptations and is therefore not always located in the core of the initial range. Levin (2019) argued that existing range contraction hypotheses oversimplify extinction precisely because they do not take local adaptation into account. Instead, Levin (2019) proposes that distinct intraspecific lineages are the focal point of species’ extinction, and that only by combining all intraspecific lineage responses can we understand the extinction of species.

Population-specific response to environmental change increases chances of survival of at least some populations (Levin, 2019). In our data however, low abundance and suboptimal adaptation to local environmental change in edge populations likely led to an increasingly fragmented species distribution. Temporary extirpations of the equatorial population (Site 925) at 2.8 Ma and 2.7 Ma (Figure 3d) resulted in an isolated Northern Hemisphere population with reduced global gene flow. The gradual extinction chain of the core populations briefly resulted in a subtropical refugia at Site 659 which persisted for another 30,000 years after the other populations had become extinct. The resistance to extinction at this site may be attributed to a combination of high abundance and phenotypic variability typical for a core population which reduces extinction risk (i.e. demographic hypothesis). Alternatively, being located furthest away from northern hemisphere ice influences of all northern hemisphere core populations (i.e. contagion hypothesis), or an optimal, site-specific microclimate may have made Site 659 a refuge for *G. puncticulata*. Populations in refugia, no matter how well adapted initially, are vulnerable to extinction because of their small population size and low genetic diversity, and absence of gene flow and migration from other areas following population decline (Blomqvist et al., 2010; Rubidge et al., 2012). Abundance at Site 659 is highly variable in its final 100,000 years (Figure 3c), suggesting increased population instability before extinction.

## Directions for future work

To understand extinction dynamics in deep time, data from many more microfossil groups are needed. Studies of extinction through time and space require high-resolution fossil records from multiple locations with high abundance of complete fossils. Most fossil records lack this resolution, but those of marine microfossils are an exception. The application of marine microfossils to biostratigraphy has already provided insights in spatial dynamics preceding extinction: it is well known that some species go extinct almost simultaneously around the world and can be used as global biohorizons (e.g. Falzoni et al., 2018; Gibbs et al., 2005; Huber et al., 2017; Wade et al., 2011) whereas others undergo extinction diachronously and are of local stratigraphic use only (e.g. Du et al., 2020; Falzoni et al., 2018; Gibbs et al., 2005; Huber et al., 2017; Kawagata et al., 2005; Prell and Damuth, 1978). However, these patterns are rarely extensively studied from a biological perspective. Global climate change is occassionally mentioned as a cause of varying population extinction timings (Gibbs et al., 2005; Prell and Damuth, 1978), but range contractions and individual population response to local environmental change are rarely quantified. Here, we have provided an example of how data on microfossils and climate can provide insights into population dynamics preceding extinction in geologic time.

# Conclusion

The extinction of *G. puncticulata* was preceded bychanges in the geographic range, abundance and body size that varied among populations. Range contraction prior to extinction showed characteristics of both the demographic (contraction towards the core) and contagion hypotheses (shrinking away from unfavourable conditions). The two northern-most populations disappeared first in keeping with the contagion hypothesis but a sub-tropical core population outlasted a tropical edge population before ultimately becoming extinct, suggesting influences of the demographic hypothesis as well. These two broad competing hypotheses are very likely a false dichotomy in many cases – here the response to local environmental change varied between a subpolar and a mid-latitude population, supporting previous inferences that populations are differentially adapted to temperature, nutrient availabibility and other microclimatic factors. Ultimately, global extinction can only be understood by unravelling the dynamics of individual populations and by understanding how how their dynamics are linked through space and time. To test the generality of our results, more data are needed from foraminifera and other fossil groups through different time intervals and climatic settings. The method of co-registered marine microfossil and palaeoclimate data is well-poised to yield transformative new insights into extinction dynamics.

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# Open Research

Data for this research will be available in this in-text citation reference: Brombacher et al. (2021) The dynamics of diachronous extinction associated with climatic deterioration near the Neogene/Quaternary boundary (<https://figshare.com/s/29f093ebba6abee22f18>, DOI: 10.6084/m9.figshare.7735094 (DOI will become active after publication)). All data are currently privately accessible through the link above and will be made publicly available upon acceptance of the manuscript.

# Conflict of interest

The authors declare no conflict of interests

References

Aze, T., Ezard, T.H., Purvis, A., Coxall, H.K., Stewart, D.R., Wade, B.S., Pearson, P.N., 2011. A phylogeny of Cenozoic macroperforate planktonic foraminifera from fossil data. Biological reviews of the Cambridge Philosophical Society 86, 900-927.

Bailey, I., Hole, G.M., Foster, G.L., Wilson, P.A., Storey, C.D., Trueman, C.N., Raymo, M.E., 2013. An alternative suggestion for the Pliocene onset of major Northern Hemisphere glaciation based on the geochemical provenance of North Atlantic Ocean ice-rafted debris. Quaternary Science Reviews 75, 181-194.

Baruah, G., Clements, C.F., Guillaume, F., Ozgul, A., 2019. When do shifts in trait dynamics precede population declines? American Naturalist 193, 633-644.

Bé, A.W., Caron, D.A., Anderson, O.R., 1981. Effects of feeding frequency on life processes of the planktonic foraminifer *Globigerinoides sacculifer* in laboratory culture. Journal of the Marine Biological Association of the United Kingdom 61, 257-277.

Bell, D.A., Jung, S.J.A., Kroon, D., Lourens, L.J., Hodell, D.A., 2014. Local and regional trends in Plio-Pleistocene d18O records from benthic foraminifera. Geochemistry, Geophysics, Geosystems 15, 3304 - 3321.

Bickert, T., Curry, W.B., Wefer, G., 1997. Late Pliocene to Holocene (2.6 - 0 Ma) Western Equatorial Atlantic deep-water circulation: inferences from benthic stable isotopes. Proceedings of the Ocean Drilling Program, Scientific Results 154, 239-254.

Blomqvist, D., Pauliny, A., Larsson, M., Flodin, L.-Å., 2010. Trapped in the extinction vortex? Strong genetic effects in a declining vertebrate population. Evolutionary Biology 10, 1-9.

Boakes, E.H., Isaac, N.J.B., Fuller, R.A., Mace, G.M., McGowan, P.J.K., 2018. Examining the relationship between local extinction risk and position in range. Conservation biology 32, 229-239.

Bolton, C.T., Bailey, I., Friedrich, O., Tachikawa, K., de Garidel‐Thoron, T., Vidal, L., Sonzogni, C., Marino, G., Rohling, E.J., Robinson, M.M., 2018. North Atlantic Midlatitude Surface‐Circulation Changes Through the Plio‐Pleistocene Intensification of Northern Hemisphere Glaciation. Paleoceanography and Paleoclimatology 33, 1186-1205.

Bolton, C.T., Lawrence, K.T., Gibbs, S.J., Wilson, P.A., Herbert, T.D., 2011. Biotic and geochemical evidence for a global latitudinal shift in ocean biogeochemistry and export productivity during the late Pliocene. Earth and Planetary Science Letters 308, 200-210.

Bolton, C.T., Wilson, P.A., Bailey, I., Friedrich, O., Beer, C.J., Becker, J., Baranwal, S., Schiebel, R., 2010. Millennial-scale climate variability in the subpolar North Atlantic Ocean during the late Pliocene. Paleoceanography 25.

Brombacher, A., Elder, L.E., Hull, P.M., Wilson, P.A., Ezard, T.H.G., 2018a. Calibration of test diameter and area as proxies for body size in the planktonic foraminifer *Globoconella puncticulata*. Journal of Foraminiferal Research 48, 241-245.

Brombacher, A., Wilson, P.A., Bailey, I., Ezard, T.H., 2017a. The breakdown of static and evolutionary allometries during climatic upheaval. The American Naturalist 190, 350-362.

Brombacher, A., Wilson, P.A., Bailey, I., Ezard, T.H., 2018b. Temperature is a poor proxy for synergistic climate forcing of plankton evolution. Proceedings of the Royal Society B: Biological Sciences 285, 20180665.

Brombacher, A., Wilson, P.A., Ezard, T.H., 2017b. Calibration of the repeatability of foraminiferal test size and shape measures with recommendations for future use. Marine Micropaleontology 133, 21-27.

Channell, J.E.T., Kanamatsu, T., Sato, T., Stein, R., Alvarez Zarikian, C.A., Malone, M.J., Expedition 303/306 Scientists, 2006. Site U1313. Proceedings of the Integrated Ocean Drilling Program 303/306.

Channell, R., Lomolino, M.V., 2000. Trajectories to extinction: spatal dynamics of the contraction of geographical ranges. Journal of Biogeography 27, 169-179.

Chapman, M.R., Funnell, B.M., Weaver, P.P.E., 1998. Isolation, extinction and migration within Late Pliocene populations of the planktonic foraminiferal lineage *Globorotalia* (*Globoconella*) in the North Atlantic. Marine Micropaleontology 33, 203-222.

Clements, C.F., Ozgul, A., 2016. Including trait-based early warning signals helps predict population collapse. Nature Communications 7, 1-8.

Curry, W.B., Shackleton, N.J., Richter, C., Expedition 154 Scientists, 1995. Site 925. Proceedings of the Ocean Drilling Program, Initial Reports 154.

Deshayes, G.P., 1832. Histoire Naturelle des Vers, Tome Deuxième. Agasse Imprimeur, Paris.

Du, Y., Chiari, M., Karádi, V., Nicora, A., Onoue, T., Pálfy, J., Roghi, G., Tomimatsu, Y., Rigo, M., 2020. The asynchronous disappearance of conodonts: New constraints from Triassic-Jurassic boundary sections in the Tethys and Panthalassa. Earth-Science Reviews 203, 1-18.

Falzoni, F., Petrizzo, M.R., Caron, M., Leckie, M.R., Elderbak, K., 2018. Age and synchronicity of planktonic foraminiferal bioevents across the Cenomanian-Turonian boundary interval (Late Cretaceous). Newsletters on Stratigraphy 51, 343-380.

Franz, S.-O., Tiedemann, R., 2002. Pliozäne Zeitreihen zur Rekonstruktion der Tiefenwasserzirkulation und der siliziklastischen Amazonasfracht im äquatorialen Westatlantik (Ceara Schwelle, ODP Leg 154). GEOMAR Report 84, 183.

Friedrich, O., Wilson, P.A., Bolton, C.T., Beer, C.J., Schiebel, R., 2013. Late Pliocene to early Pleistocene changes in the North Atlantic Current and suborbital-scale sea-surface temperature variability. Paleoceanography 28, 274-282.

García Joral, F., Baeza-Carratalá, J.F., Goy, A., 2018. Changes in brachiopod body size prior to the Early Toarcian (Jurassic) Mass Extinction. Palaeogeography, Palaeoclimatology, Palaeoecology 506, 242-249.

Gibbs, S.J., Young, J.R., Bralower, T.J., Shackleton, N.J., 2005. Nannofossil evolutionary events in the mid-Pliocene: an assessment of the degree of synchrony in the extinctions of *Reticulofenestra pseudoumbilicus* and *Sphenolithus abies*. Palaeogeography, Palaeoclimatology, Palaeoecology 217, 155-172.

Gilpin, M.E., 1987. Spatial structure and population vulnerability, in: Soulé, M.E. (Ed.), Viable populations for conservation. Cambridge University Press, Cambridge, UK.

Gilpin, M.E., Soulé, M.E., 1986. Minimum viable populations: processes of species extinction, in: Soulé, M.E. (Ed.), Conservation biology: the science of scarcity and diversity. Sinauer Associates, Inc., Sunderland, Massachussets, pp. 19-34.

Guthrie, R.D., 2003. Rapid body size decline in Alaskan Pleistocene horses before extinction. Nature 426, 166-169.

Hedrick, P.W., Lacy, R.C., Allendorf, F.W., Soulé, M.E., 1996. Directions in conservation biology: comments on Caughly. Conservation biology 10, 1312-1320.

Hemleben, C., Spindler, M., Anderson, O.R., 1989. Modern planktic formaminifera. Springer, New York.

Herbert, T.D., Peterson, L.C., Lawrence, K.T., Liu, Z., 2010. Tropical ocean temperatures over the past 3.5 million years. Science 328, 1530-1534.

Huber, B.T., Petrizzo, M.R., Watkins, D.K., Haynes, S.J., MacLeod, K.G., 2017. Correlation of Turonian continental margin and deep-sea sequences in the subtropical Indian Ocean sediments by integrated planktonic foraminiferal and calcareous nannofossil biostratigraphy. Newsletters on Stratigraphy 50, 141-185.

Jansen, E., Raymo, M.E., Blum, P., Expedition 162 Scientists, 1996. Sites 980/981. Proceedings of the Ocean Drilling Program, Initial Reports 162.

Kawagata, S., Hayward, B.W., Grenfell, H.R., Sabaa, A., 2005. Mid-Pleistocene extinction of deep-sea foraminifera in the North Atlantic Gateway (ODP sites 980 and 982). Palaeogeography, Palaeoclimatology, Palaeoecology 221, 267-291.

Kennett, J.P., Srinivasan, M.S., 1983. Neogene planktonic foraminifera. A phylogenetic atlas. Hutchinson Ross Publishing Company, Stroudsburg, Pennsylvania.

Kiessling, W., Schobben, M., Ghaderi, A., Hairapetian, V., Leda, L., Korn, D., 2018. Pre–mass extinction decline of latest Permian ammonoids. Geology 46, 283-286.

Lang, D.C., Bailey, I., Wilson, P.A., Beer, C.J., Bolton, C.T., Friedrich, O., Newsam, C., Spencer, M.R., Gutjahr, M., Foster, G.L., Cooper, M.J., Milton, J.A., 2014. The transition on North America from the warm humid Pliocene to the glaciated Quaternary traced by eolian dust deposition at a benchmark North Atlantic Ocean drill site. Quaternary Science Reviews 93, 125-141.

Lawrence, K.T., Herbert, T.D., Brown, C.M., Raymo, M.E., Haywood, A.M., 2009. High-amplitude variations in North Atlantic sea surface temperature during the early Pliocene warm period. Paleoceanography 24.

Levin, D.A., 2019. Intraspecific lineages as focal points in the extinction and persistence of species. Plant Systematics and Evolution 305, 719-726.

Lisiecki, L.E., Raymo, M.E., 2005. A Pliocene-Pleistocene stack of 57 globally distributed benthic δ18O records. Paleoceanography 20.

Lombard, F., Labeyrie, L., Michel, E., Bopp, L., Cortijo, E., Retailleau, S., Howa, H., Jorissen, F., 2011. Modelling planktic foraminifer growth and distribution using an ecophysiological multi-species approach. Biogeosciences 8, 853-873.

Lombard, F., Labeyrie, L., Michel, E., Spero, H.J., Lea, D.W., 2009. Modelling the temperature dependent growth rates of planktic foraminifera. Marine Micropaleontology 70, 1-7.

Lomolino, M.V., Channell, R., 1995. Splendid isolation: patterns of geographical range collapse in endangered mammals. Journal of Mammology 76, 335-347.

Martínez-Botí, M., Foster, G., Chalk, T., Rohling, E., Sexton, P., Lunt, D., Pancost, R., Badger, M., Schmidt, D., 2015. Plio-Pleistocene climate sensitivity evaluated using high-resolution CO2 records. Nature 518, 49-54.

Mc Intyre, K., Ravelo, A., Delaney, M., 1999. North Atlantic intermediate waters in the late Pliocene to early Pleistocene. Paleoceanography 14, 324-335.

Naafs, B.D.A., Hefter, J., Acton, G., Haug, G.H., Martínez-Garcia, A., Pancost, R., Stein, R., 2012. Strengthening of North American dust sources during the late Pliocene (2.7 Ma). Earth and Planetary Science Letters 317-318, 8-19.

Naafs, B.D.A., Stein, R., Hefter, J., Khélifi, N., De Schepper, S., Haug, G.H., 2010. Late Pliocene changes in the North Atlantic Current. Earth and Planetary Science Letters 298, 434-442.

Piazza, V., Duarte, L.V., Renaudie, J., Aberhan, M., 2019. Reductions in body size of benthic macroinvertebrates as a precursor of the early Toarcian (Early Jurassic) extinction event in the Lusitanian Basin, Portugal. Paleobiology 45, 296-316.

Pinheiro, J., Bates, D., 2000. Linear mixed-effect models: basic concepts and examples, Mixed-effect models in S and S-Plus. Springer-Verlag, New York, pp. 3-56.

Pinheiro, J., Bates, D., DebRoy, S., Sarkar, D., Team, R.C., 2021. nlme: Linear and Nonlinear Mixed Effects Models, 3.1.152 ed, CRAN.

Prell, W., Damuth, J., 1978. The climate-related diachronous disappearance of *Pulleniatina obliquiloculata* in Lte Quaternary sediments of the Atlantic and Caribbean. Marine Micropaleontology 3, 267-277.

Purvis, A., Gittleman, J.L., Cowlishaw, G., Mace, G.M., 2000. Predicting extinction risk in declining species. Proceedings of the Royal Society B: Biological Sciences 267, 1947-1952.

R Core Team, 2013. R: A language and environment for statistical computing, R Foundation for Statiastical Computing, Vienna, Austria.

Rando, J.C., Alcover, J.A., Michaux, J., Hutterer, R., Navarro, J.F., 2011. Late-Holocene asynchronous extinction of endemic mammals on the eastern Canary Islands. The Holocene 22, 801-808.

Raup, D.M., 1981. Extinction: bad genes or bad luck? Acta Geològica Hispànica: Concept and method in Paleontology 16, 25-33.

Rubidge, E.M., Patton, J.L., Lim, M., Burton, A.C., Brashares, J.S., Moritz, C., 2012. Climate-induced range contraction drives genetic erosion in an alpine mammal. Nature Climate Change 2, 285-288.

Ruddiman, W.F., Kidd, R.B., Thomas, E., Expedition 94 Scientists, 1987. Site 606. Initial Reports of the Deep Sea Drilling Project 94.

Ruddiman, W.F., Sarnthein, M., Baldauf, J.G., Expedition 108 Scientists, 1988. Site 659. Proceedings of the Ocean Drilling Program, Initial Reports 108.

Saupe, E.E., Qiao, H., Hendricks, J.R., Portell, R.W., Hunter, S.J., Soberón, J., Lieberman, B.S., 2015. Niche breadth and geographic range size as determinants of species survival on geological time scales. Global Ecology and Biogeography 24, 1159-1169.

Scott, G.H., Kennett, J.P., Wilson, K.J., Hayward, B.W., 2007. *Globorotalia puncticulata*: Population divergence, dispersal and extinction related to Pliocene–Quaternary water masses. Marine Micropaleontology 62, 235-253.

Shackleton, N.J., Backman, J., Zimmerman, H., Kent, D.V., Hall, M.A., Roberts, D.G., Schnitker, D., Baldauf, J.G., Desprairies, A., Horighausen, R., Huddlestun, P., Keene, J.B., Kaltenback, A.J., Krumsiek, K.A.O., Morton, A.C., Murray, J.W., Westberg-Smith, J., 1984. Oxygen isotope calibration of the onset of ice-rafting and history of glaciation in the North Atlantic region. Nature 307, 620-623.

Shakun, J.D., Raymo, M.E., Lea, D.W., 2016. An early Pleistocene Mg/Ca-δ18O record from the Gulf of Mexico: Evaluating ice sheet size and pacing in the 41-kyr world. Paleoceanography 31, 1011-1027.

Sikes, E.L., Keigwin, L.D., Curry, W.B., 1991. Pliocene paleoceanography: circulation and oceanographic changes associated with the 2.4 Ma glacial event. Paleoceanography 6, 245 - 257.

Spray, J.F., Bohaty, S.M., Davies, A., Bailey, I., Romans, B.W., Cooper, M.J., Milton, J.A., Wilson, P.A., 2019. North Atlantic evidence for a unipolar icehouse climate state at the Eocene‐Oligocene Transition. Paleoceanography and Paleoclimatology 34, 1124-1138.

Steadman, D.W., Martin, P.S., MacPhee, R.D., Jull, A.J., McDonald, H.G., Woods, C.A., Iturralde-Vinent, M., Hodgins, G.W., 2005. Asynchronous extinction of late Quaternary sloths on continents and islands. PNAS 102, 11763-11768.

Stewart, J.R., 2009. The evolutionary consequence of the individualistic response to climate change. Journal of Evolutionary Biology 22, 2363-2375.

Stewart, J.R., Lister, A.M., Barnes, I., Dalén, L., 2010. Refugia revisited: individualistic responses of species in space and time. Proceedings of the Royal Society B: Biological Sciences 277, 661-671.

Takagi, H., Kimoto, K., Fujiki, T., Moriya, K., 2018. Effect of nutritional condition on photosymbiotic consortium of cultured *Globigerinoides sacculifer* (Rhizaria, Foraminifera). Symbiosis 76, 25-39.

Tiedemann, R., Sarnthein, M., Shackleton, N.J., 1994. Astronomic timescale for the Pliocene Atlantic δ18O and dust flux records of Ocean Drilling Program site 659. Paleoceanography 9, 619 - 638.

Wade, B.S., Olsson, R.K., 2009. Investigation of pre-extinction dwarfing in Cenozoic planktonic foraminifera. Palaeogeography, Palaeoclimatology, Palaeoecology 284, 39-46.

Wade, B.S., Pearson, P.N., Berggren, W.A., Pälike, H., 2011. Review and revision of Cenozoic tropical planktonic foraminiferal biostratigraphy and calibration to the geomagnetic polarity and astronomical time scale. Earth-Science Reviews 104, 111-142.

Wei, K.-Y., 1994. Stratophenetic tracing of phylogeny using SIMCA pattern recognition technique: a case study of the Late Neogene planktic foraminifera *Globoconella* clade. Paleobiology 20, 52-65.

Westerhold, T., Marwan, N., Drury, A.J., Liebrand, D., Agnini, C., Anagnostou, E., Barnet, J.S.K., Bohaty, S.M., De Vleeschouwer, D., Florindo, F., Frederichs, T., Hodell, D.A., Holbourn, A., Kroon, D., Lauretano, V., Littler, K., Lourens, L.J., Lyle, M., Pälike, H., Röhl, U., Tian, J., Wilkens, R.H., Wilson, P.A., Zachos, J.C., 2020. An astronomically dated record of Earth’s climate and its predictability over the last 66 million years. Science 369, 1383-1387.

Wood, S.N., 2006. Generalized Additive Models: An Introduction with R. Chapman and Hall/CRC.

Wu, H., He, W., Weldon, E.A., 2018. Prelude of benthic community collapse during the end-Permian mass extinction in siliciclastic offshore sub-basin: Brachiopod evidence from South China. Global and Planetary Change 163, 158-170.

Zachos, J., Kroon, D., Blum, P., Expedition 208 Scientists, 2004. Site 1264. Proceedings of the Ocean Drilling Program, Initial Reports 208.

Zhang, Y., Shi, G.R., He, W.-h., Wu, H.-t., Lei, Y., Zhang, K.-x., Du, C.-c., Yang, T.-l., Yue, M.-l., Xiao, Y.-f., 2016. Significant pre-mass extinction animal body-size changes: Evidences from the Permian–Triassic boundary brachiopod faunas of South China. Palaeogeography, Palaeoclimatology, Palaeoecology 448, 85-95.

Zhang, Y., Shi, G.R., Wu, H.-t., Yang, T.-l., He, W.-h., Yuan, A.-h., Lei, Y., 2017. Community replacement, ecological shift and early warning signals prior to the end-Permian mass extinction: A case study from a nearshore clastic-shelf section in South China. Palaeogeography, Palaeoclimatology, Palaeoecology 487, 118-135.



Figure 1 Geographic range contraction according to the demographic (left) and contagion hypothesis (right). Light areas indicate lost range area and dark areas represent the remaining range. Red arrows show the direction of range contraction.



Figure 2 Study site locations and SEM of study species *Globoconella puncticulata* (a) and study interval (b, c). Oxygen isotopes (black) from the Lisiecki and Raymo global benthic stack (Lisiecki and Raymo, 2005) with key Marine Isotope Stages (MIS) labeled, and ice rafted debris (IRD, orange) supplied to the open ocean by icebergs calved from ice sheets large enough to reach the coastlines, at Site U1313 (Bolton et al., 2010; Lang et al., 2014) in our study interval (b) and oxygen isotopes in the past 5 million years (Lisiecki and Raymo, 2005) with the temporal range of the study species (red) (c).



Figure 3 Abundance (a-d) and box and whisker plots of shell size (e-h) of *Globoconella puncticulata* in the 600,000 years leading up to its extinction at six Atlantic Ocean sites. Boxes (thick lines) in panels e-h represent the interquartile range (IQR), the whiskers (thin lines) include all data within 1.5\*IQR and any datapoints outside the 1.5\*IQR are represented by open circles. Diamonds represent the median of each sample. Size data of Site U1313 from Brombacher et al. (2017a) all other data, this study.



Figure 4 Population abundance and timing of their extinction in the Atlantic Ocean. The global benthic oxygen isotope stack (Lisiecki and Raymo, 2005) is provided for reference with major Marine Isotope Stages (MIS) indicated in bold. The extinctions of edge sites 981 and 925 resemble the demographic hypothesis, whereas core sites U1313, 606, 1264 and 659 more closely follow the contagion hypothesis. See Supplementary Figure S6 for a spatial representation of these data.



Figure 5 Shell size and abundance of *Globoconella puncticulata* populations at Sites U1313 (Brombacher et al., 2017a) and 981 (this study) with records of local export productivity (C37 mass accumulation rates (MAR)) (Bolton et al., 2011; Naafs et al., 2010) and sea surface temperature (Lawrence et al., 2009; Naafs et al., 2010), and global CO­2  (Martínez-Botí et al., 2015). LR04 (Lisiecki and Raymo, 2005) is provided for reference with key Marine Isotope Stages indicated at the bottom and was not included in the Linear Mixed Effect models. Size is plotted as box-and-whiskers plots, with boxes (thick lines) representing the interquartile range (IQR), whiskers (thin dashed lines) including all data within 1.5 \* Interquartile Range (IQR) and open circles indicating any datapoints outside the 1.5\*IQR range. Diamonds represent the median of each sample.