

Maastrichtian-Rupelian paleoclimates in the southwest Pacific – a critical re-evaluation of biomarker paleothermometry and dinoflagellate cyst paleoecology at Ocean Drilling Program Site 1172

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Abstract. Sea surface temperature (SST) reconstructions based on isoprenoid glycerol dialkyl glycerol tetraether (isoGDGT) distributions from the Eocene southwest (SW) Pacific Ocean are unequivocally warmer than can be reconciled with state-of-the-art fully coupled climate models. However, the SST signal preserved in sedimentary archives can be affected by contributions of additional isoGDGT sources. Methods now exist to identify and possibly correct for overprinting effects on the isoGDGT distribution in marine sediments. We here use the current proxy insights to (re-)assess the reliability of the isoGDGT-based SST signal in 69 newly analysed and 242 re-analysed sediments ODP Site 1172 (East Tasman Plateau, Australia) following state-of-the-art chromatographic techniques. We compare our results with paleo-environmental and paleoclimatologic reconstructions based on dinoflagellate cysts. The resulting ~130 kyr-resolution Maastrichtian-Oligocene TEX₈₆-based SST record confirms previous conclusions of anomalous warmth in the early Eocene sw Pacific and remarkably cool conditions during the mid-Paleocene. Dinocyst diversity and assemblages show a strong response to the local SST evolution, supporting the robustness of the TEX₈₆ record. Soil-derived branched GDGTs stored in the same sediments are used to reconstruct mean annual air temperature (MAAT) of the nearby land using the MBT'_{5me} proxy. MAAT is consistently lower than SST during the early Eocene, independent of the calibration chosen. General trends in SST and MAAT are similar, except for 1) an enigmatic absence of MAAT rise during the Paleocene-Eocene Thermal Maximum and Middle Eocene Climatic Optimum, and 2) a subdued middle-late Eocene MAAT cooling relative to SST. Both dinocysts and GDGT signals suggest a mid-shelf depositional environment with strong river-runoff during the Paleocene-early Eocene, progressively becoming more marine thereafter. This trend reflects gradual subsidence and more pronounced wet/dry seasons in the northward drifting Australian hinterland, which may also explain the subdued middle Eocene MAAT cooling relative to that of SST. The overall correlation between dinocyst assemblages, marine biodiversity and SST changes suggests that temperature exerted a strong influence on the surface-water ecosystem. Finally, we find support for a potential temperature control on compositional changes of branched glycerol monoalkyl glycerol tetraethers (brGMGTs) in marine

sediments. It is encouraging that a critical evaluation of the GDGT signals confirms that most of the generated data is reliable. However, this also implies that the high TEX₈₆-based SSTs for the Eocene SW Pacific, and the systematic offset between absolute TEX₈₆-based SST and MBT'_{5me}-based MAAT estimates remain without definitive explanation.

35 1 Introduction

1.1 The Paleogene Southwest Pacific Ocean

Reconstructions of deep-sea (Westerhold et al., 2020) and sea surface temperature (Bijl et al., 2009; 2013a; Hollis et al., 2012; 2019; Frieling et al., 2014; Inglis et al., 2015; O'Brien et al., 2017; Evans et al., 2018; Cramwinckel et al., 2018; O'Connor et al., 2019; Sluijs et al., 2020) have revealed overall cool climate in the Maastrichtian and Paleocene, long-term
40 warming towards the early Eocene Climatic Optimum (EECO; 53.4 – 49.2 Ma), and subsequent cooling during the middle and late Eocene (48.6 – 33.6 Ma). The EECO stands out as particularly warm, with ice-free polar regions (Hollis et al., 2009; 2012; Pross et al., 2012; Bijl et al., 2013a; Frieling et al., 2014; Hines et al., 2017). Certain southern high-latitude regions retain warm-temperate conditions into the late Eocene (Bijl et al., 2009; Liu et al., 2009; Houben et al., 2019) and, despite ample evidence for pronounced Antarctic glacial expansion across the Eocene-Oligocene transition (Salamy and Zachos,
45 1999; Bohaty et al., 2012), even into the early Oligocene (Passchier et al., 2013; 2017; Hartman et al., 2018; O'Brien et al., 2020). Variations in atmospheric CO₂ concentrations (Anagnostou et al., 2016; 2020; Foster et al., 2017) are likely the primary driver of these multi-million-year climatic trends (Cramwinckel et al., 2018). While equatorial proxy-based sea surface temperatures (SSTs) and deep-sea temperatures, assumed to reflect high-latitude SSTs, show good correspondence with numerical model simulations under Eocene boundary conditions and with varying CO₂ forcing (Cramwinckel et al.,
50 2018), proxy-based SST reconstructions of the southwest (SW) Pacific remain warmer than those from model simulations (Bijl et al., 2009; Cramwinckel et al., 2018; Hollis et al., 2019; Crouch et al., 2020; Lunt et al., 2021), despite proposed zonal heterogeneity (Douglas et al., 2014). Specifically, numerical climate models are currently unable to simulate a paleoclimate in which the annual SST difference between the equatorial Atlantic Ocean (Cramwinckel et al. 2018) and the sw Pacific Ocean (Hollis et al., 2012; Bijl et al., 2013a) is as small as the proxy data suggest. Mechanisms explaining this mismatch
55 might be found in the geographic boundary conditions and spatial resolution of the numerical models, and the limitations of the proxies to reflect absolute temperatures. The absolute SST estimates for the SW Pacific are closer to those from the equatorial regions than they are to the deep-sea (Cramwinckel et al., 2018), which is implausible given that the South Pacific is inferred to have been the dominant region of deep-water formation during the Eocene (Huber and Thomas, 2010; Thomas et al., 2003; 2014). Without this model-data mismatch resolved, it remains unclear to what extent climate models properly
60 simulate polar amplification or, alternatively, if the current proxies properly reflect high-latitude temperatures under greenhouse conditions (Lunt et al., 2012).

Proxy evidence for warmth in the Eocene derives from a suite of organic and calcite-based proxies for marine or terrestrial temperature. SST reconstructions of the SW Pacific Ocean have primarily relied on organic geochemical proxies, mainly

TEX₈₆ (Sluijs et al., 2011; Hollis et al., 2009; 2012; Bijl et al., 2009; 2013a; Cramwinckel et al., 2020; Crouch et al., 2020).
65 Although absolute temperatures have significant uncertainties, they have been broadly confirmed by robust oxygen isotope
ratios ($\delta^{18}\text{O}$) and trace element ratios (Mg/Ca) from well-preserved planktonic foraminifera from Eocene sections in New
Zealand (Hollis et al., 2009; 2012; Creech et al., 2010; Hines et al., 2017; Crouch et al., 2020). These proxies require
assumptions regarding seawater chemistry (Mg/Ca, $\delta^{18}\text{O}$ seawater, pH) that also carry significant uncertainty (e.g., Evans et
al. 2018; Kozdon et al., 2020) and these uncertainties may affect trends in multi-million year carbonate-based
70 reconstructions. The application of clumped isotope paleothermometry can reduce some of the uncertainties surrounding
carbonate-based proxies and provide additional SST constraints (Evans et al., 2018), as has been demonstrated in work on
Seymour Island (Douglas et al., 2014).

Pollen-based vegetation reconstructions from New Zealand, Tasmania and Wilkes Land (Carpenter et al., 2012; Pross et al.,
2012; Contreras et al., 2013; 2014; Huurdeman et al., 2020), present mean annual air temperature (MAAT) reconstructions
75 which confirm the warm conditions from SST proxies. Here, uncertainties on absolute temperatures are dependent on the
diversity of the pollen assemblage, and the temperature tolerances of nearest living relatives (NLRs). Uncertainties increase
when evolutionary distances to NLRs are large. In general, vegetation-based reconstructions arguably deliver the best
constraints on winter temperatures. This is because of fundamental physiological restrictions in individual tolerances of
plants (e.g., Reichgelt et al., 2018), whereas MAAT exerts much less control on the standing vegetation than seasonal
80 temperature and (seasonal) hydrological extremes. Absolute air temperature reconstructions from soil-derived lipid
biomarkers (e.g., using brGDGT-based paleothermometry; e.g., Weijers et al., 2007a; De Jonge et al., 2014a) have also
yielded MAAT reconstructions for the Eocene (Pancost et al., 2013; Bijl et al., 2013a; Huurdeman et al., 2020; Lauretano et
al., 2021). The resulting MAAT records are more in line with numerical model simulations (Lunt et al., 2021), but
considerably colder than the SSTs from the same sections. This is difficult to reconcile in a coastal climate setting, where
85 SST and adjacent air temperature should be broadly consistent.

These uncertainties in lipid-based absolute temperature reconstructions have been put forward as underlying reason for the
data-model mismatch in past climate reconstructions. This sparked major research efforts to improve understanding of proxy
functioning and potential confounding factors, ultimately to improve the accuracy of biomarker-derived temperature
estimates, and to resolve the mismatch between model simulations and proxy data.

90 **1.2 Advances in GDGT paleothermometry**

TEX₈₆ utilizes the correspondence of proportionally higher numbers of cyclopentane rings in sedimentary archaeal
membrane lipids, the isoprenoid glycerol dialkyl glycerol tetraethers (isoGDGTs), with higher SST (Schouten et al., 2002).
This relationship is attributed to a viscoelastic adaptation of the membrane of pelagic Thaumarcheota, the dominant source
organisms of isoGDGTs, to temperature (Schouten et al., 2002; 2013). For some periods in geological deep time, including
95 the Paleocene and Eocene, TEX₈₆ calibrations based on GDGTs in core top sediments need to be extrapolated above the
modern SST range ($\sim 30^\circ\text{C}$) to estimate SST. It remains unclear if the relationship between SST and TEX₈₆ is linear or

exponential beyond the high-end of the core-top calibration (e.g., Cramwinckel et al., 2018; Hollis et al., 2019). It is therefore recommended that both linear and exponential calibrations are presented, which typically lead to higher and lower SST estimates, respectively (Hollis et al., 2019). Yet, as the absolute TEX₈₆ values of many Eocene sediments exceed those
100 observed for modern core-tops, even the most conservative calibrations yield SSTs >30°C in the warmest intervals. Increasingly, such temperatures are corroborated by estimates from other SST proxies (e.g., Bijl et al., 2010; Douglas et al., 2014; Hines et al., 2017; Crouch et al., 2020).

Along with calibration uncertainties, a number of confounding factors have been identified since the first publication of the isoGDGT-based SST records from the SW Pacific. These are related to a suite of pre-, syn-, and post-depositional processes
105 that might alter the pelagic isoGDGT signal in marine sediments. In addition to the pelagic contribution, isoGDGT assemblages may include contributions from methanogenic (Blaga et al., 2009) and methanotrophic (Weijers et al., 2011; Zhang et al., 2011) archaea, deep-dwelling archaea (Taylor et al., 2013) and terrestrial sources (Hopmans et al., 2004; Weijers et al., 2006). Various approaches have been developed to recognize these contributions, leading to more reliable SST reconstructions. However, the influence of growth phase (Elling et al., 2014), and environmental ammonium and
110 oxygen concentrations (Qin et al., 2015; Hurley et al., 2016) on sedimentary isoGDGT distributions are still poorly constrained.

Branched GDGTs (brGDGTs) produced by soil bacteria can be used to reconstruct mean annual air temperature (MAAT). The MBT'_{5me} index (De Jonge et al., 2014a; Naafs et al., 2019; Dearing Crampton-Flood et al., 2020) represents the degree of methylation of the 5-methyl brGDGTs which strongly responds to MAAT. Studies on brGDGTs in marine sediments
115 reveal that the MAAT evolution of Australian-New Zealand land masses followed the trends in deep- and surface ocean temperature remarkably well, but with lower absolute values than SST (Bijl et al., 2013a; Pancost et al., 2013). With improved analytical techniques (Hopmans et al., 2016), brGDGT isomers with a methylation on the 5- or the 6-position of the alkyl chain can now be separated and quantified (De Jonge et al., 2013). With this separation, the pH co-dependence of the brGDGT signal can be removed by only considering 5-methyl brGDGTs that only respond to temperature (De Jonge et
120 al., 2014a; Naafs et al., 2017a). The separation of brGDGT isomers also allows the recognition and correction of potential contributions of aquatic brGDGTs to the soil-derived brGDGT signal stored in marine sediments that complicate the use of brGDGTs as continental paleothermometer in continental margin sediments (Peterse et al., 2009; Tierney and Russell, 2009; Zell et al., 2013; De Jonge et al., 2014b; 2015; Sinninghe Damsté, 2016; Dearing Crampton-Flood et al., 2018).

Lipids from a related biomarker family, the branched glycerol monoalkyl glycerol tetraethers (brGMGTs), were identified in
125 the marine realm in core-top sediments (Liu et al., 2012), oxygen minimum zones (Xie et al. 2014), and Paleogene sediments (Sluijs et al., 2020). In the terrestrial realm, they were described in peats (Naafs et al., 2018a) and East African lake sediments (Baxter et al 2019), the latter identifying 7 individual brGMGTs. The brGMGTs contain a covalent bond connecting the two alkyl chains, which is thought to enhance membrane stability at higher temperatures (Morii et al., 1998). The abundance of brGMGTs relative to that of brGDGTs, as well as the relative distribution of brGMGT isomers seems to
130 vary with temperature in both the terrestrial (Naafs et al., 2018a; Baxter et al., 2019; Tang et al., 2021) and marine realms

(Sluijs et al., 2020), although this is thus far only based on empirical relationships. The exact sources of these compounds in sedimentary archives and their temperature relation are as yet not fully understood. For example, brGMTs in Paleogene Arctic Ocean sediments seem to be produced in a marine system with a strong oxygen minimum zone and substantial input of penecontemporaneous terrestrial matter (Sluijs et al., 2020). Similar to the temperature response of 5-methyl brGDGTs, the degree of methylation of acyclic brGMTs (HMBT_{acyclic}) follows the TEX₈₆ in the Paleogene Arctic record. However, the index brGMTI, which was empirically calibrated to temperature using a suite of East African lake sediments (Baxter et al., 2019) does not (Sluijs et al., 2020).

Fig. 1

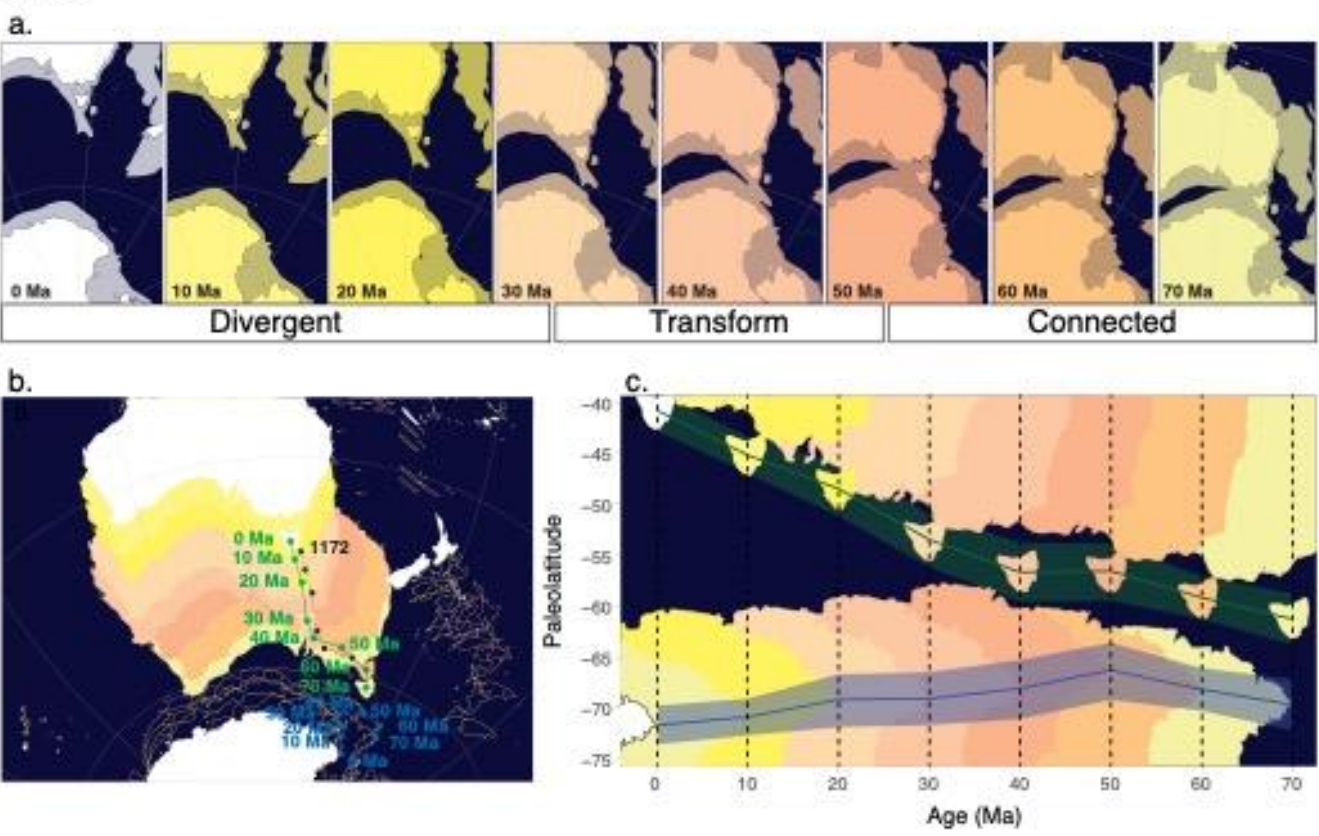


Figure 1. Absolute plate tectonic changes around Tasmania during the Maastrichtian to recent. a. Tasmanian Gateway (TG) opening in 8 time slices, present to 70 Ma. Relative tectonic motion between Australia and Antarctica is limited until about 53 Ma, transform until 34 Ma and divergent from 34 Ma onwards. b. Like a, but on an orthographic projection (compiled with Gplates, using paleomagnetic reference frame (Torsvik et al., 2012) from Seton et al. (2012)). Green and blue lines and dots represent the pathways of Tasmania and Cape Adare, respectively, black dashed line and dots represent the paleo-position of Site 1172. c. The absolute paleolatitude of Cape Adare, Antarctica (blue line, blue shading = uncertainty), as indicative of the plate tectonic motion around the pathway of the Tasman Current, and of Tasmania (green line, green shading = uncertainty), as indicative of the plate tectonic motion of the source area of the terrestrial organic matter, in 10 Myr time steps from 70–0 Ma. obtained from paleolatitude.org (Van Hinsbergen et al., 2015). Plate contours represent paleolatitude of present-day shorelines, for orientation (obtained from Gplates). Note that in visualizations b and c, submerged continental crust is not shown, but does limit TG opening.

1.3 Revisiting GDGTs, and investigating brGMGTs at Site 1172

150 From the new insights about the GDGT-based proxies, it is clear that assessing the reliability of previously published
GDGT-based temperature reconstructions requires a detailed review of the depositional conditions. Moreover, the presence
and proxy potential of brGMGTs the early Paleogene SW Pacific has not yet been studied. To this end, we have revisited the
sedimentary record at ODP Site 1172 on the East Tasman Plateau (ETP), which contains an almost complete succession of
late Cretaceous (Maastrichtian) to early Oligocene sediments (Brinkhuis et al., 2003; Schellenberg et al., 2004; Stickley et
155 al., 2004a; Bijl et al., 2013b; 2014). We have undertaken new analyses of isoGDGTs, brGDGTs, and brGMGTs from the
Maastrichtian and Paleocene section of the record, and re-analysed previously published sample sets using the
chromatography method with improved compound separation (Hopmans et al., 2016). We critically evaluate the biomarker
results using established indicators for pre-, syn- or post-depositional overprint of the primary sea surface and air
temperature signals. In addition, we use dinoflagellate cyst assemblages and terrestrial palynomorph abundance as recorders
160 of paleoenvironmental change on the continental shelf, for constraints on depositional, environmental and hydrological
changes, in order to aid interpretations of the GDGT proxies. We also evaluate the nature, source and possible temperature
affinity of the brGMGTs in our record. After this critical (re-)evaluation, we interpret the Maastrichtian to early Oligocene
sea surface and air temperature, and paleoenvironmental evolution of the sw Pacific region.

2 Material

165 2.1 Site locality and tectonic evolution

Since the Cenomanian, the continental complex including Australia, Tasmania, ETP and the South Tasman Rise (STR)
tectonically moved as one continental plate (Müller et al., 2000) – here referred to as Australia. True polar wander, although
relatively poorly constrained, has caused absolute motions of the spin axis relative to the paleo-positions of the plate circuit
(Torsvik et al., 2012; Van Hinsbergen et al., 2015). On a paleomagnetic reference frame, Antarctica and Australia rotated
170 northwestward by 3° of latitude and over 3° longitude between 70 and 50 Ma (Fig. 1), with only little transform
displacement between them. From 50 Ma onwards, the tectonic drift orientation of Antarctica shifted slightly more
southwards than that of Australia, causing left transform motion between notably west of Cape Adare and west Tasmania
(Fig. 1; Williams et al., 2019). Although this divergence effectively opened the Tasmanian Gateway (TG) to surface flow of
the Antarctic Counter Current close to the early-middle Eocene boundary (Bijl et al., 2013a), probably through subsidence
175 of the Antarctic continental margin and STR, rapid northward movement of Australia and southward movement of East
Antarctica did not start before about 40 Ma (Fig. 1; Cande and Stock, 2004; Seton et al., 2012). Yet, a connection
between Australian and Antarctic continental blocks persisted until 34 Ma when transform motion between STR and
Wilkes Land changed into spreading and ocean crust formation (Cande and Stock, 2004; Seton et al., 2012).

Fig. 2

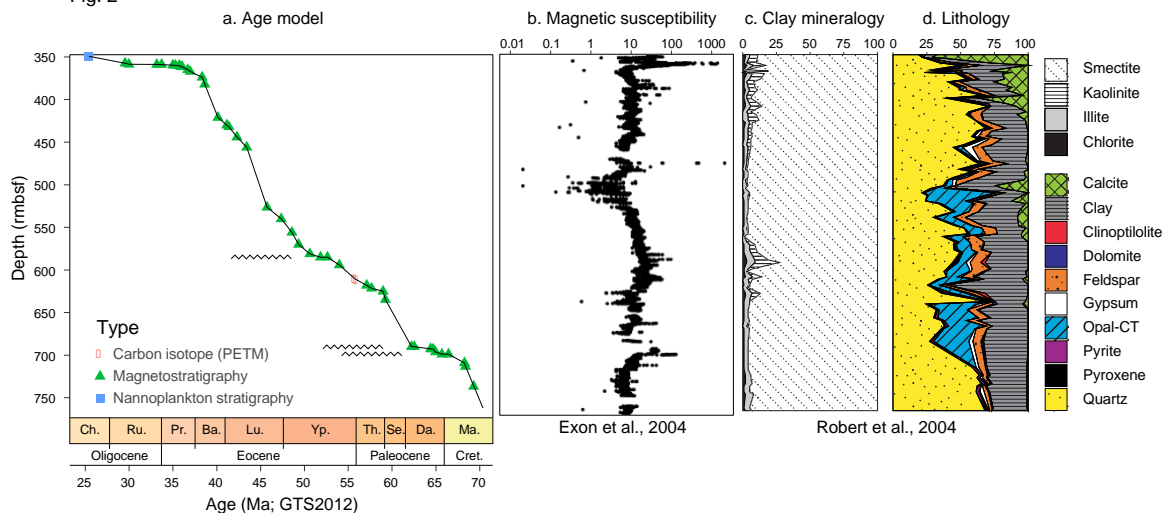


Figure 2. a. Age tie points used to construct the age model for ODP Site 1172, based on stable carbon isotopic (red dot; Sluijs et al., 2011), magnetostratigraphic (green triangles; Stickley et al., 2004a; Bijl et al., 2010; Houben et al., 2019), and nannofossil (blue square; Houben et al., 2019) age constraints. Black wobbly lines represent hiatuses. **b.** Shipboard magnetic susceptibility from Exon et al. (2001). **c.** Clay mineralogy and **d.** smear slide-based lithological observations from Robert (2004).

2.2 Lithology

A 760 meter thick sediment sequence was recovered at ODP Site 1172, on the western part of the ETP, east of Tasmania, Australia (Exon et al., 2001). We studied the interval from 760 meters below sea floor (mbsf) to about 350 mbsf in a composite (Röhl et al., 2004a) of Hole A and D. The carbonate ooze from 350 mbsf upwards did not yield any GDGTs. The studied succession consists broadly of green-grey silty claystones from 760 to ~505 mbsf, grading into dark grey to black silty claystones from 636 – 625 mbsf, then a return of the greenish grey silty claystones, which is interrupted by a more lithified unit at 610 mbsf (Exon et al., 2001; Röhl et al., 2004a, b; Schellenberg et al., 2004; Sluijs et al., 2011). The silty claystones turn gradually and cyclically lighter from 570 mbsf upwards until ~361 mbsf, with gradually more calcium carbonate and biogenic silica preserved (Röhl et al., 2004a). The clay mineralogy is dominated by smectite but contains progressively more kaolinite above ~500 mbsf (Fig. 2; Exon et al., 2001; Robert, 2004). Grey silty claystones give way to a green glauconitic horizon between 360.1 and 357.3 mbfs (Exon et al., 2001). Above this greensand, carbonate oozes continue further upwards. The record is quasi-continuously bioturbated by zoophycos (Exon et al., 2001). We follow the depth corrections published in (Sluijs et al., 2011) for Cores 12R–31R, based on the correlation of core-log and downhole log magnetic susceptibility.

2.3 Age model

For age tie points (Table S1; Fig. 2), we use the identification of the PETM (Sluijs et al., 2011) and Middle Eocene Climatic Optimum (MECO; Bijl et al., 2010) and robust biostratigraphical constraints from the Eocene–Oligocene Boundary interval

(Houben et al., 2019; Sluijs et al., 2003). In the intervals in between these tie points, the age model is based on magnetostratigraphy, which in the early Eocene suffers from a strong normal overprint (Dallanave et al., 2016). Despite the overprint, there is a paleomagnetic signal preserved in the inclination data (Fuller and Touchard, 2004). Biostratigraphic constraints from calcareous nannofossils, diatoms and dinocysts (Stickley et al., 2004a; Bijl et al., 2013b) are broadly consistent with nearby sites (Dallanave et al., 2016).

The nature of the sediments and depositional setting (see below) implies that small hiatuses may exist between the age tie points (Röhl et al., 2004a). Indeed, hiatuses were already identified across the Cretaceous–Paleogene Boundary (Schellenberg et al., 2004), in the mid-Paleocene (Bijl et al., 2013b; Hollis et al., 2014), early Eocene (Bijl et al., 2013b) and in the middle Eocene (Röhl et al., 2004a), corresponding to spikes in magnetic susceptibility (Fig. 2). The section across the Eocene–Oligocene transition is strongly condensed (Stickley et al., 2004a, b; Houben et al., 2019).

2.4 Depositional setting

From the lithology (Robert, 2004) and palynological content (Brinkhuis et al., 2003), the Maastrichtian–Eocene part of the record is interpreted to represent a shallow-marine, mid-continental shelf depositional setting, with gradual deepening in the middle Eocene based on an increase in calcium carbonate content (Fig. 2; Röhl et al., 2004a). The late Eocene greensands were initially interpreted as indicating deepening and current inception as a result of widening of the TG (Stickley et al., 2004b). Later studies have related the greensands to invigorated ocean circulation and winnowing (Houben et al., 2019), with no deepening required. Volcanic activity of the Balleny plume in the late Eocene (Hill and Moore, 2001) might have influenced bathymetric changes of the ETP, in addition to the tectonic stresses that act on diverging plates in the TG area.

The regional tectonic evolution has implications for interpreting the marine and terrestrial temperature record at this site, because it influenced regional oceanography and climatic conditions in the hinterland. Field data and model simulations indicate that with a closed TG, the Tasman Current, a strong western boundary current of the proto-Ross gyre, bathed the plateau with Antarctic-derived surface waters (Huber et al., 2004; Bijl et al., 2011; 2013b; Sijp et al., 2014; 2016). Palynological evidence confirms that the Proto-Ross Gyre influence persisted at the ETP until the late Eocene (Warnaar et al., 2009; Bijl et al., 2011). This means that despite northward tectonic drift, the same strong western boundary current bathed the site during the Maastrichtian to late Eocene (Sijp et al., 2016), with perhaps some intermittent influence of East Australian Current waters from the north (Bijl et al., 2010; Cramwinckel et al., 2020). This ended when the proto-Leeuwin Current started to flow through the progressively widening TG (Fig. 1), bringing the ETP under the influence of more northerly sourced surface waters (Stickley et al., 2004b; Houben et al., 2019).

Persistent terrigenous input (Brinkhuis et al., 2003) arguably requires a large terrestrial catchment area, and the ETP seems too small (~50,000 km²) to have had vast areas above sea level. Seismic information from the East Tasman Saddle, connecting the Tasmanian Margin to the ETP, does not suggest there was a deep basin in between (Hill and Exon, 2004). Therefore, Tasmanian-sourced detrital material could reach the ETP. Paleocene–Eocene terrestrial palynomorph assemblages

contain common Permian–Triassic elements (Contreras et al., 2014); the Permian–Triassic upper Parmeener group contains
235 thick terrestrial (coal) deposits and comprises the surface lithology of most of eastern Tasmania today. Therefore, the source
area for the terrestrial organic matter (OM) and detrital input was likely Tasmania. Minor contributions could have derived
from rivers flowing from southeast Australia. Also, the Cretaceous–Paleocene ETP might have received perhaps a minor
component of Antarctic-sourced terrestrial OM input, in addition to the dominant Tasmanian source. The regional
palynology (Truswell, 1997; Macphail, 2000; 2002; Carpenter et al., 2012; Pross et al., 2012; Contreras et al., 2013;
240 2014), the abundance of peatlands (Holdgate et al., 2009), and the felsic lithology (Moore et al., 2013) suggests the
hinterland catchment contained acidic, wet soils and peats.

2.5 Samples

For this study, we used lipid extracts that have been analysed for GDGTs previously (Bijl et al., 2009; Sluijs et al., 2011;
Hollis et al., 2014; Houben et al., 2019). We augmented these with 69 new extracts of sediments from the Maastrichtian
245 and the Paleocene, to extend and improve the temporal resolution of the record. Unfortunately, not all archived samples from
the PETM interval (Sluijs et al., 2011) could be located, and the interval was reanalysed in a lower resolution (6 samples
over the PETM interval). For the remaining PETM, we used the published peak areas (Sluijs et al., 2011) to calculate
TEX₈₆, which is warranted given the new analytical technique does not affect isoGDGT peak area ratios (Hopmans et al.,
2016). For palynology, we collated and revisited data presented in (Brinkhuis et al., 2003; Bijl et al., 2010; 2013b; Sluijs
250 et al., 2011; Houben et al., 2019) and generated higher-resolution data for the Maastrichtian and Paleocene.

3. Methods

3.1 Organic geochemistry

3.1.1 Extraction, column separation and analysis

Earlier work (Bijl et al., 2009; 2010; 2013a; Sluijs et al., 2011; Houben et al., 2019) presented in detail the extraction,
255 Al₂O₃ column separation and filtering techniques used for the samples. We followed the same procedures for the processing
of the new samples. We reanalysed all available polar fractions using the double column UHPLC-MS approach as described
in (Hopmans et al., 2016). In short, processing involved extraction with a Dionex accelerated solvent extractor using
dichloromethane:methanol (DCM:MeOH) 9:1 (v/v), column separation of the total lipid extract using solvent mixtures
hexane:DCM 9:1 (v/v), hexane:DCM 1:1 (v/v) and DCM:MeOH 1:1 (v/v) for apolar, ketone and polar fractions,
260 respectively. Polar fractions were filtered using a 0.45 µm polytetrafluorethylene filter and analysed using an Agilent 1260
Infinity series HPLC system coupled to an Agilent 6130 single-quadrupole mass spectrometer.

3.1.2 Data analysis: indices, overprints

Since the discovery of isoGDGTs as proxy for SST (see Schouten et al. (2013) for a review), several non-SST effects have been identified that may affect the distribution of isoGDGTs in the sediment. Several indices have been developed to identify most of the known sources of overprints (Table 1). As well as identifying potential overprints, these indices can provide additional paleoenvironmental information.

Table 1: Indices from iso- and brGDGTs and brGMGTs. For the chemical structure of these components see Fig. 3.

| Index name | Equation | Proxy for | Cut-off value | Source |
|-------------------------|--|---|----------------------|--------------------------|
| TEX ₈₆ | $\frac{GDGT2 + GDGT3 + Cren'}{GDGT1 + GDGT2 + GDGT3 + Cren'}$ | Sea surface temperature | - | (Schouten et al., 2002) |
| BIT index | $\frac{IIIa + IIIa' + IIa + IIa' + Ia}{Cren + IIIa + IIIa' + IIa + IIa' + Ia}$ | Terrestrial input | >0.4? Site-dependent | (Hopmans et al., 2004) |
| fcren' | $\frac{\%Cren'}{\%Cren' + \%Cren}$ | Non-thermal contribution of crenarchaeol isomer | 0.25 | (O'Brien et al., 2017) |
| Methane Index | $\frac{GDGT1 + GDGT2 + GDGT3}{GDGT1 + GDGT2 + GDGT3 + Cren + Cren'}$ | Contribution by methane-metabolising archaea | >0.3 | (Zhang et al., 2011) |
| AOM ratio | $GDGT2/Cren$ | Contribution by anaerobic methane oxydizers | >0.2 | (Weijers et al., 2011) |
| GDGT2/3 ratio | $GDGT2/GDGT3$ | Contribution by deep-dwelling archaea | >5 | (Taylor et al., 2013) |
| Methanogenesis | $GDGT0/Cren$ | Contribution by methanogenic archaea | >2.0 | (Blaga et al., 2009) |
| Ring index (RI) | $0*\%GDGT0 + 1*\%GDGT1 + 2*\%GDGT2 + 3*\%GDGT3 + 4*\%Cren + 4*\%Cren'$ | Non-pelagic GDGT composition | $\Delta RI > 0.3^*$ | (Zhang et al., 2016) |
| MBT' _{5me} | $\frac{IIIa + IIIb + IIIc}{IIIa + IIIb + IIIc + IIa + IIb + IIc + Ia}$ | Mean annual air temperature | - | (De Jonge et al., 2014a) |
| CBT' | $\frac{IIIb + IIb' + IIc' + Ia'}{IIIa + IIa + Ia}$ | (soil-)pH | - | (De Jonge et al., 2014a) |
| #rings _{Tetra} | $\frac{Ib + 2 * Ic}{Ia + Ib + Ic}$ | Marine in situ | >0.7 | (Sinninghe Damsté, 2016) |
| #rings _{Penta} | $\frac{IIb + IIb' + 2 * IIc + 2 * IIc'}{IIa + IIa' + IIb + IIb' + IIc + IIc'}$ | brGDGT production | - | |

| | | | | |
|--------------------------------------|--|---------------------------------|---------------------------|---|
| #rings _{penta5} | $\frac{IIb + 2 * IIc}{IIa + IIb + IIc}$ | | - | |
| #rings _{penta6} | $\frac{IIb' + 2 * IIc'}{IIa' + IIb' + IIc'}$ | | - | |
| IR | $\frac{IIa' + IIb' + IIc' + Ia'}{IIa + IIa' + IIb + IIb' + IIc + IIc' + Ia + Ia'}$ | River in situ brGDGT production | Depends on soil value | (De Jonge et al., 2014b) |
| IR _{hexa} | $\frac{Ia'}{Ia + Ia'}$ | | - | (Sinninghe Damsté, 2016) |
| IR _{penta} | $\frac{IIa' + IIb' + IIc'}{IIa + IIa' + IIb + IIb' + IIc + IIc'}$ | | | |
| brGMGTI | $\frac{H1020c + H1034a + H1034c}{H1020b + H1020c + H1034a + H1034b + H1034c}$ | Temperature, oxygenation? | In lakes | (Baxter et al., 2019) |
| %brGMGT, %brGMGT _a cyclic | $\frac{[brGMGT]}{[brGMGT] + [brGDGT]} \times 100$ | | In peats, lakes | (Baxter et al., 2019) |
| | $\frac{[brGMGT]}{[Ia + IIa + IIIa + brGMGT]} \times 100$ | | | (Naafs et al., 2018a) |
| HMBT _{acyclic} , all | $\frac{H1020a + H1020b + H1020c}{H1020a + H1020b + H1020c + H1034a + H1034b + H1034c}$ | | In peats | (Naafs et al., 2018a) |
| HMBT _{acyclic} | $\frac{H1020c}{H1020c + 1034b + H1048}$ | | In Arctic Ocean sediments | Sluijs et al., 2020, sensu Naafs et al., 2018a) |

* Cutoff depends on TEX₈₆ value. $\Delta RI = RI_{\text{TEX}} - RI$, where $RI_{\text{TEX}} = -0.77 * \text{TEX}_{86} + 3.32 * \text{TEX}_{86}^2 + 1.59$

All of the ratios that signal biases express the influence of this bias relative to modern “normal” or idealized compositions.

- 270 For the branched and isoprenoid tetraether (BIT) index, it was proposed that the closed sum ratio between brGDGTs, which were thought to derive exclusively from soils, and the isoGDGT crenarchaeol, which was thought to be produced exclusively by marine archaea, could be an indicator for the relative contribution of soil organic matter (OM) into marine sediments (Hopmans et al., 2004). However, it was subsequently shown that crenarchaeol, but critically also other isoGDGTs are also produced on land (Weijers et al. (2006), and that a large contribution of soil-OM to marine sediments can cause a bias in
- 275 TEX₈₆. Weijers et al. (2006) used an end-member model to calculate the potential bias in TEX₈₆-based SST reconstructions in the Congo Fan and this shows significant bias (>2°C) for BIT above 0.3. However, it is rather arbitrary to assign a cut-off for BIT based on the magnitude of the SST bias alone, as this bias depends primarily on the difference between the isoGDGT composition of the soil and marine endmembers. As a consequence, there is no uniform value for BIT index above which TEX₈₆ should be discarded. Secondly, because the BIT index is a closed sum, the index is equally affected by
- 280 increased production of crenarchaeol as it is by the input of brGDGTs. Although this could be verified with absolute concentrations of GDGTs, these are not always presented. Third, brGDGTs are produced in situ in the marine realm

(Peterse et al., 2009; Sinninghe Damsté, 2016). The corollary of this is that the cut-off value for BIT to infer an overprint in the isoGDGT-based SST signal should be assumed to differ per environmental setting, and perhaps also through time if the setting changes significantly (e.g., Sluijs et al., 2020). As a result, cut-offs have been set higher (e.g., 0.4; Bijl et al., 2013a) when no relationship between TEX₈₆ and BIT index values is detected. This approach is also not without complications, because the abundance and composition of isoGDGTs (including crenarchaeol) in soil material varies (Weijers et al., 2006; De Jonge et al., 2015; De Jonge et al., 2016; Naafs et al., 2019; Sluijs et al., 2020). This implies that for samples with intermediate to high BIT (Weijers et al., 2006), a terrestrially-derived isoGDGT contribution with a variable isoGDGT composition or a TEX₈₆-signature close to the original pelagic isoGDGT signal would become undetectable in a TEX₈₆-BIT index crossplot. In some paleo-environmental settings that evidently lack high input of soil-OM, BIT index values are high (Leutert et al., 2020). For reasons given above, this could either mean very low crenarchaeol production in the marine realm, or in situ production of brGDGTs. These two scenarios make for a completely different paleoenvironmental interpretation, and fail to indicate whether the brGDGTs can be used to reconstruct MAAT of the hinterland, or whether isoGDGTs can be used for reliable paleothermometry. For that, it is important to assess the sources of brGDGTs, as brGDGTs produced in rivers (Zell et al., 2013; De Jonge et al., 2014b) or the (coastal) marine environment (Peterse et al., 2009; Sinninghe Damsté, 2016) may contribute and thereby bias the initial soil-derived brGDGT signal, affecting the reliability of MAAT reconstructions. Fortunately, production of brGDGTs in the marine realm can be recognized based on the weighed number of rings of the tetramethylated brGDGTs, quantified in the #rings_{tetra}, where values >0.7 indicate a purely marine source of the brGDGTs (Sinninghe Damsté, 2016). The #rings_{tetra} in coastal marine sediments can subsequently be used to determine and correct for the contribution of marine brGDGTs to the temperature signal of the soil-derived brGDGTs (Dearing Crampton-Flood et al., 2018). Similarly, brGDGTs that are produced in rivers are characterized by a relatively high abundance of the 6-methyl brGDGTs relative to the 5-methyl brGDGTs, expressed as the Isomerization Ratio (IR; De Jonge et al., 2014b).

For the other indices for overprints, cut-off values may not be as uniformly applicable to all depositional settings either. Leutert et al. (2020) showed that modern samples exceeding cut-off values for the GDGT2/3 ratio and methanogenesis index do not have anomalous TEX₈₆ index values in the modern core-top dataset based on the Ring index. This suggests that the TEX₈₆ index value is not equally influenced by non-thermal contributions in all depositional settings. As a result, evaluating non-thermal contributions on TEX₈₆ index values should consider the depositional and environmental setting along with the cut-off values.

Fig. 3

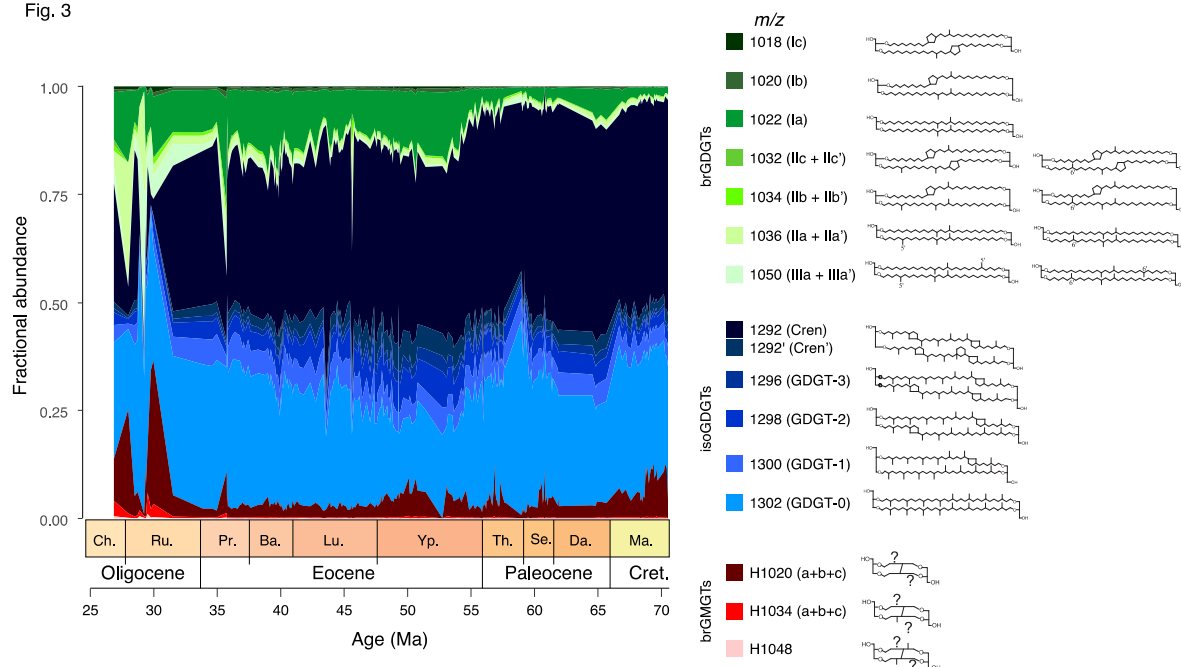


Figure 3. Variations in fractional abundances of the various branched (green) and isoprenoid (blue) GDGTs, and branched GMTs (red) at ODP Site 1172.

3.1.3 Calibrations for TEX_{86} and $\text{MBT}'_{5\text{me}}$

Several calibrations have been proposed to convert TEX_{86} index values to sea surface temperatures. For higher temperatures, it remains uncertain if the TEX_{86} -SST relationship is linear or exponential for values above the modern calibration (O'Brien et al., 2017; Tierney et al., 2017; Cramwinckel et al., 2018; Hollis et al., 2019). Following recent recommendations (Hollis et al. 2019), we here apply several calibrations to convert our TEX_{86} values to SSTs. We apply an exponential calibration (Kim et al., 2010) even though it suffers from a regression dilution bias (Tierney and Tingley, 2015), and two linear calibrations; one on the warm part ($>15^\circ\text{C}$) of the core-top dataset (O'Brien et al., 2017) and one using Bayesian statistics (BAYSPAR; Tierney and Tingley, 2015) to convert our TEX_{86} values to SSTs. We chose these calibrations to present the appropriate uncertainty related to calibration function, particularly for TEX_{86} values exceeding those of the modern core-top data. For MAAT reconstructions, we use the Bayesian BayMBT₀ of Dearing Crampton-Flood et al. (2020). We will compare the calibrations for both proxies to middle Eocene $U^{K'}_{37}$ -based SSTs (from Bijl et al., 2010, but using BAYSPLINE calibration of Tierney and Tingley, 2018) and to Paleocene–early Eocene sporomorph-based MAAT estimates (Contreras et al., 2014) from the same site.

Table 2. Calibration equations for TEX_{86} and MAAT

| Calibration | Equation | Type | Proxy for | Source |
|-------------|----------|------|-----------|--------|
|-------------|----------|------|-----------|--------|

| | | | | |
|---------------------------------|---|-----------------|---|---------------------------------------|
| SST_{exp} (TEX_{86}^H) | $68.4 * \log_{10}(TEX_{86}) + 38.6$ | Exponential | Mean annual SST (0–20m) | (Kim et al., 2010) |
| SST_{lin} | $58.8 * TEX_{86} - 11.18$ | Linear | Mean annual SST (0–20m) | (O'Brien et al., 2017) |
| BAYSPAR | TEX_{86} , Prior mean = 20, prior std = 10, search tolerance = 0.15 | Bayesian linear | Mean annual SST (0–20m) | (Tierney and Tingley, 2015) |
| BayMBT ₀ | MBT' _{5me} , Prior mean = 20, prior std = 15 | Bayesian linear | Mean annual air temperature (excluding days >0°C) | (Dearing Crampton-Flood et al., 2020) |

3.1.4 R-script for data analysis and evaluation

To facilitate systematic calculation of GDGT ratios, data analysis, visualization, and evaluation, we constructed a set of R
 330 markdowns (<https://github.com/bijlpeter83/RGDGT.git>) that can be applied to any time- or depth series of isoGDGT, brGDGT and/or brGMGT data. The R script loads peak areas of GDGTs/GMGTs from Microsoft excel spreadsheets, calculates and plots fractional abundances, overprinting indices and paleotemperature time or depth series.

3.2 Palynology

3.2.1 Sample processing

335 Palynological sample processing techniques were published in the original papers describing these datasets (Brinkhuis et al., 2003; Bijl et al., 2010; 2013b; Sluijs et al., 2011; Houben et al., 2019). We followed the same procedure for the new Maastrichtian–Paleocene samples. These publications do utilize different sieve mesh sizes (10 or 15 µm) but given all counted dinocysts are larger than 15 µm, this has not led to differences in the dinocyst results.

3.2.2 Taxonomy

340 We used the taxonomic framework cited in Williams et al., (2017) in our counts down to the species level, with one exception. For the Wetzelielloidae subfamily, we follow Bijl et al. (2017), for reasons stated therein. We follow the supra-generic classification of Fensome et al. (1993), also for genera described post-1993.

3.2.3 Ecological affinities of dinocyst eco-groups

In this paper, dinocysts are plotted in eco-groups and complexes, which group species with similar ecologic affinities based
 345 on modern (for extant taxa) and empirical (for extant and extinct taxa) data (Sluijs et al., 2005; Frieling and Sluijs, 2018). The species of which the ecologic affinity was not assessed previously were included into larger groups based on shared

morphological characteristics; primarily tabulation, archaeopyle and cyst- and process shapes. For example, the “*Apectodinium* complex” is equated to “Wetzellioids” following Frieling et al. (2014; see Table S2 for taxonomic grouping). We directly compare dinocyst- and terrestrial palynomorph indices with GDGT-based indices (following the approach of Frieling and Sluijs, 2018) to arrive at multi-proxy reconstructions of SST, river runoff and distance to shore (Table 3). For this, we resampled and binned GDGT-based indices to the sample resolution of the dinocyst data. Some dinocyst groups have affinities for multiple environmental parameters, due to the strong co-variance of environmental gradients on an inshore-to-offshore transect. Therefore, the complete dinocyst assemblage (including absence of presence) and knowledge of the geographic setting is used to reconstruct changes in paleoenvironmental conditions from dinocysts.

Table 3. Environmental parameters, and their corresponding GDGT indices and dinocyst eco-groups, based on Sluijs et al. (2005), Frieling and Sluijs (2018) and Sluijs and Brinkhuis (2009).

| Environmental parameter | GDGT index | Dinocyst eco-group |
|-------------------------------|-------------------|---|
| Sea surface temperature | TEX ₈₆ | %Thermophiles (Wetzellioideae, <i>Hafniasphaera</i> spp., <i>Florentinia reichartii</i> , <i>Polysphaeridium</i> spp., <i>Homotryblum</i> spp., <i>Heteraulacacysta</i> spp., <i>Eocladopyxis</i> spp., <i>Dinopterygium</i> spp.) |
| Runoff, fresh water, salinity | IR, BIT | % <i>Senegalinium</i> cpx, % <i>Phthanoperidinium</i> spp., %Terrestrial palynomorphs |
| Distance to shore | BIT, #ringtetra | %Open marine (<i>Apectodinium</i> spp., <i>Impagidinium</i> spp., <i>Operculodinium</i> spp., <i>Spiniferites</i> spp.), % <i>Glaphyrocysta</i> cpx., %Epicystal Goniodomids (<i>Polysphaeridium</i> spp., <i>Homotryblum</i> spp., <i>Heteraulacacysta</i> spp., <i>Eocladopyxis</i> spp., <i>Dinopterygium</i> spp.), %Terrestrial palynomorphs |

3.2.4 Diversity and variability indices

To gain insight into the diversity and variability of the dinocyst assemblage through time, we employ several indices (Table 4), and compare their results at species and eco-group level. We note that diversity in dinocyst taxonomy is complicated because dinocysts are dominantly produced during the hypnozygotic phase of the dinoflagellate life cycle following sexual reproduction. Only ~15% of modern dinoflagellates have this cyst stage (Fensome et al., 1993). This leaves a large proportion of the biological group unrepresented in the fossil record, including, for example, coral and foraminifer symbionts, but also many free-swimming taxa. Dinocyst diversity is therefore a biased indicator for plankton diversity. As a further complication, taxonomic divisions of cysts, although strongly related to dinoflagellate morphology and taxonomy, is fully based on the morphology of dinocysts. Even modern dinoflagellate-dinocyst relationships are often ambiguous. Some

dinoflagellate species produce various cyst morphologies (Rochon et al., 2008) that represent multiple cyst genera or species. Some of these cyst morphological variations are the result of ecology (Mertens et al., 2011). This complicates comparing cyst datasets on the species level and affects assessing biological diversity based on cysts. On the genus level, taxonomic division of cysts is in most cases based on plate tabulation, which is the morphologic feature that has the closest relationship to dinoflagellate biological diversity (Fensome et al., 1993). However, ecologic or biogeographic affinities have been established for some dinocysts on a species level (e.g., Frieling and Sluijs, 2018), which does suggest subtle morphological features may have biologic and ecologic significance. The eco-groups we use here are the result of extensive reviews of empirical data. These groups combine dinocyst genera and species with fundamentally similar plate tabulations, and thus probably group cysts of closely affiliated biological dinoflagellate species, and, as such also ecologic affinities. For our diversity calculations we use both the ungrouped data on species level and the diversity in dinocyst eco-groups. As the simplest approximation of biological diversity, the richness (R) sums dinocyst taxa. Furthermore, several diversity measures were calculated using the R package Vegan (Oksanen et al., 2015): Fisher's alpha (α), Shannon index (H') and Simpson index (D). Finally, we employed the Σcv metric (Gibbs et al., 2012), which we here dub the "Gibbs index", which provides a measure for assemblage variability. Together, these metrics give insight into changing stability and diversity of these regional dinocyst assemblages over the Maastrichtian to early Oligocene.

Table 4. Diversity and variability indices for (fossil) assemblages. In the Shannon and Simpson indices, p_i represents the proportional abundance of the i^{th} taxon of the total amount of taxa R. The Gibbs index summates the coefficients of variation (SD/mean) of all taxa (i to R) over a certain rolling window $t_1 \rightarrow t_2$. In Fisher's alpha, the α parameter is estimated from the dataset in which S is the expected number of species with an abundance of n . x represents a nuisance parameter estimated from the dataset, generally between 0.9 and 1. While the Gibbs index is calculated over a rolling window, the other indices are calculated per sample.

| Index | Equation | Source |
|-----------------------------|--|-----------------------|
| Shannon index (H') | $H' = \sum_{i=1}^R p_i * \ln p_i$ | (Shannon, 1948) |
| Simpson index (D) | $D = \sum_{i=1}^R p_i^2$ | (Simpson, 1949) |
| Gibbs index (Σcv) | $\Sigma cv = \sum_{i=1}^R \left(\frac{SD_{i,t_1 \rightarrow t_2}}{\text{mean}_{i,t_1 \rightarrow t_2}} \right)$ | (Gibbs et al., 2012) |
| Fisher's alpha (α) | $S_n = \frac{\alpha x^n}{n}$ | (Fisher et al., 1943) |

3.3 Comparison of GDGT and dinocyst assemblage data

Both dinocyst eco-groups and GDGT indices bear information on SST, runoff (~salinity), nutrients and marine primary productivity, and relative distance to shore (Table 4). As dinocyst and GDGT analyses were performed on partly separate datasets, the highest resolution dataset (dinocysts) was linearly resampled to the depth intervals of the GDGT data to facilitate inter-comparison.

4. Results

4.1 GDGTs

The resulting dataset has an average temporal resolution of ~130 kyrs for the time interval between 70 and 30 Ma. IsoGDGTs dominate the pool of GDGTs (particularly GDGT-0 and crenarchaeol), with a gradual increasing relative abundance of brGDGTs (particularly Ia) throughout the record (Fig. 3). BrGMGTs are in low (<10%) relative abundance, except in the Oligocene, where they account for up to 30% of the total pool of tetraether lipids.

4.1.1 Isoprenoidal GDGTs

Before the TEX₈₆ index results (Fig. 4a) can be interpreted in terms of SST, we assess whether the isoGDGTs are primarily derived from the sea surface, or have received contributions from non-pelagic sources. Across the Ypresian, the BIT index (Fig. 4b) increases from values around 0.1 in the underlying sediments to values at times exceeding 0.4. Most EECO and middle Eocene values, however, have BIT index values between 0.2 and 0.35. The remaining sediments have BIT index values of around 0.3, with the exception of some upper Eocene and Oligocene sediments that occasionally have values around 0.75. There is no correlation between TEX₈₆ and BIT index in our entire dataset ($R^2=0.0067$; $p=0.15$). None of the samples have f_{cren'} (Fig. 4c) values above the cut-off of 0.25, which excludes non-thermal contributions of the crenarchaeol isomer to the isoGDGTs. In fact, the consistent trend in f_{cren'} and the TEX₈₆ (Fig. 4) demonstrates the temperature sensitivity of cren'. With the exception of 6 samples in the Oligocene, all samples have methane index values below the conservative cut-off value of 0.3 (Fig. 4d), although in the early and middle Eocene, methane index values do approach the cut-off. AOM ratio values suggest an overprint from anaerobic methane oxidisers in one sample in the early Eocene that also has a high BIT index, and 5 samples from the Oligocene where isoGDGTs supposedly received a contribution of anaerobic methane oxidisers (Fig. 4e). As a result, these samples were not used for SST reconstruction.

Values signalling a deeper water column overprint (GDGT2/3 ratio) fall below 5 for most of the data (Fig. 4f), which was expected given the depositional setting on a continental shelf. A total of 5 samples has GDGT2/3 values exceeding 5, 4 of which also had other indications of overprinted isoGDGT signals. We discard eight TEX₈₆ datapoints because GDGT-0/crenarchaeol ratios are above 2 (Fig. 4g). One of these is from the PETM, with a value just above the cut-off. The others represent Oligocene samples.

Fig. 4

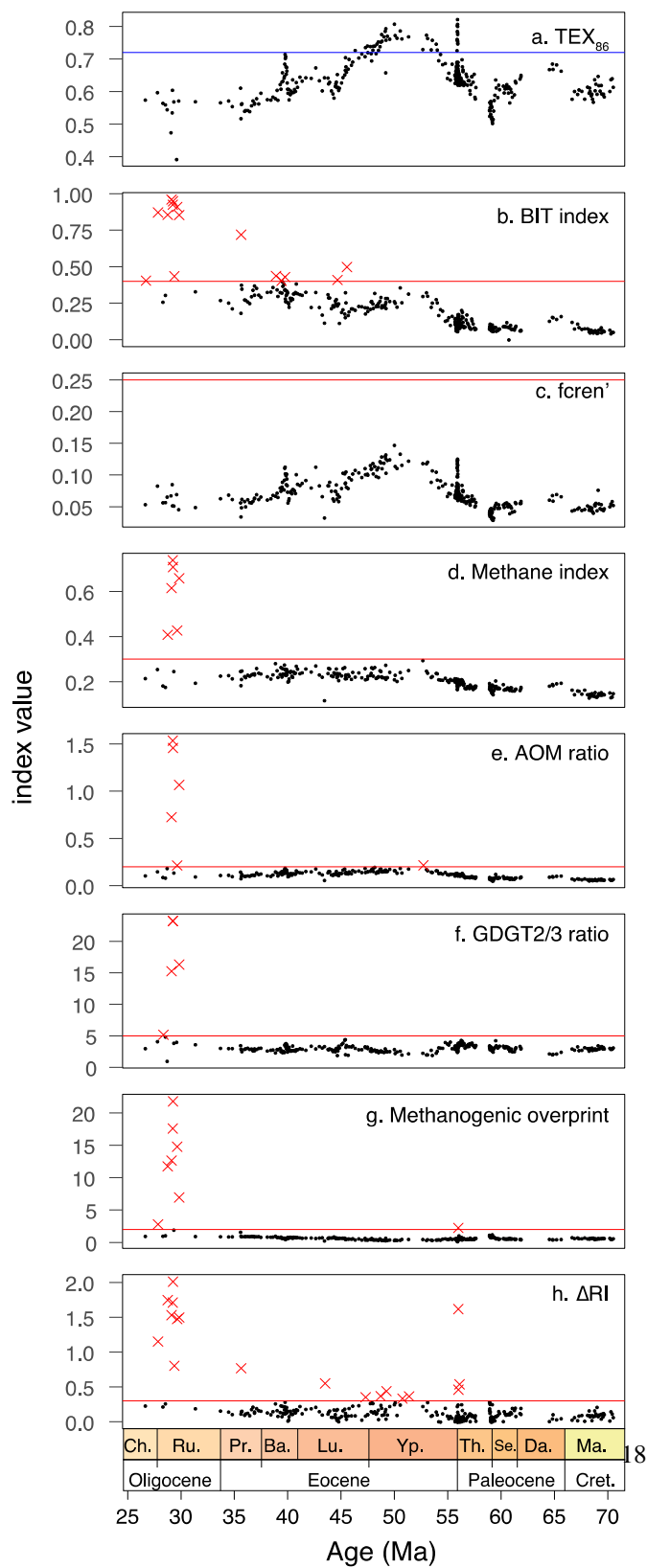


Figure 4. TEX_{86} index and indices to identify overprints. For the equations of these indices see Table 1. Red line indicates the cut-off value, red crosses indicate samples exceed the cutoff a. TEX_{86} (Schouten et al., 2002), with blue line indicating the maximum modern core-top value (~ 0.72). b. BIT index (Hopmans et al., 2004). c. feren' (O'Brien et al., 2017), no samples discarded; d. Methane index (Zhang et al., 2011), $n_{\text{discarded}} = 6$. e. AOM ratio (Weijers et al., 2011), $n_{\text{discarded}} = 6$. f. GDGT2/3 (Taylor et al., 2013), conservative cut-off of 5, $n_{\text{discarded}} = 4$, one sample retained for absence of anomalous Ring index values (see text). g. Methanogenesis (Blaga et al., 2009), cut-off of 2.0, $n_{\text{discarded}} = 8$. h. ΔRI $n_{\text{discarded}} = 18$.

In total, 18 samples fall outside the range of RI values that characterize modern pelagic-derived isoGDGT compositions ($\Delta\text{RI} > 0.3$; Fig. 4h, or 95% confidence interval; Fig. 5). Nine of these 18 come from the interval > 43 Ma and have no other indications of overprints. These samples fall just outside the 95% confidence interval limits (Fig. 5) and do follow the overall RI/ TEX_{86} trend, but we here discard them to be conservative. The other 9 samples with too high ΔRI come from the late Eocene-Oligocene and have BIT index values over 0.4. Two of these samples have no other overprint indication, the other 7 have multiple other overprints. One sample has RI values above the range of modern samples (with no other overprinting indications), the others have too low RI values. Eight samples with normal RI values do have overprints based on the other indices: BIT index values slightly over 0.4 (6 samples), GDGT2/3 ratio values over 5 (1 sample), and AOM ratio over 0.2 (1 sample). Following Leutert et al. (2020) we retained the samples with normal RI values and high BIT or GDGT2/3 ratio values. After considering all these potential biases, and retaining those with normal isoGDGT distributions, we discard a total of 19 samples. Most discarded samples are from the Oligocene and are excluded based on multiple overprint criteria.

Fig.5

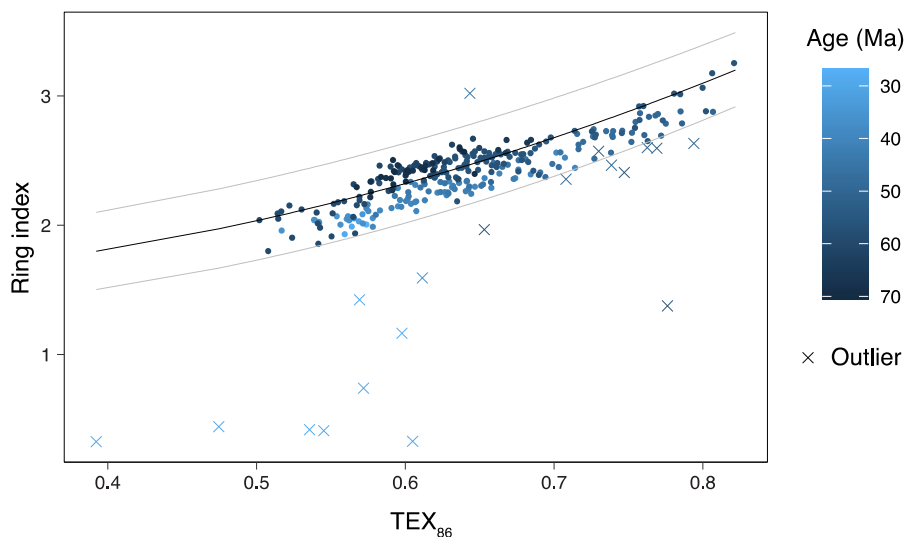
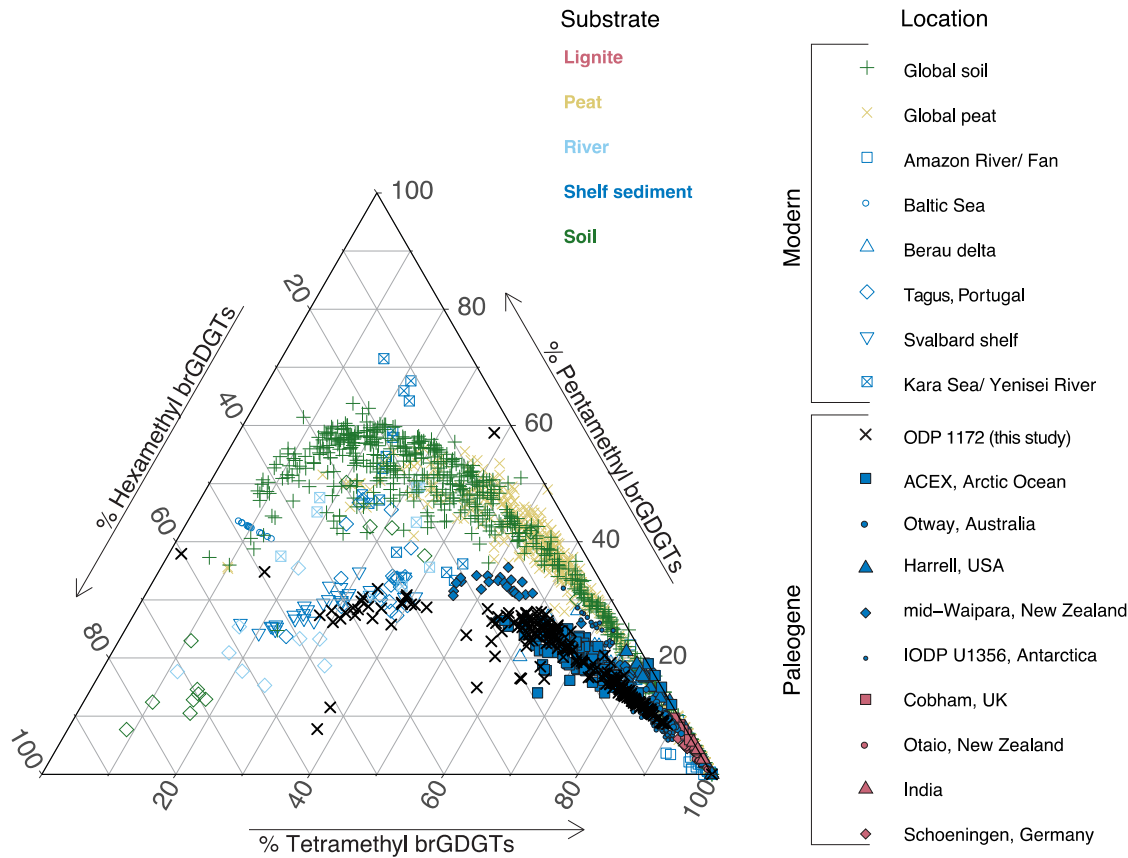


Figure 5. Ring index (RI) versus TEX_{86} . Dots indicate RI/ TEX_{86} values for samples that are retained, crosses indicate samples that are discarded (see Fig. 4). Color of all datapoints indicates age of the sample, from Maastrichtian (dark blue) to Oligocene (light blue). Black and grey curves represent the RI/ TEX_{86} ratio in modern core top samples and 95% confidence interval, respectively (see Table 1).

Fig. 6



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Figure 6. Ternary diagram showing fractional abundance of tetra-, penta- and hexamethylated brGDGTs from modern soils (Dearing Crampton-Flood et al., 2020), peats (Naafs et al., 2017b), rivers (Tagus River, Portugal (Warden et al., 2016) and Yenisei River (De Jonge et al., 2015)) and marine sediments (Kara Sea (De Jonge et al., 2015), Svalbard (Dearing Crampton-Flood et al., 2019), Berau delta (Sinninghe Damsté, 2016), Portugal (Warden et al., 2016), Baltic Sea (Warden et al., 2018), and Amazon Fan (Zell et al., 2014)), Paleogene lignites (Cobham, Schoeningen, India and Otaio (Naafs et al., 2018b)) and Paleogene marine sediments (Arctic Ocean, IODP Arctic Coring Expedition ACEX (Willard et al., 2019), Otway Basin, Australia (Hurdeman et al., 2020), Gulf of Mexico coastal plain, Harrel core (Sluijs et al., 2014), New Zealand (Pancost et al., 2013) and Antarctic Margin IODP Site U1356 (Bijl et al., 2013a)). Data from ODP Site 1172 in black crosses.

4.1.2 Branched GDGTs

450

Before brGDGT distributions can be interpreted in terms of air temperature, we assess whether they are primarily derived from soils, or have received contributions from river-, or marine in situ production. The relative abundance of tetra-, penta- and hexamethylated brGDGTs in the samples from 1172 deviates from that in modern and Paleogene soils and peats (Fig. 6): it has either comparatively less tetra-, less penta-, or more hexamethylated brGDGTs, and the data split into 2 clusters. The largest cluster follows the distribution observed in modern soils and peats with an offset (Fig. 6).

Fig.7

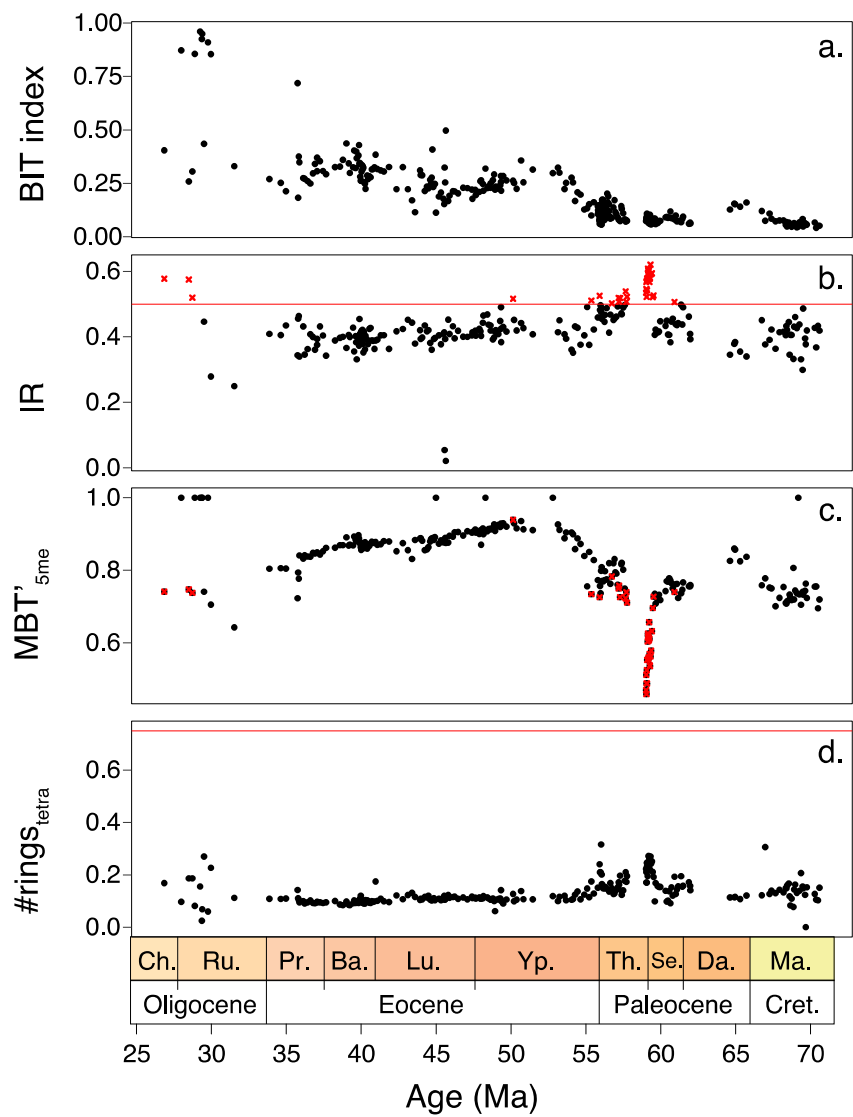


Figure 7. a. BIT index, b. IR (with cut-off value of the Australian soil IR of 0.5 and discarded MAAT samples in red crosses), c. MBT'5me and d. #rings_{tetra} from Site 1172.

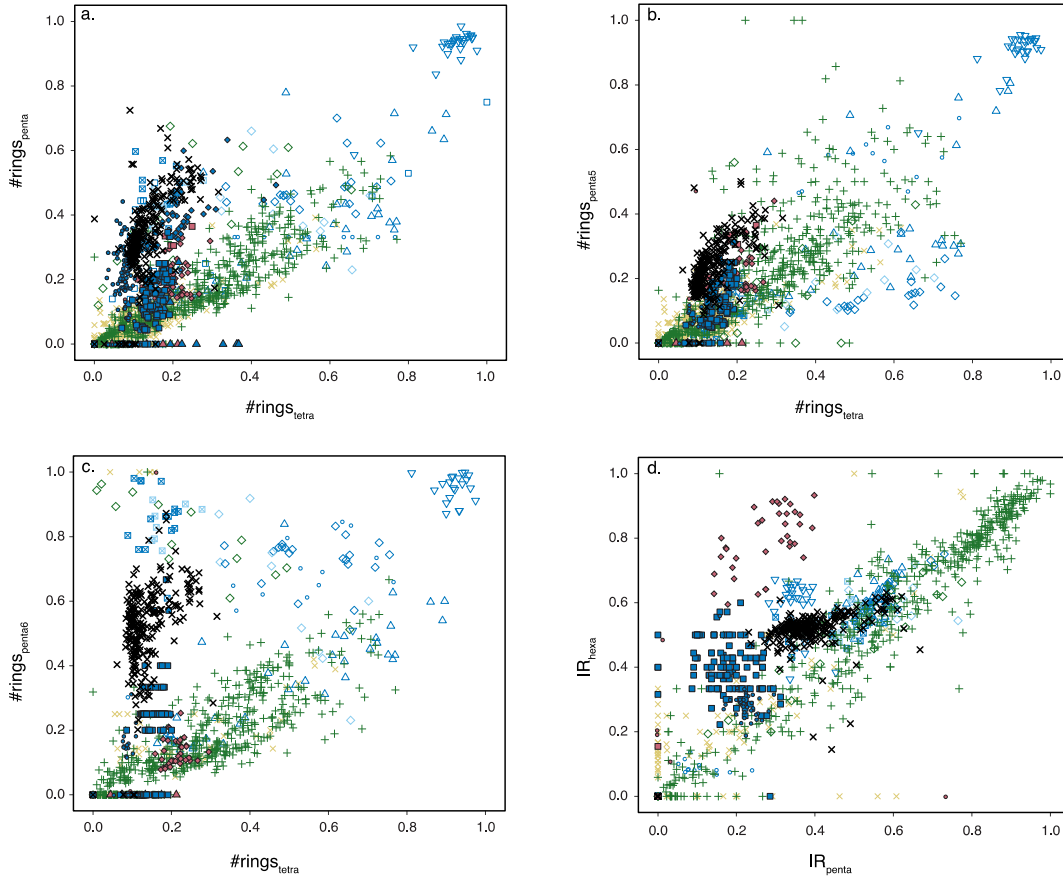
460 This generally indicates brGDGT contributions from sources other than soils. The smaller second cluster, containing exclusively mid-to-late Paleocene samples, is indeed close to that of modern continental shelf sediments (Berau delta and Svalbard; Sinninghe Damsté, 2016; Dearing Crampton-Flood et al., 2019) for which in situ brGDGT production

substantially exceeds soil contributions. This would generally suggest that the smaller cluster, and perhaps also the larger cluster of samples, have contributions from marine *in situ* brGDGT production. However, unlike those modern marine
465 sediments, our entire record does not show elevated #rings_{tetra} values that are associated to marine *in situ*-produced brGDGTs (Fig. 7). Instead, the low #rings_{tetra} values on our record (<0.3) suggest that brGDGTs have a terrestrial source. In the modern soil- and peat brGDGT dataset #rings_{penta} and #rings_{tetra} follow a coherent 1:1 correlation (Fig. 8a). At Site 1172, however, #rings_{tetra} is below 0.3, whereas #rings_{penta} ranges up to 0.6. This diverts from the 1:1 line but brings the brGDGT dataset
470 abundances of tetra-, penta- and hexamethylated brGDGTs plotted in the ternary diagram, where all Paleogene data plotted offset to that in modern soils. #rings_{tetra} and #rings_{penta} for the 5- (Fig. 8b) and 6-methyl brGDGTs (Fig. 8c) separately shows that the diversion of the 1172 data from modern soils and peats is primarily caused by a larger degree of cyclisation of the 6-methyl brGDGTs.

The samples within the Paleocene cluster have relatively high IR. The IR in our record averages ~0.4, but is elevated
475 between 62 Ma and 54 Ma, with peak values in the mid-Paleocene up to 0.6 (Fig. 7). Although both the IR_{penta} and IR_{hexa} are well within the range of the modern soil and peat data (Fig. 8d) they exceed the two available datapoints from modern southeast Australian soils, which have IR values of maximum 0.5 (De Jonge et al., 2014a), suggesting a contribution by non-soil sources (be it marine or river *in situ* production; De Jonge et al., 2014a). Given that the relatively low #rings_{tetra} values argue against a large contribution from marine *in situ* production, we suggest that river-produced 6-me brGDGTs
480 have contributed to the GDGT pool in the Paleocene samples with high IR. Therefore, for MBT'_{5me} calculation, we eliminate all brGDGT samples in our record that have IR values above that of Australian soils.

Finally, the consistent offset in brGDGT composition in Paleogene marine sediments relative to modern soils can potentially be explained by non-soil contributions even though many (most) indices suggest soil-dominated sourcing. Notably, not only Paleogene marine sediments from Site 1172 are offset from present-day soils, but this extends to other sites as well,
485 including Paleogene lignites that are not influenced by marine contributions. We suggest that there may be an additional evolutionary- or bacterial community factor that is at least in part responsible for the observed offset brGDGT distributions during the Paleogene. Except where indices suggest non-soil contributions, Paleogene marine sediments may contain a dominant soil-derived brGDGT signal in spite of their offset composition.

Fig.8



490 **Figure 8. Crossplots of various brGDGT indices. a. #rings_{tetra} vs. #rings_{penta}, #rings_{tetra} vs. #rings_{penta5}, #rings_{tetra} vs. #rings_{penta6}, IR_{penta} vs. IR_{hexa}. For color and symbol legend see Figure 6.**

4.2 SST and MAAT reconstructions

4.2.1 SST

By removing all samples with potential overprint from the record we can now assess the trends in TEX₈₆-based SST (Fig. 9a). Reconstructed SSTs differ by up to 4 °C between the exponential SST_{exp} and the linear SST_{lin} calibration in the warmest intervals, and progressively less with lower SSTs (Fig. 9a). The BAYSPAR method, which uses a linear relation between SST and TEX₈₆ yields the lowest SSTs for the low TEX₈₆ values in the Maastrichtian, Paleocene and middle–late Eocene, and SSTs between those based on the linear and exponential calibrations for the highest TEX₈₆ index values (Fig. 9a). U^K₃₇-based SSTs (Bijl et al., 2010, with BAYSPLINE calibration) around the MECO are in best agreement with the high SSTs derived from the linear TEX_{OBL} calibration, although in this TEX₈₆ range all calibrations yield SSTs within error (Fig. 9a).

In general, Maastrichtian (70.5–66.7 Ma) SSTs show a gradual cooling trend towards the late Maastrichtian–early Danian hiatus of ~800 kyrs. The early Danian (~65 Ma) is roughly 3 °C warmer than the late Maastrichtian (~67 Ma). The mid-Paleocene (62–59 Ma) shows two cooling steps: a ~2 °C cooling at 62 Ma, followed by a return to higher SSTs roughly at 60 Ma, and a ~5 °C cooling to minimum values of around 18°C around 59.5 Ma, where the record is truncated by a hiatus. A hiatus straddling the early Thanetian (59.0–57.7 Ma) likely obscures the onset of late Paleocene warming out of the mid-Paleocene temperature minimum. Background latest Paleocene SSTs fluctuate considerably, but the PETM warming clearly stands out (Sluijs et al., 2011), with a magnitude (5-7 °C depending on the calibration) comparable to that in other mid- to high latitude sections and similar to the global average (Frieling et al., 2017). Post-PETM SSTs drop back to pre-PETM values completely, followed by renewed warming towards the EECO. The magnitude of warming is 6 to 11 °C depending on the choice of calibration. Smaller early Eocene hyperthermal events, cannot be confidently identified at Site 1172. Highest SSTs of the EECO are slightly lower than peak PETM values in all calibrations, consistent with other records (Hollis et al., 2019; Inglis et al., 2020). However, a hiatus between 52.7 and 51.4 Ma may conceal peak EECO temperatures (Hollis et al., 2019; Lauretano et al., 2018). A gradual SST decline marks the end of the EECO around 49.2 Ma, which is broadly coeval with cooling inferred from benthic foraminiferal isotope records (Lauretano et al., 2018) and proxy records from New Zealand (Dallanave et al., 2016; Crouch et al., 2020). Post-EECO cooling is gradual and of small magnitude (2 °C from peak EECO values) until about 46.4 Ma, after which it accelerates (5 °C) between 46.2 Ma and 44.4 Ma. The age model in this particular interval is uncertain, due to a questionable assignment of chron C21n (Bijl et al., 2010; 2013b). Following the minimum SSTs at 44.4 Ma, SSTs rise again towards a plateau at 42.7 Ma. An SST minimum of ~22 °C is observed just prior to the MECO, at 40.2 Ma, followed by MECO warming of about 5–7 °C at 39.9 Ma. Post-MECO cooling seems more gradual than MECO warming, although this might be the result of fine-scale sedimentation rate changes (Bijl et al., 2010), which are not accounted for in our age model. SSTs are almost identical to those for the MECO at nearby Site 1170 on the South Tasman Rise (Cramwinckel et al., 2020). Post-MECO cooling transitions into gradual late Eocene cooling, down to a minimum of ~19 °C at 35.7 Ma. The following latest Eocene to Oligocene TEX₈₆-based SSTs (35–27 Ma) are 3 °C warmer, with surprisingly no cooling across the Eocene–Oligocene transition (Houben et al., 2019).

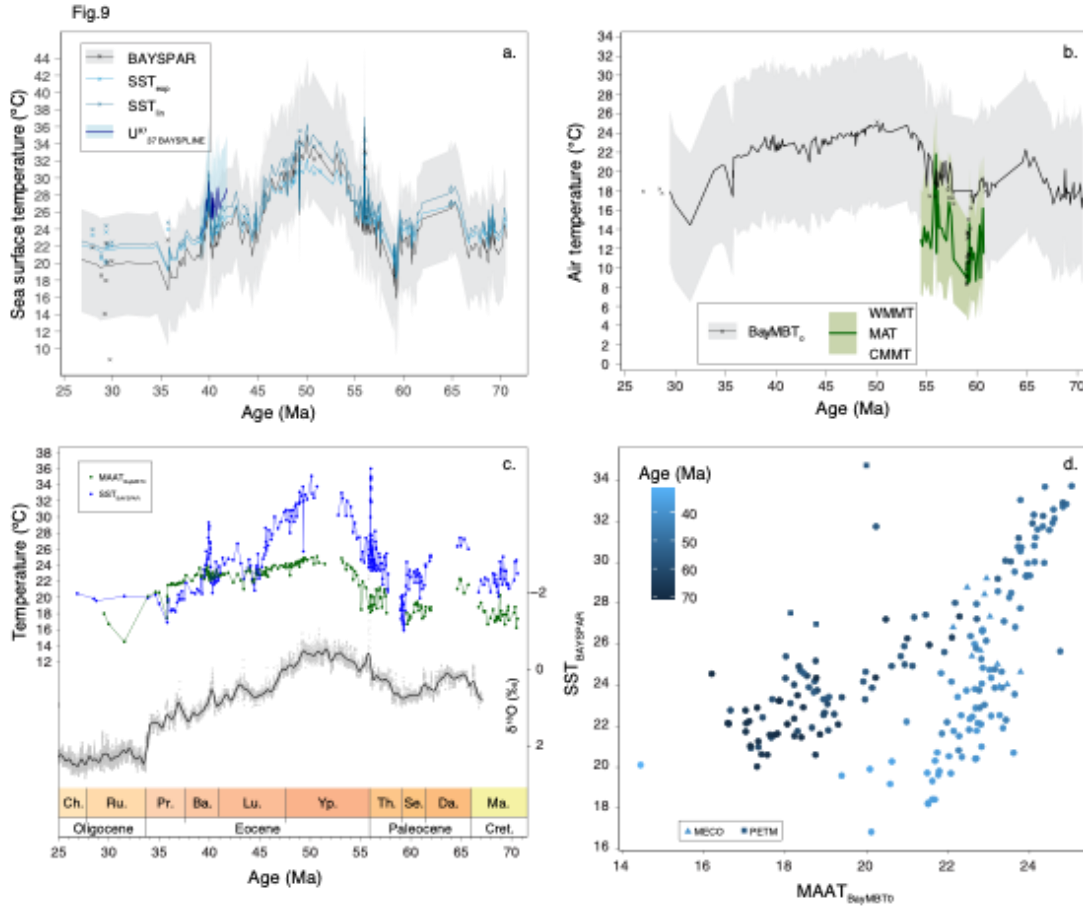


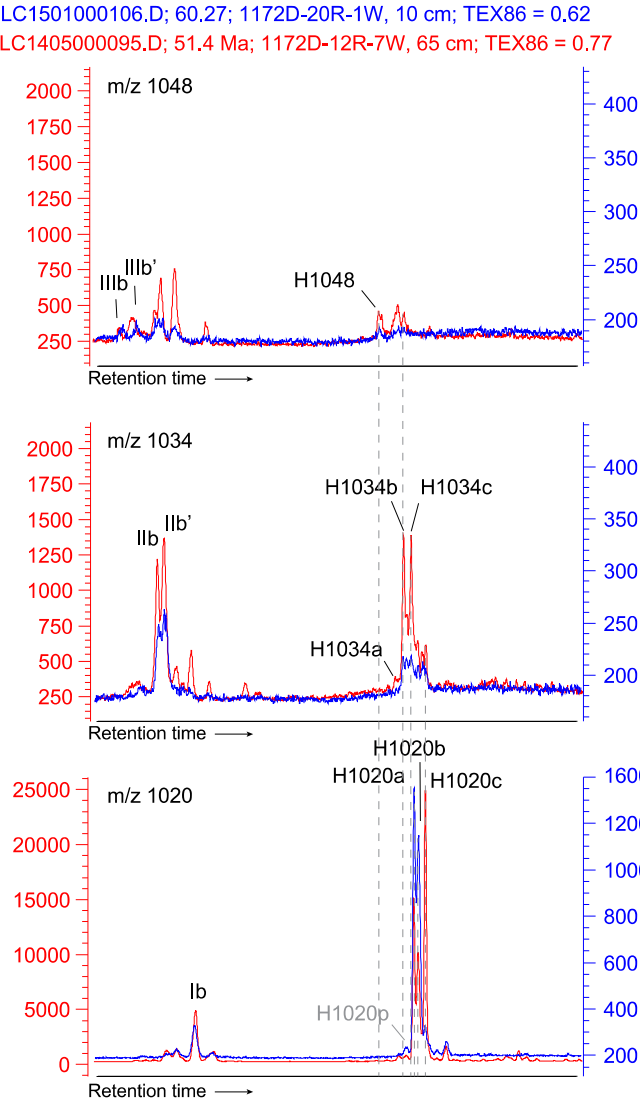
Figure 9. Paleothermometers. a. TEX_{86} based SST reconstructions, using a Bayesian (BAYSPAR; black, with 90% confidence interval in grey envelope; Tierney and Tingley, 2015) an exponential (SST_{exp} ; light blue; Kim et al., 2010) and a linear (SST_{lin} ; dark blue; O'Brien et al., 2017) calibration. Also plotted are $\text{U}^{\text{K}'}_{37}$ -based SSTs (thick blue line; from Bijl et al., 2010, recalculated using BAYSPLINE of Tierney and Tingley, 2018). Discarded samples are indicated by the crosses. b. brGDGT-based MAAT reconstructions using the Bayesian calibration (BayMBT_0 ; black, with 90% uncertainty interval in grey envelope; Dearing Crampton-Flood et al., 2020). Discarded samples are indicated in crosses. Also plotted are pollen-based NLR-approach mean annual (MAT) reconstructions (thick dark green, with light green envelope representing coldest and warmest month mean temperatures; from Contreras et al., 2014). c. SST and MAAT reconstructions using the Bayesian calibrations, with the CENOGRID benthic foraminifer oxygen isotope splice of Westerhold et al., 2020 (10pnt LOESS smooth in grey, 500 pnt LOESS smooth in black). d. cross-correlation between SST and MAAT, using the calibrations shown in c.

4.2.2 MAAT

After excluding samples with IR values above those of local soils, we present $\text{MBT}'_{5\text{me}}$ -based MAAT reconstructions. MAAT evolution reveals late Maastrichtian-early Danian warming of 3°C , gradual 4°C cooling towards the mid-Paleocene, gradual 8°C warming towards the EECO, with a conspicuous absence of warming at the PETM. Gradual cooling of 4°C starts at 49.5 Ma and continues into the Late Eocene. MAAT reaches a minimum at 43 Ma, followed by a plateau-phase. There is little warming that can be associated to the MECO. MAAT shows a conspicuous transient 3°C cooling in the Late

545 Eocene, at the onset of glauconite deposition. The Eocene-Oligocene Transition is characterized by another 5°C cooling. except during the late Eocene, SST is overall warmer than MAAT (Fig. 9c). This difference is largest during the EECO (up to 9°C).

Fig. 10



550 **Figure 10. Stack of 2 UHPLC-MS chromatograms (1172D-12R-7W, 65cm; 51.41 Ma in red and 1172D-20R-1W, 10cm; 60.27 Ma in blue) of m/z 1048, 1034 and 1020 traces, between ~40 and ~60 minutes retention time, showing the pattern of peaks corresponding to brGDGTs and brGMGTs. Peak labels refer to the molecular structures in Fig. 3. Note that not all peaks are 3x baseline.**

4.3 brGMGTs

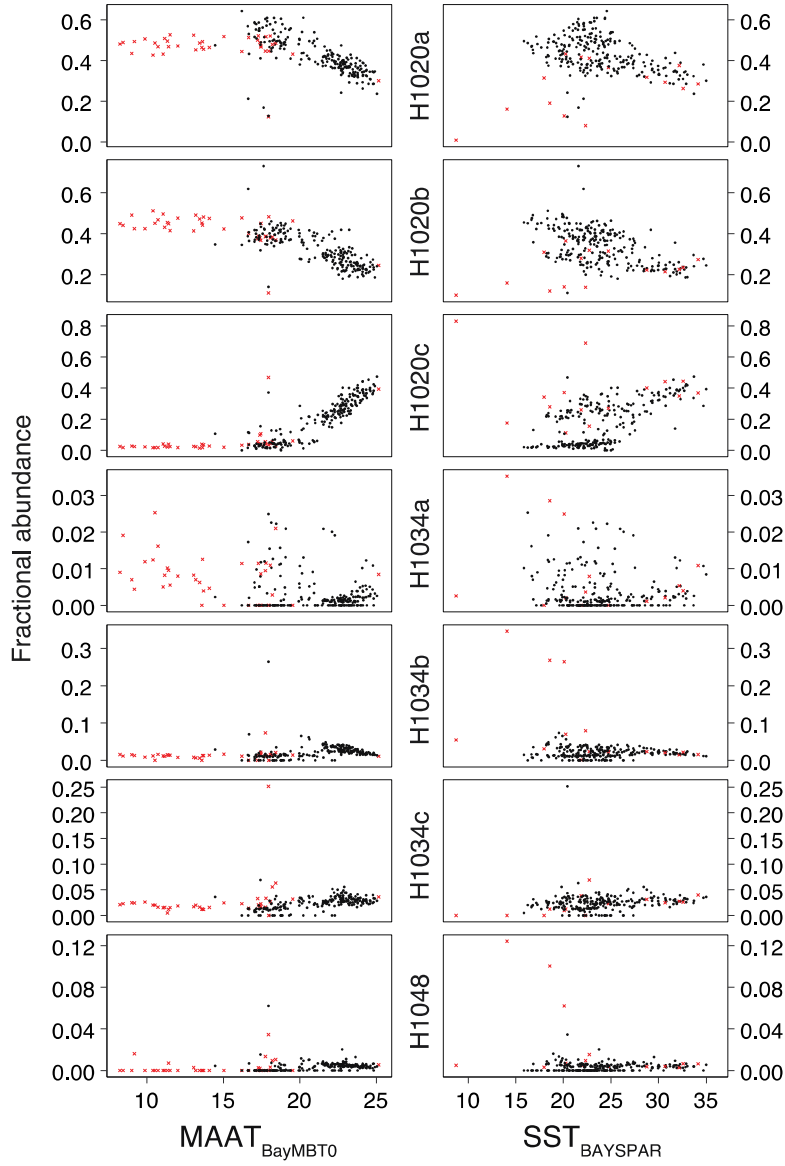
We detected brGMGTs throughout the record (see Fig. 10 for typical chromatograms). The exact molecular structure and position of the C-C bridge is still unknown but based on visual comparison with chromatograms of brGMGTs in African lake sediments (Baxter et al., 2019), (traces of) all 7 known brGMGTs can be identified. Next to H1020a, b, and c as most abundant brGMGTs, also H1034a, b and c are detected, although in some samples in the Maastrichtian, individual peaks could not be separated. Traces of H1048 can be seen at times, suggesting presence of this compound as well, albeit in low relative abundance. Next to these previously described peaks, we here recognized several other peaks that consistently occur at Site 1172: one compound that elutes just prior to H1020a, which we term H1020p (Fig. 10).

The fractional abundances of the H1020 isomers show qualitatively similar relations to temperature as in the East African lake dataset of Baxter et al. (2019): the relative abundance of H1020b has a negative relationship with MAAT, while H1020c has a positive relationship (Fig. 11). In addition, H1020a seems negatively correlated with MAAT in our data, although this compound does not show any relation to temperature in lakes (Baxter et al., 2019). The relative abundances of the H1034 isomers do not show a clear relationship with MAAT (Fig. 11), similar as in the tropical lake dataset, where the scatter increases at lower (H1034b) or higher (H1034a, c) temperatures (Baxter et al., 2019).

Downcore trends in brGMGTI are primarily controlled by the relative abundance of H1020c and closely follow those in both SST and MAAT. Upon cross-correlation, we note that brGMGTI has a higher correlation with MAAT ($R = 0.89$) than with SST ($R = 0.57$; Fig. 12). Interestingly, the correlations between brGMGTI and temperature indices only hold for part of the dataset. Samples with high IR values, for which brGDGT-based MAAT is influenced by a fluvial brGDGT contribution, fall outside the correlation (grey crosses in Fig. 12). In addition, we also note that brGMGTI approaches 0 at reconstructed MAAT of 15°C. The correlation with temperature (both MAAT and SST) is divided into 2 clusters. BrGMGTI values for samples from the younger part of the record, with BIT index $< \sim 0.3$, seem to correlate better to temperature than those from the older part of the record, with BIT index values $> \sim 0.3$ (Fig. 12). In the Oligocene, where many isoGDGT signals were evidently overprinted, the %brGMGTs increases (Fig. 3, 12d), and the brGMGT composition is characterized by a relatively higher abundance of H1020c, H1034b and H1048 compared to the rest of the record (Fig. 3; 12).

There is no clear relationship between %brGMGTs and brGDGT-based MAAT (Fig. 12h, i). This differs from the Paleocene-Eocene Arctic Ocean record (Sluijs et al., 2020), where the brGMGTI has no correlation to temperature, while %brGMGTs do. This may indicate differences in sources of brGMGTs between the sites, differences in oceanographic settings, or a non-temperature control on their distribution. In contrast, HMBT_{acyclic} appears to show similar trends to the brGDGT-based MAAT (Fig. 12g), as is also seen in the Arctic Ocean (Sluijs et al., 2020).

Fig. 11



585 **Figure 11. Fractional abundances of brGMGTs plotted against MAAT_{soil} and SST_{lin}. Red crosses are those samples that have non-thermal overprints on the paleotemperatures.**

Fig. 12

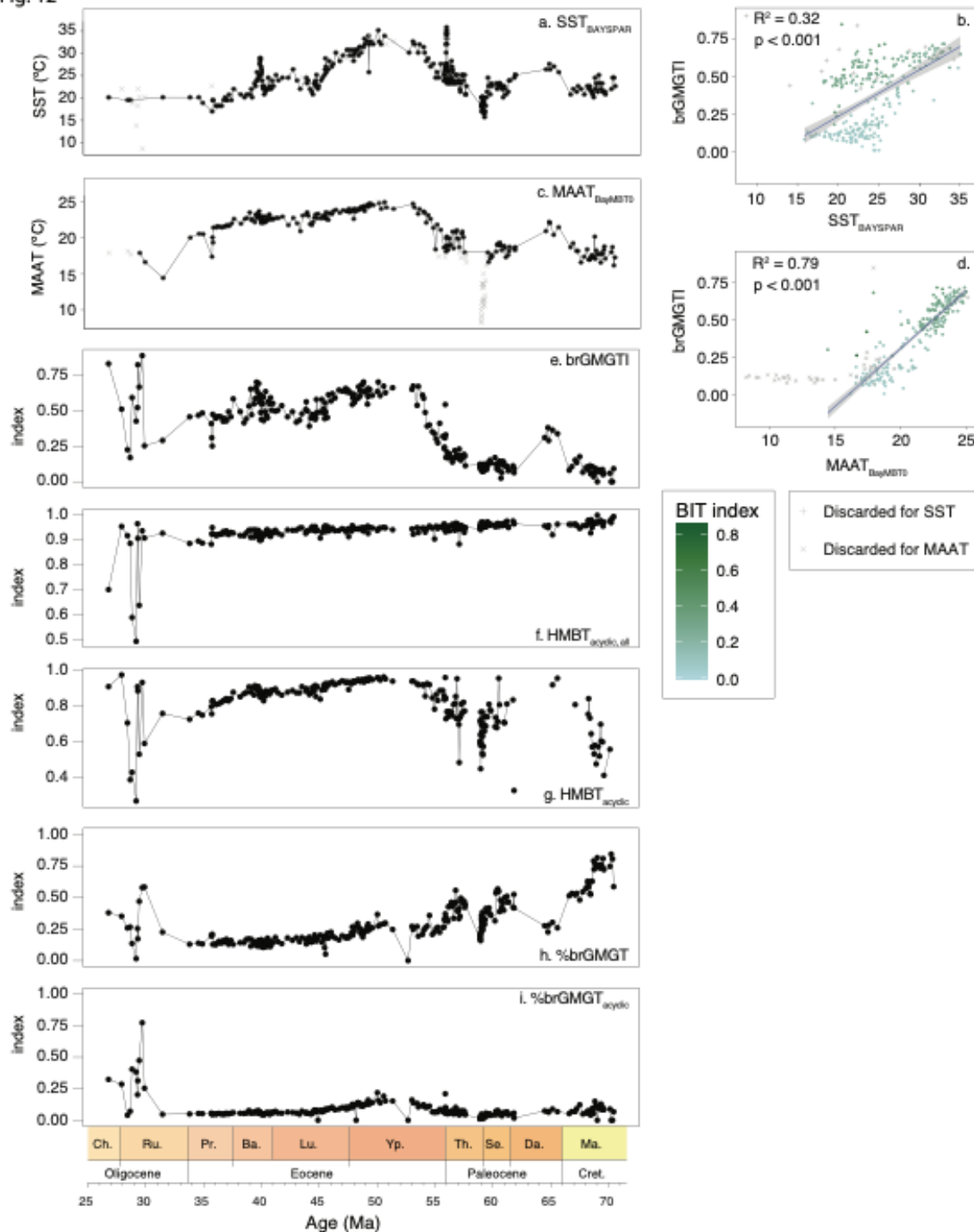


Figure 12. Time series and crossplots of brGMGT-based indices with MAAT and SST reconstructions. (a) SST_{in} (b) crossplot of SST_{in} and brGMGTI, (c) MAAT_{soil}, (d) crossplot of MAAT_{soil} and brGMGTI, (e) brGMGTI, (f) HMBT_{acyclic, all}, (g) HMBT_{acyclic}, (h)

590 %brGMGT (Baxter et al., 2019), (i) %brGMGT_{acyclic} (Naafs et al., 2018a). For equations see Table 1). Colour variable in the crossplots indicate the BIT index value.

4.4 Palynology

4.4.1 Assemblages

Dinocyst assemblages are dominated by *Manumiella* spp. throughout the Maastrichtian, together with Goniodomideae (notably *Dinopterygium* spp.), and *Cerebrocysta* cpx. (Fig. 13). From the late Maastrichtian onwards, *Senegalinium* cpx. increases gradually in relative abundance, interrupted by acmes of *Palaeoperidinium pyrophorum* in the Danian and *Glaphyrocysta* cpx. in the lower Selandian. *Senegalinium* cpx. is most abundant during the mid-Paleocene and the Paleocene–Eocene transition, while the PETM itself is characterised by abundant *Apectodinium* (~30%) and common *Senegalinium* cpx. and Goniodomideae (Sluijs et al., 2011). After the PETM, *Senegalinium* cpx. abundance temporarily increases, and followed by high abundance of *Deflandrea* cpx, *Spinidinium* cpx and *Elytrocysta* spp. with minor contributions of various other genera, e.g., *Wetzeliiella*, *Hystriochokolpoma*, Goniodomideae (Fig. 13). From 45 Ma to ~37 Ma, *Enneadocysta* spp. abundance oscillated with *Deflandrea* cpx. and *Spinidinium/Vozzhennikovia* cpx, with contributions of *Phthanoperidinium* spp. (Röhl et al., 2004a). Protoperidinioid dinocysts become common in the late Eocene, around 35.5 Ma (Houben et al., 2019; Sluijs et al., 2003; Fig. 13).

Fig. 13

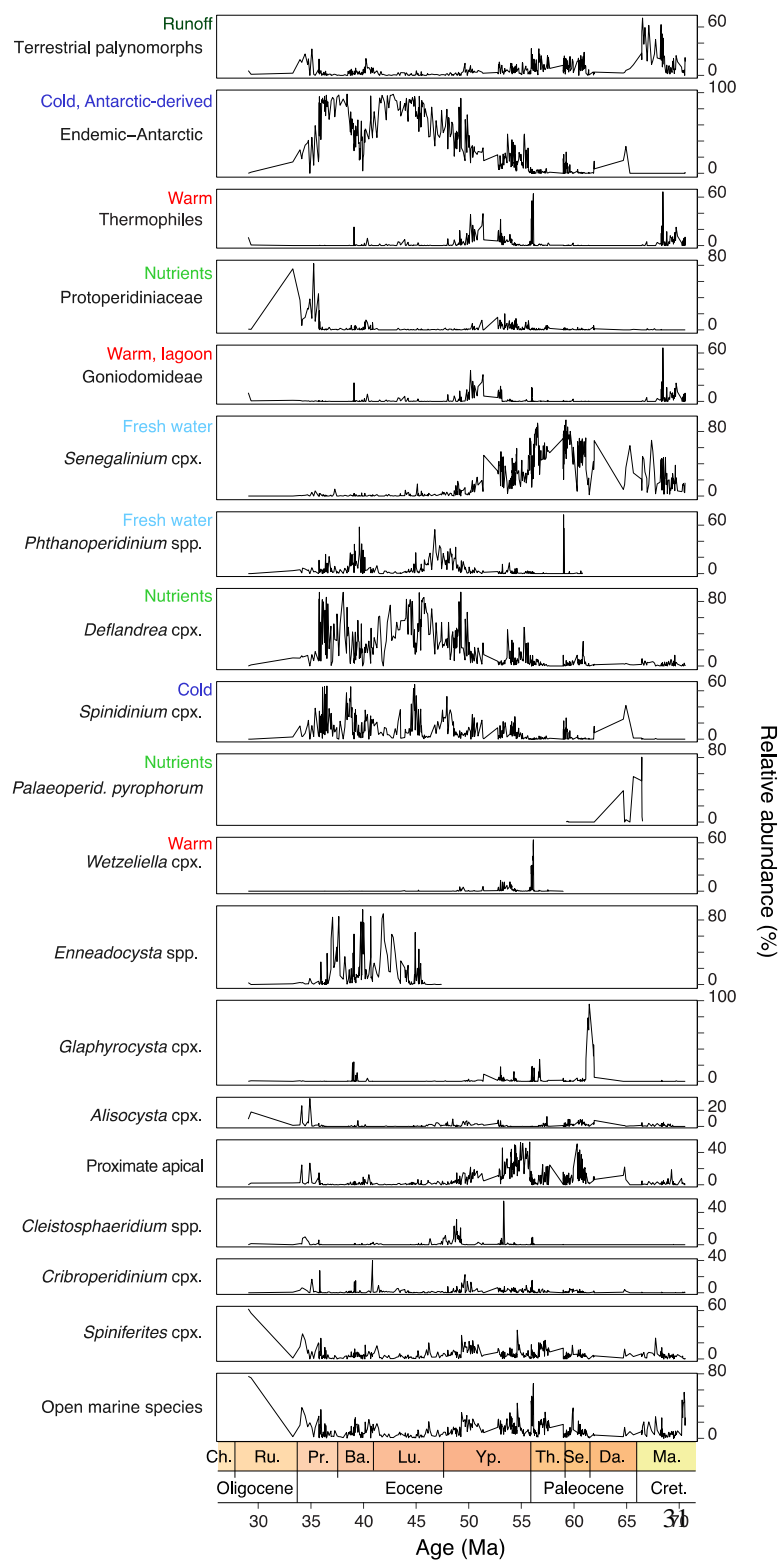


Figure 13. Dinocyst eco-groups and %terrestrial palynomorphs from Site 1172. From left to right, eco-groups are ordered in an in-shore-to-offshore transect. Relative abundance ticks represent 20%.

4.4.2 Diversity and variability

Together with the changing assemblage composition, some strong and coherent changes in diversity and variability indices are observed (Fig. 14). First, results are highly similar for the full species-level dataset and the grouped assemblages, indicating robustness of both the grouping and diversity analysis. The Maastrichtian–Paleocene assemblages are characterized by relatively low diversity, which increases towards the EECO, as variability indicated by the Gibbs index decreases. After this, diversity drops and variability increases during middle–late Eocene cooling ~49–38 Ma, with an interruption around 41–39 Ma. Towards the top of the record, both diversity and variability increase.

Fig. 14

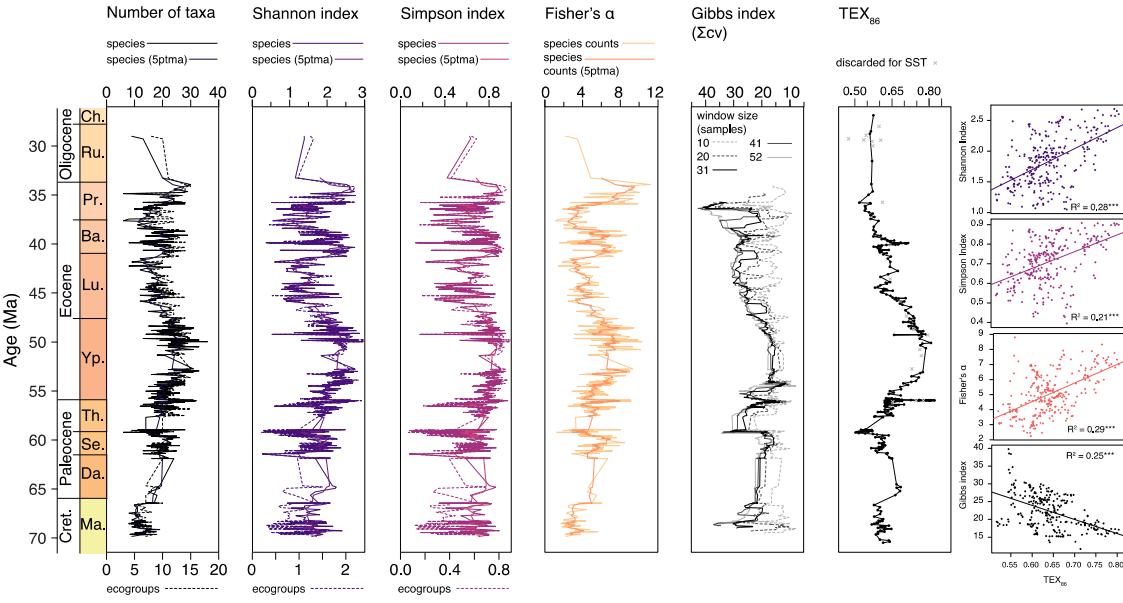


Figure 14. Dinocyst taxonomic diversity and variability through time. From left to right are plotted number of taxa (black), Shannon index (purple), Simpson index (pink), Fisher’s alpha (orange) (species-based dataset in solid lines; eco-grouped dataset in dashed lines) and Gibbs index (note the reversed axis; different window sizes in grey-black). TEX₈₆ plotted for comparison. Grey crosses indicate samples that have non-pelagic or non-thermal contributions. Crossplots show correlation of these indices with TEX₈₆.

5. Discussion

5.1 Air and sea surface temperature evolution compared

The near-shore depositional setting of our record should have recorded a coastal terrestrial environment, with MAATs similar to the local SSTs. Indeed, the brGDGT-based MAAT record shows strong similarities to SST particularly in multi-million year trends (Fig. 9). Specifically, the early Danian is warmer than the Maastrichtian, and both records show a 2-step

Paleocene cooling, late Paleocene warming towards peak values in the EECO, and middle–late Eocene cooling. On shorter time scales both records, occasionally even on sample-to-sample level, show synchronous variability (Fig. 9). The Bayesian calibrations for MAAT and SST generate overlapping median temperatures for the colder middle-late Eocene and Paleocene, but in the warmer Eocene time intervals SSTs remain consistently warmer (by $\sim 8^\circ\text{C}$). The lower MAATs cannot be explained by a saturation of the MBT'_{5me} index that is underlying the paleothermometer, as maximum recorded MBT'_{5me} values are around 0.85 in the EECO. The large calibration errors of BAYSPAR ($\pm 6^\circ\text{C}$) and BayMBT₀ ($\pm 8^\circ\text{C}$) in principle could resolve the offset. However, applying the extremes in calibration error to make them consistent with each other would make both profoundly inconsistent to the independent SST and MAAT estimates from the same samples. More importantly, the colder MAAT compared to SST seems to be a consistent feature in many regions where TEX₈₆ and MBT'_{5me} were applied together (see e.g., Hollis et al., 2019 and compare Willard et al., 2019 and Sluijs et al., 2020). The offset between MAAT and SST would have been more variable between sites if it were only the result of calibration errors. TEX₈₆-based SSTs in the southwest Pacific realm have been on the high end of many multi-proxy-based temperature reconstructions for the Eocene (Hollis et al., 2012), and therefore may be assumed to have an as yet poorly understood warm bias. However, the consistency of TEX₈₆-based SSTs with other SST proxies for specific time intervals and locations (Bijl et al., 2010; Hines et al., 2017) suggests the SST-MAAT offset cannot be easily reconciled by only invoking a warm bias in TEX₈₆, as was also shown by Hollis et al. (2012). MAAT reconstructions on the other hand have been broadly consistent with nearest living relative based temperature reconstructions on pollen assemblages in many applications in the Eocene (Pross et al., 2012; Contreras et al., 2013; 2014; Pancost et al., 2013; Willard et al., 2019), but both brGDGT and plant-based temperature estimates are arguably prone to cold biases at the high-temperature end (e.g. Naafs et al. 2018b, Van Dijk et al., 2020). In addition, it may be that the same evolutionary or bacterial community factors that make Paleogene brGDGT assemblages deviate from that in modern soils, may also cause a deviation in the calibration to MAAT. This may resolve some of the offset between MAAT and SST reconstructions. Although this implies that quantitative MAAT estimates based on MBT'_{5me} in non-analogue settings such as the present one should be interpreted with care, the strong temperature dependence in Paleogene brGDGTs cannot be ignored.

Another surprising result is that the brGDGT-based MAAT record does not capture PETM and MECO warming (Fig. 9b), while pollen-based MAAT does. This is remarkable for several reasons: (1) brGDGTs were measured on the same samples as isoGDGTs, which precludes a sampling bias; (2) SST changes of longer duration (Maastrichtian–Danian warming, mid-Paleocene cooling and early Eocene warming towards the EECO) are represented in the MAAT record; (3) The duration of PETM ($\sim 150\text{--}250$ kyrs; Röhl et al., 2007, Zeebe & Lourens, 2019) and MECO (~ 400 kyrs; Bohaty et al, 2009) is too long to explain the absence in MAAT warming with a lag in soil-derived OM delivery to the ocean (up to several kyr, see, e.g., Feng et al., 2013; Schefuß et al., 2016; Huurdeman et al., 2020); (4) seems incompatible with the fact that other low-amplitude, shorter-term SST changes are reflected in the MAAT record (Fig. 9), which suggests that soil-derived OM did capture short-term climate variability on the time resolution of the samples; (5) Other brGDGT-based records from the region do show a temperature response during the PETM (Hollis et al., 2012; Pancost et al., 2013, Huurdeman et al. 2020). A lack of MAAT

660 rise during the PETM and the MECO could be explained by a switch from brGDGT sourcing from soils to (peaty) lakes, which are notoriously cold-biased (Tierney et al., 2010). Paleocene–Eocene peats are abundant in southeast Australia (Holdgate et al., 2009), and it is possible that a contribution from peat lakes at times of global warming, contributed to a dampening of the proxy-response. This may also explain the lack of response at the EECO and explain why MBT'_{5me} does not reach saturation in that interval. The final option, which, although unlikely, cannot be excluded, is that a dominant
665 fraction of the brGDGT was produced in situ and subdues the temperature response during these warming phases.

The subdued middle-late Eocene cooling in the air temperature record compared to the strong decrease in SST is particularly clear when cross-plotting SST *versus* MAAT (Fig. 9d): the relationship between the two proxies is different in the Maastrichtian–early Eocene compared to the middle–late Eocene. This may be related to the start of a progressive northward tectonic drift of the Tasmanian hinterland, which occurred around the same time (Fig. 1). A similar subdued MAAT record
670 relative to SST cooling was observed in records from New Zealand (Pancost et al., 2013; Crouch et al., 2020), which was also moving northward at that time. This puts the hinterland of the soil-derived brGDGTs into warmer climate zones throughout the middle–late Eocene, while the ETP remained under influence of the Antarctic-derived Tasman Current (TC) through that time (Huber et al., 2004; Sijp et al., 2016). The TC cools and likely strengthens in the middle and late Eocene. The terrestrial climate cooled as well, but this signal will be subdued if the soil material is sourced from a progressively
675 lower-latitude environment as Australia drifted northward. Yet, one would expect that the close coupling between coastal and offshore temperatures in near-shore environments would subdue such tectonic effects in mean annual temperature reconstructions.

Both marine and terrestrial temperature records diverge once more during the late Eocene–Oligocene, when they show a sharp cooling of 2–3 °C at the onset of greensand deposition, at ~35.5 Ma. Whereas this is merely a continuation of
680 progressive late Eocene cooling in the SST record, the decrease in MAAT is more pronounced compared to the middle–late Eocene trends. Another 3–4 °C MAAT drop occurs sometime between the late Eocene and the early Oligocene (likely associated with the Eocene–Oligocene transition), while SSTs revert to warmer, middle Eocene-like values. The continued sea surface warmth across the EOT has been related to persistent influence of the proto-Leeuwin Current (Houben et al., 2019), which apparently retained a constant temperature across the EOT. Since MAAT decreased while SSTs remained high,
685 the influence of regional oceanographic changes did not appear to impact climate changes in the source region of brGDGTs during this time interval. Later, at 30 Ma, MAAT does rise in the region, which could be related to either the progressive northward movement of the site, or a response to widespread warming in the early Oligocene (e.g., Westerhold et al., 2020). This is surprising given the proximity of the sediment record to the coastline, but it represents further evidence that hinterland MAAT and near-shore SST trends can be decoupled under certain circumstances. MAAT temperatures from Site
690 1172 around EOT are somewhat colder than those from Southern Australia (Lauretano et al., 2021) although calibration error bars do overlap.

Minimum mid-Paleocene SSTs are lower than those for the early Oligocene at Site 1172. However, the site migrated northward by ~7 ° of latitude between the mid-Paleocene and the early Oligocene (Fig. 1) and the oceanographic regime

changed with throughflow of the proto-Leeuwin Current through the TG in the Late Eocene (Houben et al., 2019). More remarkably, the mid-Paleocene SSTs approach those of the warmest interglacial intervals of Oligocene Wilkes Land Margin, east Antarctica (Hartman et al., 2018), at a similar paleolatitude as the Paleocene ETP. This suggests that mid-Paleocene Antarctic-proximal temperatures were similar to those at times of major Antarctic glaciation during EOT. Previous work has indeed presented widespread evidence for low sea level during this time interval (e.g., Guasti et al., 2006; Frieling et al., 2014; 2018; Hollis et al., 2014), which combined with low SSTs suggests the presence of some continental ice on the Antarctic continent during this mid-Paleocene interval.

5.2 BrGMGTs

The continuous presence of brGMGTs in our record appears unrelated to depositional conditions, and distributional changes of brGMGT isomers do not correlate to varying relative contributions of terrestrial material. This suggests that at least part of the brGMGTs has a marine origin, in line with observations from modern sediments and water columns (Liu et al., 2012, Xie et al., 2014) and from Paleogene Arctic Ocean sediments (Sluijs et al., 2020). Their sparse presence in modern mineral soils (Peterse, F., pers obs., based on metadata from De Jonge et al., 2019; Kirkels et al., 2020) seems to confirm this, although abundant occurrence in peats and lakes (Naafs et al., 2018a, Baxter et al. 2019, Tang et al., 2021) shows clear potential for terrestrial input. Despite the uncertainties in sourcing of brGMGTs, we find signals in the brGMGTs at Site 1172 resemble those from East African lakes. For example, the brGMGT index (brGMGTI), which was derived to quantify the relationship between brGMGT composition and MAAT in surface sediments of East African lakes (Baxter et al., 2019), seems to be correlated to temperature in our record as well (Fig. 12b, d). In addition, H1020c is practically absent in the cold mid-Paleocene part of our record, similar to in cold ($<12^{\circ}\text{C}$), high-altitude lakes (Baxter et al. 2019), which suggests that the biosynthesis of H1020c occurs only above a certain temperature, or that the microbes that biosynthesize H1020c do not live at low temperatures. However, contrasting to in lakes, site 1172 lacks H1048 isomers in the cold mid-Paleocene interval, illustrating that there are also differences between fresh water and marine depositional settings.

The HMBT record does not reflect a temperature signal when using all brGMGT isomers in the calculation (Fig. 12f), as H1020 isomers are by far the most abundant brGMGTs at Site 1172, and any change in the abundance of H1020c is compensated by both H1020b and H1020a in our record. In contrast, only using the H1020c and H1034b isomers does show some correlation to temperature in our record (Fig. 12g).

%brGMGT does not show a clear relationship with temperature (Fig. 12h, i), in contrast to observations in modern peats (Naafs et al., 2018a) and lake sediments (Baxter et al., 2019). We do note the extremely high %brGMGT (~80%) in the Maastrichtian and Oligocene, much higher than seen elsewhere. We also do not see a clear inverse relationship between %brGMGT and BIT index, as in the Arctic Eocene record of Sluijs et al. (2020). There, it does correlate with SST, which we do not see either. These discrepancies may be because of the complexity in the sources of brGDGTs in the records.

BrGMGTs are more abundant and different in composition in the Oligocene, compared to the rest of the record. This coincides with abundant signs of overprints in all br- and isoGDGT indices, which may indicate a source change of

brGMGTs in this interval, e.g., related to the same changes in contribution that cause the overprints in isoGDGTs. The differences in brGMGT signals between our record, the Eocene Arctic Ocean record and modern sediments demonstrate that more research is needed on their modern sources and environmental dependencies to further assess their use and value in paleoclimate studies.

5.3 Integrated environmental and climatological reconstruction

5.3.1 Sea level and ecosystem response to SST changes

The abundance of temperature-responsive dinocyst eco-groups qualitatively confirm our SST trends. The thermophile dinocyst eco-group shows higher relative abundance at higher SSTs (Fig. 15). This eco-group has been empirically linked to SST in a global compilation of late Paleocene-early Eocene records (Frieling and Sluijs, 2018), and we here see the same relationship for the longer early Paleogene. The somewhat muted response of the dinocyst assemblages to the large EECO SST rise is consistent with that in other SW Pacific records and microfossil groups (Crouch et al., 2020), and might be related to the already warm background conditions upon which this warming took place. Moreover, the abundance of SST-insensitive dinocysts in the assemblage and absence of typical cold-indicators could further explain the muted response. The relationship between isoGDGT-based SSTs and relative abundance of the supposedly cold-affiliated endemic dinocysts (Bijl et al., 2011) is not as clear. This might be because the majority of dinocyst species within the Endemic-Antarctic dinocyst eco-group are restricted to the Eocene (Bijl et al., 2011). The absence of endemic-Antarctic dinocysts in the cold mid-Paleocene is interesting, given that some endemic species had already appeared (Bijl et al., 2013b). It might be that the strong fresh-water influence negatively affected their proliferation in the mid-Paleocene. Indeed, from 50 Ma onwards, SST has a negative correlation with the %endemic Antarctic dinocyst abundance in the record (Fig. 15; see Bijl et al., 2011), which may indicate a more distinct biogeographical separation through the development of stronger meridional gradients or evolutive adaptation of the endemic dinocyst community to more polar environments following the extreme warmth of the EECO.

We particularly note the correlation between TEX_{86} -based SST and the relative abundance of more open ocean-affiliated dinocysts (Fig. 15), which suggests that SST correlates with regional sea level. The thermophilic and Open Ocean eco-groups share some outer neritic species. However, the strong covariance is caused by the absence of other Open Ocean species, such as *Impagidinium* spp., *Spiniferites* spp., and *Operculodinium* spp. Moreover, sea level trends derived from Open Ocean dinocysts are supported by weight % CaCO_3 in these sediments in the middle Eocene (Röhl et al., 2004a), on orbital and longer time scales. SST-induced sea level changes hint at steric effects as driver. Indeed, because of the flatness of continental shelf areas during long-term greenhouse climates (Somme et al., 2009) small changes in regional sea level will cause large changes in distance to shore and associated characteristics, including e.g. salinity, nutrients and suspended sediment loads, which subsequently shapes the dinocyst assemblages.

The SST control on dinocyst assemblages is further demonstrated in diversity and assemblage variability indices. Throughout the long time interval covered by our record, dinocyst richness and diversity show a remarkably similar pattern to TEX₈₆-based SST. Species richness, as well as diversity expressed in the Shannon and Simpson indices and Fisher's alpha reach a maximum during the prolonged warmth of the EECO and drop during middle-late Eocene cooling. These relationships are demonstrated by modest correlation coefficient between diversity and TEX₈₆ (Fig. 14). This pattern holds for the long-term trends, as well as for the short-term PETM, but, surprisingly, not for the MECO. Dinocyst diversity is known to peak mid-way an inshore-to-offshore transect in the modern and Eocene oceans (Brinkhuis, 1994; Pross and Brinkhuis, 2005). As our assemblages indeed consist of many mid-shelf species, the correlation between diversity and SST might thus be indirectly related through habitat size, with expansion of the flat shelf area and thereby increase in shelf niches during the EECO (Somme et al., 2009). Variability as indicated by the Gibbs index records the opposite pattern to diversity for the long-term trends into and out of the EECO (Fig 14). That is, the lower-diversity assemblages in general have a higher variability, or sample-to-sample fluctuations within the relative abundances. This indicates that the higher diversity dinocyst assemblages might be more ecologically resilient.

5.3.2 Marine environmental response to runoff changes

Senegalinium cpx. abundance broadly correlates with the IR (Fig. 16), which signals input of river-produced brGDGTs. *Senegalinium* cpx. is generally thought to have tolerated low salinity environments (Sluijs and Brinkhuis, 2009; Barke et al., 2011). These two proxies together confirm a large input of fresh water at this site during the mid-to-late Paleocene in line with high relative abundance of terrestrial palynomorphs, low abundances of open ocean dinocysts and high IR values (Fig. 16). Interestingly, *Phthanoperidinium* spp., which is also generally associated with low-salinity to near fresh-water conditions (Sluijs and Brinkhuis, 2009; Barke et al., 2011; Frieling and Sluijs, 2018) does not correlate with IR (Fig. 16). Because we find different species of *Phthanoperidinium* spp. than those previously associated to fresh-water conditions (e.g., Barke et al., 2011), it could be that the fresh-water tolerance of *Phthanoperidinium* spp. was not shared among all species. It could also be that some species of *Phthanoperidinium* spp. were slightly less euryhaline than *Senegalinium*, contrary to the species noted by Barke et al. (2011).

Fig.15
GDGT- and palynology-based proxies for SST

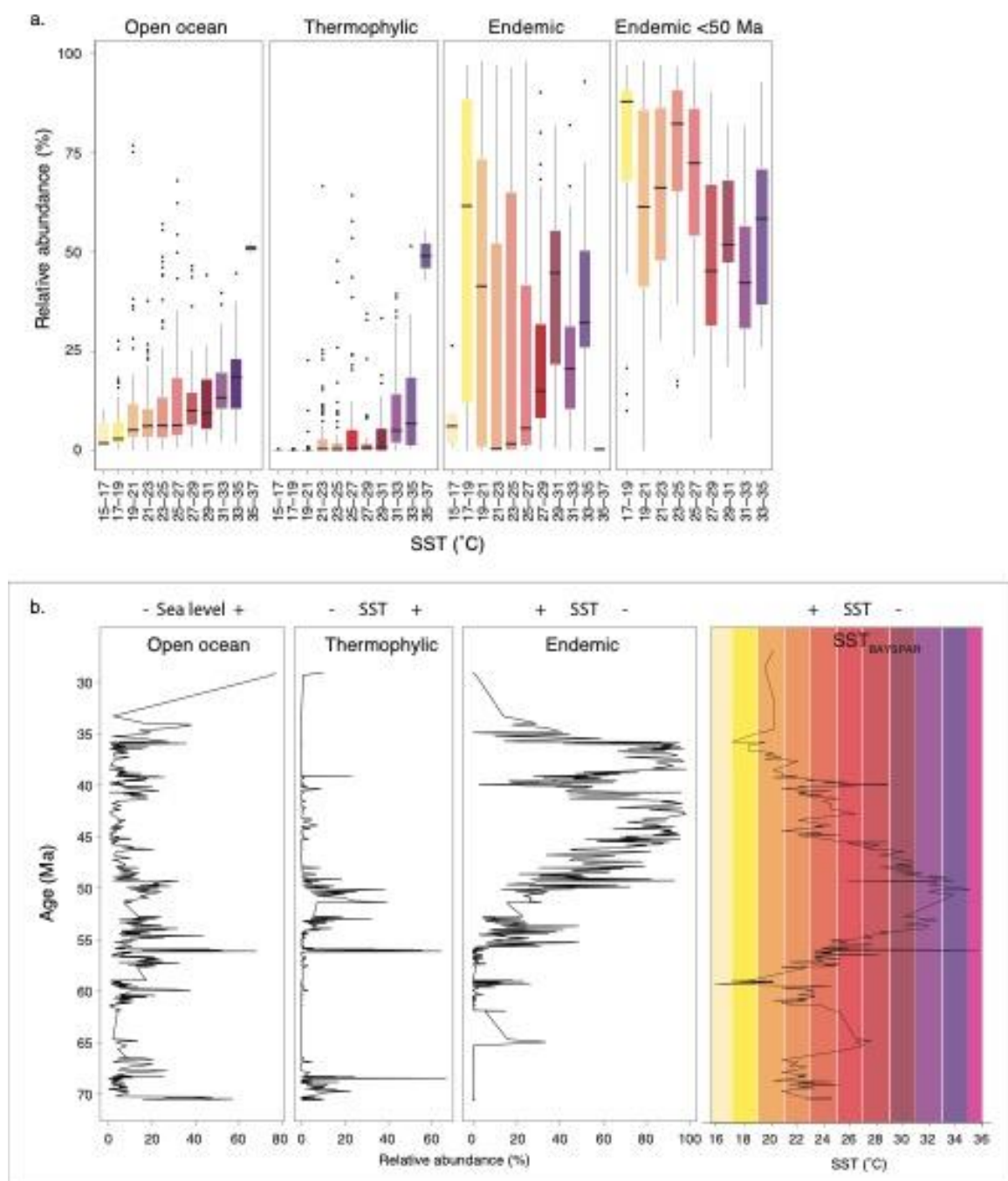


Figure 15. isoGDGT- and palynology-based proxies for SST. a. relative abundance of Open ocean, thermophylic and endemic-Antarctic dinocyst eco-groups (for groups see Table 3; total dataset and samples <50Ma only) plotted against linearly interpolated (resampled) TEX₈₆-based SST, in 2 degree bins. Median (black bars), lower-to-upper quartiles (25–75 percentile; coloured envelopes), lower to upper whiskers (95% confidence intervals; black lines), outlier samples (black dots). b. Time series of the same dinocyst eco-groups and SST.

Tasmania was located at latitudes between 55° and 60° S, in the middle of the region of strong westerly winds, within the range of low atmospheric pressure and received a large amount of precipitation during the Paleocene (Huber and Caballero, 2011). River input from Tasmania and perhaps also Australia could have reached the site. Interestingly, while Tasmania migrated northwards only a few degrees latitude between 60 and 40 Ma (Fig. 1), under warmer temperature conditions as in the mid-Paleocene, the freshwater input decreased, based on both *Senegalinium* cpx., terrestrial palynomorph abundance (decimated abundance from 50 Ma onwards) and the IR (decrease to normal values around 54 Ma; Fig. 16). This suggests either a rerouting of river input or a change in hinterland precipitation regime. Support for the latter comes from clay mineralogical data, showing an increase in kaolinite starting at 50 Ma (Fig. 2; Robert, 2004). Kaolinite forms abundantly at the base of acidic peats (Staub and Cohen, 1978; Korasidis et al., 2019) and in old, leached soils, which were ubiquitous in the Eocene Australian hinterland (Holdgate et al., 2009). While the hinterland could well retain the kaolinite-rich clays during the Maastrichtian and Paleocene, because the site was under a year-round high precipitation regime, it was less efficient in doing so when the hinterland experienced seasonally more variable climatic conditions as Australia migrated northward. The increase in kaolinite delivery to Site 1172 is hence interpreted as a signal of enhanced soil (or peat) erosion from the catchment areas in Tasmania and SW, as a result of a (seasonally?) more variable climate regime.

5.4 Environmental and climatological changes in the catchment

The long-term trends in the BIT index are at odds with all the river runoff indicators in our data (Fig. 7). BIT remains low during the interval in which increased river water discharge is inferred, but increases when this river-runoff signal decreases. This is unexpected because the BIT index should reflect high input of soil-derived brGDGTs into the marine sediments, associated with increased discharge (Hopmans et al., 2004). There are two ways to explain this signal. The BIT index could be responding to marine crenarchaeol production, rather than to the terrestrial brGDGT flux. Indeed, marine productivity could have been spurred by runoff and associated nutrient delivery to the coastal zone during the Paleocene, and decreased in the Eocene, thereby lowering and increasing BIT index values, respectively. Another explanation is that the BIT index has been influenced by in-river crenarchaeol production, although crenarchaeol concentrations in rivers are normally negligible compared to those produced in the marine realm (Zell et al., 2013). Decreasing BIT index values with increasing river- and soil input has been related to climate in the catchment in the late Quaternary Zambezi river (Lattaud et al., 2017), where hinterland vegetation and seasonality in precipitation controls soil-OM input to the marine realm. Studies of modern soil-river-ocean pathways in the Amazon River system show that whereas river-suspended brGDGTs in its upper tributaries during the wet season reflect catchment soils (Kirkels et al., 2020), the lower Amazon contains increasing contributions of *in situ* produced brGDGTs, especially during the dry season, when soil mobilisation is limited (Zell et al., 2013). In addition,

soil-derived brGDGTs appear to be rapidly lost upon entering a river, and are only effectively transferred to the ocean floor in catchments with a distinct rainy season or during periods with heavy rainfall (e.g. Weijers et al., 2007b; Guo et al., 2020).

820 The marine surface sediments of the Amazon fan consequently contain a mix of soil, fluvial, and marine brGDGTs (Zell et al., 2014). Thus, we here propose the following scenario to explain the BIT index trends for Site 1172: In the early Paleocene–early Eocene, year-round precipitation caused strong fresh-water delivery into the SW Pacific. Soil-derived brGDGTs were partly remineralized in the river catchment, and the strongly diluted soil-derived OM was dominated by river-produced brGDGTs, which suppressed the BIT index. This explains the unexpectedly cold MAATs derived from the
825 mid-Paleocene samples with high IR. The high fluvial discharge also promoted pelagic isoGDGT production, including crenarchaeol, due to associated nutrient delivery, further lowering the BIT index. Later in the early Eocene, precipitation in the hinterland decreased and became more seasonal as the hinterland gradually drifted out of the zone of intense precipitation. This led to a more effective transfer of soil-derived brGDGTs and kaolinite to the marine realm, increasing the BIT index at the ETP.

830

Fig. 16

GDGT- and palynology-based proxies for runoff

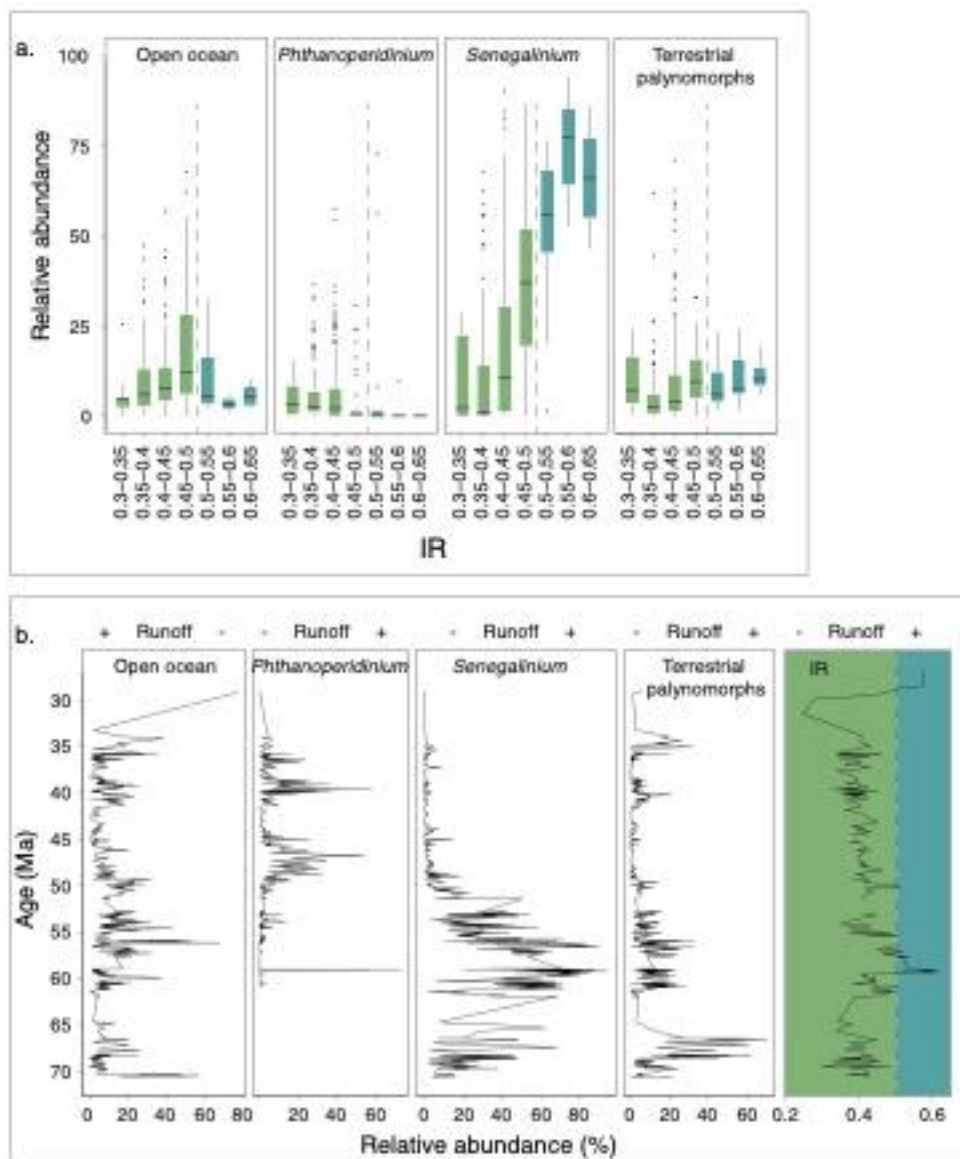


Figure 16. BrGDGT- and palynology-based proxies for river runoff. a. relative abundance of Open Ocean, *Phthanoperidinium* spp., *Senegalinium* cpx. (for groups see Table 3) and terrestrial palynomorphs in 0.05 index unit bins of the resampled IR. Dashed line separates elevated IR bins from non-elevated IR bins. Median (black bars), lower-to-upper quartiles (25–75 percentile; coloured envelopes), lower to upper whiskers (95% confidence intervals; black lines), outlier samples (black dots). b. Time series of the same palynology eco-groups and the IR.

6. Conclusions

We have analysed GDGT- and dinoflagellate cyst data in samples from the Maastrichtian–lower Oligocene sediments at ODP Site 1172. IsoGDGTs represent a pelagic signal throughout the Maastrichtian–Eocene but are influenced by sedimentary-produced isoGDGTs in the Oligocene. BrGDGTs at this site are likely primarily soil- or peat-derived, albeit evolutionary changes in brGDGT producers may be responsible for Paleogene-specific brGDGT signals. Exceptions are the mid-Paleocene, where in river-produced brGDGTs influence the record, and the Oligocene, where marine in situ brGDGTs dominate. TEX₈₆ and MBT'_{5me} records reveal the SST and MAAT evolution of the region, respectively, within the range of independent existing paleotemperature reconstructions from the same samples. A temperature offset between the SST and MAAT proxies remains poorly reconciled with the coastal proximity of the site but is likely a combined effect of evolutionary trends in MBT'_{5me}-MAAT relationships and a disparate integration of climate signals by both proxies. The subdued air temperature cooling in the middle Eocene compared to the SST cooling could in part reflect progressive northward tectonic drift of the hinterland, while SSTs responded to stronger progressive cooling of southern-sourced surface waters. Strong MAAT cooling occurred step-wise at 35.5 Ma and around the Eocene–Oligocene transition. The absence of SST cooling around the Eocene–Oligocene transition seems to suggest some disconnection between oceanographic reorganisations and regional MAAT changes. The absence of a MAAT response at the PETM and MECO remains unresolved but might be related to shifting sources of brGDGTs during these warm phases.

The relatively high IR and low #rings_{tetra} confirm inferences from dinocyst assemblages of an inner mid-shelf depositional setting. We reconstruct strong river-runoff in the mid-Paleocene–earliest Eocene at this site, and normal open-marine conditions thereafter. The latter reflects increased seasonality in precipitation in the catchment as the continent tectonically migrates northwards during the middle and late Eocene, out of the region of year-round precipitation. The correlation between SST changes, dinoflagellate cyst assemblages and biodiversity metrics suggest temperature-controlled sea level changes influenced the site, probably through steric effects. We find corroborating evidence for a temperature signal in brGMGT assemblages, further demonstrating the potential to develop novel proxies with more extensive studies of modern distributions.

7. Supplements

- Table S1: Age tie points
- Table S2: grouping of dinocysts in eco-groups

8. Data availability

865 Raw GDGT/GMGT and palynological data presented in this paper, as well as R markdown code for data analysis and visualization can be found on Github (<https://github.com/bijlpeter83/RGDGT.git>). Data was deposited at zenodo under DOI:10.5281/zenodo.4471204.

9. Author contributions

AS (PETM) and PKB (part) prepared samples for GDGT analyses. PKB (part), AS (PETM) and CB (part) integrated
870 UHPLC-MS results. AS (PETM) and PKB (part) analysed palynological samples. All authors contributed to the interpretation of the OG results, and PKB, AS, JF and MC interpreted dinocyst results. PKB wrote the R script for data analyses and visualization, drafted the figures and wrote the paper with input from all authors.

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