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Southern high-latitude terrestrial climate change during the Paleocene–Eocene derived from a marine pollen record (ODP Site 1172, East Tasman Plateau)

L. Contreras^{1,2}, J. Pross^{2,3}, P. K. Bijl⁴, R. B. O'Hara², J. I. Raine⁵, A. Sluijs⁴, and H. Brinkhuis^{4,6}

¹Paleoenvironmental Dynamics Group, Institute of Geosciences, Goethe University Frankfurt, Altenhöferallee 1, 60438 Frankfurt, Germany

²Biodiversity and Climate Research Centre, Senckenberganlage 25,

60325 Frankfurt, Germany

³Paleoenvironmental Dynamics Group, Institute of Geosciences, Heidelberg University, Im Neuenheimer Feld 234, 69120 Heidelberg, Germany

⁴Department of Earth Sciences, Faculty of Geosciences, Utrecht University, Laboratory of Palaeobotany and Palynology, Budapestlaan 4, 3584 CD Utrecht, the Netherlands

⁵Department of Palaeontology, GNS Science, P.O. Box 30368, Lower Hutt 5040, New Zealand ⁶NIOZ Royal Netherlands Institute for Sea Research, P.O. Box 59, 1790 AB Den Burg, Texel, the Netherlands





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Abstract

Reconstructing the early Paleogene climate dynamics of terrestrial settings in the high southern latitudes is important to assess the role of high-latitude physical and biogeochemical processes in the global climate system. However, whereas a number of high-quality Paleogene climate records has become available for the marine realm of the high southern latitudes over the recent past, the long-term evolution of coeval terrestrial climates and ecosystems is yet poorly known. We here explore the climate and vegetation dynamics on Tasmania from the middle Paleocene to the early Eocene

- (60.7–54.2 Ma) based on a sporomorph record from Ocean Drilling Program (ODP)
 Site 1172 on the East Tasman Plateau. Our results show that three distinctly different vegetation types thrived on Tasmania under a high-precipitation regime during the middle Paleocene to early Eocene, with each type representing different temperature conditions: (i) warm-temperate forests dominated by gymnosperms that were dominant during the middle and late Paleocene; (ii) cool-temperate forests dominated by south-
- ern beech (*Nothofagus*) and araucarians across the middle/late Paleocene transition interval (~ 59.5 to ~ 59.0 Ma); and (iii) paratropical forests rich in ferns that were established during and in the wake of the Paleocene–Eocene Thermal Maximum (PETM). The transient establishment of cool-temperate forests lacking any frost-sensitive elements (i.e., palms and cycads) across the middle/late Paleocene transition interval
- indicates markedly cooler conditions, with the occurrence of frosts in winter, on Tasmania during that time. The integration of our sporomorph data with previously published TEX₈₆-based sea-surface temperatures from ODP Site 1172 documents that the vegetation dynamics on Tasmania were closely linked with the temperature evolution in the Tasman sector of the Southwest Pacific region. Moreover, the comparison of our season-specific climate estimates for the sporomorph assemblages from ODP Site 1172 with the TEX^L₈₆- and TEX^H₈₆-based temperature data suggests a warm-season bias of both calibrations for the early Paleogene of the high southern latitudes.





1 Introduction

The Southern Ocean is an important region for early Cenozoic (65–34 Ma) climates, being the dominant region for deep-water formation during that time (Thomas et al., 2003; Bijl et al., 2009; Sijp et al., 2011; Hollis et al., 2012). South Pacific sea-surface
and global intermediate water temperatures increased from the late Paleocene to the early Eocene, with maximum warmth during the Early Eocene Climatic Optimum (EECO; 53–51 Ma), followed by a cooling trend during the middle and late Eocene (Zachos et al., 2001, 2008; Bijl et al., 2009; Hollis et al., 2012). This cooling trend ultimately culminated in the establishment of a continental-scale ice shield on Antarctica during the earliest Oligocene (e.g., Zachos et al., 1994, 2008; Barrett, 1996), which represented a decisive step in the Earth's transition from a "greenhouse" into an "icehouse" world.

Organic geochemical surface-water temperature proxy records from the high-latitude Southwest Pacific Ocean (notably TEX_{86} ; Bijl et al., 2009) closely mirror trends in

- the benthic foraminiferal oxygen isotope data from the late Paleocene to the early Oligocene (Zachos et al., 2001, 2008) which lends further support to the suggestions that the southern ocean was the main region for deep-water formation during that time. Irrespective of the calibration used, Southwest Pacific TEX₈₆-derived sea-surface temperatures (SSTs) were relatively cool during the early and middle Paleocene. During
- the late Paleocene and early Eocene, SSTs gradually rose to tropical values (> 26 °C), with maxima being reached during the Paleocene-Eocene Thermal Maximum (PETM; Sluijs et al., 2011) and the EECO (Bijl et al., 2009; Hollis et al., 2009, 2012). Towards the end of the early Eocene (49–50 Ma), a pronounced SST cooling of ~ 4 °C occurred on the Australo-Antarctic margin; this cooling has been attributed to the onset of west-
- ²⁵ bound surface-water throughflow across the Tasmanian Gateway (Bijl et al., 2013a). A similar cooling trend is registered for surface waters off New Zealand (Hollis et al., 2009, 2012) and in the Tasman sector of the Southwest Pacific Ocean (Bijl et al., 2009; Hollis et al., 2012). Strikingly low SSTs are recorded for the high-latitude Southwest





Pacific Ocean during the interval spanning the middle/late Paleocene transition (59.5–59.0 Ma); TEX₈₆-derived SST decreased by ~ 3°C during that time (Bijl et al., 2009; Hollis et al., 2012; 2014). Along with this SST drop, lowered sea level and marked bathyal erosion suggest that a transient growth of an Antarctic ice sheet may have occurred (Hollis et al., 2014).

While an increasing amount of data has become available on the marine climate evolution in the southern high latitudes during the early Paleogene, the coeval terrestrial climate dynamics of that region are yet poorly documented (e.g., Passchier et al., 2013). For the Paleocene, paleobotanical records reflect the thriving of temperate, gymnosperm-rich forests dominated by podocarps in Southeast Australia (e.g., Macphail et al., 1994; Greenwood et al., 2003; Greenwood and Christophel, 2005), on Seymour Island (Antarctica; Askin, 1990), and in New Zealand (Mildenhall, 1980; Raine et al., 2009).

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In contrast, highly diverse angiosperm forests containing taxa that today are re-¹⁵ stricted to tropical environments characterize the early Eocene vegetation in the higherlatitude Southwest Pacific region. This vegetation is widely known from Southeast Australia (Macphail et al., 1994; Greenwood et al., 2003; Greenwood and Christophel, 2005), Tasmania (Truswell, 1997; Carpenter et al., 2012) and New Zealand (Raine et al., 2009; Handley et al., 2011); notably, it also thrived on the Wilkes Land margin of the Antarctic continent, i.e., at paleo-latitudes of ~ 70° S (Pross et al., 2012; Contreras et al., 2013).

The pronounced vegetation turnover from temperate forests during the Paleocene to near-tropical forests during the early Eocene suggests a marked, climatically driven change in terrestrial environments in the high southern latitudes. However, the tran-

sitional process between these two vegetation types and the underlying change in terrestrial climate conditions of the Southwest Pacific region have remained poorly understood. The gaps in the documentation of terrestrial climate dynamics during the early Paleogene appear particularly pronounced considering the coeval datasets for the marine realm from the same region (Bijl et al., 2009, 2013a; Hollis et al., 2009,





2012, 2014). This is due to the fact that prior paleobotanical studies are mainly based on stratigraphically discontinuous outcrops that provided only limited insights into the Paleocene and early Eocene vegetation of the region (Greenwood et al., 2003). In addition, available studies on sporomorphs from the early Paleogene of the Southwest

- Pacific region have predominantly focussed on the taxonomical characterization of the assemblages and the generation of biostratigraphic schemes (e.g., Stover and Evans, 1973; Stover and Partridge, 1973; Truswell, 1997; Macphail, 1999). In any case, estimates of terrestrial temperatures in the high southern latitudes and a comparison with the currently available SST estimates from the marine realm can strongly enhance the understanding of the climate evolution in the high southern latitudes during the carky.
- ¹⁰ understanding of the climate evolution in the high southern latitudes during the early Paleogene.

In light of the above, we here explore the vegetation response to high-southernlatitude climate forcing from the Paleocene to the "hothouse" conditions of the early Eocene based on a new, chronostratigraphically well-calibrated (Bijl et al., 2013b)

- sporomorph record from Ocean Drilling Program (ODP) Site 1172 off eastern Tasmania. We quantitatively evaluate the compositional variations of the sporomorph assemblages and carry out quantitative sporomorph-based paleoclimatic reconstructions. For a further assessment of the terrestrial climatic conditions in the Southwest Pacific region, we apply the same paleoclimate reconstruction approach to previously published
- ²⁰ sporomorph records from Southeast Australia (Bass Basin, Gippsland Basin, Southeast Highlands) and New Zealand. Finally, we compare our terrestrial paleoclimate estimates with previously published TEX^L₈₆- and TEX^H₈₆-based SST reconstructions from the same site (Bijl et al., 2009, 2013b; Hollis et al., 2014) in order to contribute to a better understanding of the early Paleogene climate dynamics in the high southern latitudes.





2 Material and methods

The middle Paleocene to early Eocene strata studied here were recovered at ODP Site 1172, which is located ~ 100 km east of Tasmania on the western side of the East Tasman Plateau (ETP; 43°57.6′ S, 149°55.7′ E; Fig. 1) (Shipboard Scientific Party, 2001a).

2.1 Regional setting and paleoceanography

During the early Paleogene, Tasmania and the ETP were located at $\sim 65^{\circ}$ S, much closer to Antarctica than today (Exon et al., 2004b). Paleoceanographic patterns as determined by winds and gateway configuration were likely vital for regional climates on land (Sijp et al., 2011). The study site was located close to the Tasmanian promon-10 tory, which hampered deep ocean exchange between the Southwest Pacific and the Australo-Antarctic Gulf for most of the early Paleogene (Exon et al., 2001; Stickley et al., 2004; Fig. 1). During the Paleocene and early Eocene, the Tasman region was under the persistent influence of the Antarctic-derived Tasman Current; in contrast, the Australo-Antarctic Gulf (west of Tasmania) was bathed by the low-latitude-derived 15 Proto-Leeuwin Current (Huber et al., 2004; Sijp et al., 2011; see Fig. 1). This paleoceanographic configuration determined marine biogeographical patterns in the region (Huber et al., 2004; Bijl et al., 2011, 2013a). The onset of the deepening of the Tasmanian Gateway at ~ 49–50 Ma initiated a westbound Antarctic Counter Current flowing along the Antarctic margin from the Pacific into the Australo-Antarctic Gulf (Bijl et al., 20 2013a). Continued rifting through the Eocene and accelerated deepening of the Tasmanian Gateway (~ 35.5 Ma) led to the inflow of Australo-Antarctic Gulf waters through the Tasmanian Gateway into the southern Pacific during the early Oligocene (Stick-

ley et al., 2004; Sijp et al., 2011). It could be expected that these paleoceanographic reorganizations affected regional climates throughout the early Paleogene.





2.2 Age model, lithology and depositional environment

We here follow the corrected sample depths of 1172D published in Sluijs et al. (2011) through detailed correlation of the X-ray fractionation core scanning to the *γ*-ray downhole log. The age model of the studied sequence is based on the magnetostratigraphy, chemostratigraphy and dinoflagellate cyst (dinocyst) biostratigraphy as presented in Bijl et al. (2013b). For the interval studied (60.7–54.2 Ma), the age model is based on 3 magnetic reversals, confidently correlated to the Geomagnetic Polarity Time Scale (GPTS) of Vandenberghe et al. (2012), the PETM (~ 56 Ma), which exhibits a negative carbon isotope excursion of ~ 3‰ between 611.89 and 611.86 rmbsf (revised meters below sea floor) (Sluijs et al., 2011), and several dinocyst first and last occurrence data that have been calibrated regionally following Crouch et al. (2014). The magnetostratigraphic age model for the Paleocene section (Röhl et al., 2004) was adjusted by Bijl et al. (2010) on the basis of the recognition of the PETM in Core 1172D-15R (611.8 rmbsf; Sluijs et al., 2011). The missing interval between Cores 16R and 17R

¹⁵ (~ 620 rmbsf) represents a ~ 1.3 Ma-long hiatus that spans the time interval correlative to between infra-Subchrons C26n and C25n (57.7 to 59 Ma; Bijl et al., 2010, 2013b).

The upper 60 cm of section in Core 17R (i.e, below the hiatus) are heavily disturbed and have many sediment characteristics more consistent with Core 16R than with the underlying sediment (Röhl et al., 2004). We therefore follow Röhl et al. (2004) in their suggestion that this interval represents caved material of late Paleocene age.

The Middle Paleocene to Lower Eocene succession of ODP Site 1172 consists mainly of gray to grayish brown clay- and siltstones with low abundances of calcareous and siliceous microfossils (Shipboard Scientific Party, 2001b; Röhl et al., 2004). Environmentally, the succession is interpreted to reflect very shallow to restricted marine conditions, with marked runoff from the nearby shores (Röhl et al., 2004).

Any study on sporomorphs from marine sediments critically relies on the identification of the source region in order to provide meaningful paleoclimatic information. An

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in-depth discussion of this issue is provided in Sect. 3.2 ("Constraints on sporomorph source region").

2.3 Sample processing and data analysis

Eighty-nine samples from the Paleocene and Eocene of ODP Site 1172 were processed at the Laboratory of Palaeobotany and Palynology (Bijl et al., 2011) and were reanalysed for terrestrial palynomorphs. Sample processing followed standard palynological techniques, including treatment with HCI (10%) and HF (38%) and sieving through a 15 µm nylon mesh (e.g., Pross, 2001). The microscope slides are stored in the collection of the Laboratory of Palaeobotany and Palynology at Utrecht University. Whenever possible, 300 sporomorphs (excluding reworked specimens) were analysed per sample and determined to the species level; this required the analysis of up to six slides per sample. The analyses were performed using a light microscope at

- 200× magnification; morphological details were studied with a magnification of 1000×. Sporomorph identifications are mainly based on Couper (1960), Harris (1965), Stover
- and Partridge (1973), Truswell (1983), Raine et al. (2008), and Truswell and Macphail (2009). The botanical affinities are given following Macphail et al. (1994), Raine (1998), and Truswell and Macphail (2009). All sporomorph data (including photomicrographs of key taxa) are provided in the Supplement.

Rarefaction was applied to evaluate sporomorph diversity; this allows to estimate the number of sporomorph species at a constant sample size (Raup, 1975). Detrended Correspondence Analysis (DCA) is an ordination technique that was used to assess the overall variation in floral composition through time.

To constrain the source region of the sporomorphs from ODP Site 1172, we compared the floristic composition of our record with coeval sporomorph records from

Southeast Australia (Bass Basin – data from Blevin, 2003; Gippsland Basin – Department of Primary Industries, 1999; Southeast Highland sections – Taylor et al., 1990) and New Zealand (Middle Waipara and Otaio River sections – Pancost et al., 2014) using DCA (see Fig. 1 for locations and Table 1 for further details and references).



From all records derived from drillings, only data from core or sidewall core samples were evaluated to avoid potential contamination by caving. To exclude a bias as it could have been introduced into our comparison through different taxonomic criteria, we have standardized all datasets using broader groups of taxa (e.g., *Gleicheniidites* spp./*Clavifera* spp., *Phyllocladidites* spp., *Nothofagidites* spp. (*fusca* group)) when necessary. Due to the differences in sample sizes, sporomorph percentages were used as input for the DCA, and only samples with counts \geq 100 individuals were evaluated.

2.4 Sporomorph-based climate reconstructions

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Quantitative sporomorph-based climate estimates were carried out following the bioclimatic analysis approach of Greenwood et al. (2005). For each taxon with a known nearest living relative (NRL; Table 2), climatic profiles were generated with regard to mean annual temperature (MAT), coldest month mean temperature (CMMT), warmest month mean temperature (WMMT), and mean annual precipitation (MAP). The climate profiles are derived from (i) the dataset of Pross et al. (2012), which is mainly based on distribution data from the Australian National Herbarium online database (Australian National Herbarium, 2011) and the mathematical climate surface software ANUCLIM 5.156 (Houlder et al., 1999), and (ii) the PALAEOFLORA database, which contains climatic information for a plant taxon based on its global distribution (Utescher and Mosbrugger, 2013). Following Greenwood et al. (2005), the climatic values for each

- assemblage were calculated based on the zone of overlap of the majority of taxa from that assemblage with respect to a given climate parameter. This overlap interval was calculated using the 10th percentile (as lower limit) and 90th percentile (as upper limit) of the total range of the NLRs recorded in that assemblage. The climate estimate is given as the midpoint between the lower and upper limits, with the error spanning from
- the lower to the upper limit. Only samples with counts ≥ 100 individuals were used in the climate reconstructions.





2.5 Statistical examination of the connection between floristic composition and temperature

To examine the correlation between the floristic composition of our sporomorph record from ODP Site 1172 (as represented by DCA Axis 1 sample scores; Figs. 2 and 4) and TEX^L₈₆- and TEX^H₈₆-derived SST values from the same site (Bijl et al., 2009, 2013b; Hollis et al., 2014), we applied a state space model. State space models or dynamic linear models allow data distributed along time to be interpreted as the combination of several components, such as trend, seasonal or regressive components (Petris et al., 2009; see West and Harrison, 1997, for further details on the advantages and devel-

¹⁰ opment of the method).

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We compared 60 SST data points with our 40 values of the DCA Axis 1 scores. Because only six of the SST data points and DCA Axis 1 sample scores are from the exact same depths, we multiplied imputed values of DCA Axis 1 sample scores and SST; the imputed values are based on Eqs. (1) and (2) below. The cross-covariance between two points was calculated following Eq. (3) and the final correlation using Eq. (4). When $X_{d(i)}$ (1) is the "true" SST and $X_{d(i)}$ (2) is the "true" Axis 1 sample score, σ is the standard deviation, ρ is the correlation between the two variables and τ is the Kendall rank correlation coefficient. For Eqs. (1) and (2) we assumed that both variables are normally distributed (*N*).

²⁰ SST_i ~ N
$$\left(x_{SST}(t(i)), \tau_{SST}^{2}\right)$$

DCA_i ~ N $\left(x_{DCA}(d(i)), \tau_{DCA}^{2}\right)$
 $\Delta d \begin{pmatrix} \sigma_{11}^{2} & \rho \sigma_{11} \sigma_{22} \\ \rho \sigma_{11} & \sigma_{22} \end{pmatrix} + \begin{pmatrix} \sigma_{SST}^{2} & 0 \\ 0 & \sigma_{DCA}^{2} \end{pmatrix}$
corr(SST, DCA) = $\frac{\Delta t \rho \sigma_{DCA} \sigma_{SST}}{\sqrt{\left(\Delta dt + \tau_{DCA}^{2}\right) \left(\Delta t \sigma_{SST}^{2} + \tau_{SST}^{2}\right)}}$
301



(1)

(2)

(3)

(4)

The model was fitted using OpenBUGS run through the BRugs package (Thomas et al., 2006) of the R software for statistical computing (R Development Core Team, 2011) fitted in a Bayesian framework using the BRugs package. The R script and further explanation of the analysis is given in the Supplement.

5 3 Results

3.1 Sporomorph results from ODP Site 1172

Of the 89 palynological samples processed from the Middle Paleocene to Lower Eccene of ODP Site 1172, 40 samples yielded sporomorph counts ≥ 100 individuals and were further used in our analyses. The preservation of sporomorphs is generally good. A total of 197 sporomorph types were identified. A range chart with the 10 relative abundances of key taxa is presented in Fig. 2; a full account of the identified taxa and their abundance data along with photomicrographs of key taxa is provided in the Supplement. Based on rarefied values, the entire study interval is characterized by rich sporomorph assemblages (mean \pm s.d. = 29.2 \pm 3.4 taxa/sample at 100 individuals, n = 40). Remarkably low sporomorph species numbers are recorded for the 15 samples corresponding to the PETM (22.8 and 24.5 species/sample at 100 individuals; Fig. 2). The DCA results of our sporomorph record yield distinctly different values for the Axis 1 and Axis 2 sample scores (Fig. 3a). They allow to define three sample groups, with each sample group being characteristic for specific time intervals of our record (Fig. 3a). These intervals are: (i) the middle and late Paleocene; (ii) the mid-20 dle/late Paleocene transition; and (iii) the early Eocene including the PETM. All three sample groups comprise characteristic sporomorph assemblages that are portrayed in the following.





3.1.1 Middle Paleocene (60.7–59.5 Ma) and late Paleocene (59.0–55.6 Ma) intervals

The sporomorph assemblages from the middle and late Paleocene intervals (excluding the middle/late Paleocene transition, see below) are represented by 20 samples. They

- ⁵ are dominated by gymnosperm pollen, which on average accounts for 45 % of all sporomorphs. The gymnosperm pollen is represented mainly, in the order of decreasing abundances, by *Podocarpidites* spp. (botanical affinity: podocarpaceae; podocarps), *Dilwynites granulatus (Wollemia* (Wollemi pine)/*Agathis* (Kauri); Macphail et al., 2013), *Phyllocladidites mawsonii (Lagarostrobos franklinii;* Huon pine), and *Araucariacites* spp. (Araucariaceae: *Agathis* (Kauri) and *Araucaria*) (Fig. 2). Other abundant sporo-
- spp. (Araucariaceae; Agathis (Kauri) and Araucaria) (Fig. 2). Other abundant sporomorphs are, in the order of decreasing abundances, Cyathidites spp. (probably Cyatheaceae), Nothofagidites spp. (Nothofagus; southern beech), Gambierina spp., Gleicheniidites spp. (Gleicheniaceae), and Ceratosporites spp. (Lycopodiaceae, Selaginellaceae).
- ¹⁵ Although the sporomorph assemblages from the middle to late Paleocene have very similar floristic compositions, differences exist with regard to the percentages of some taxa (Fig. 2). For instance, the early part of the middle Paleocene exhibits relatively high (~ 30 %) percentage of podocarpaceous pollen (mainly *Podocarpidites* spp. (Podocarpaceae; podocarps) and *Phyllocladidites mawsonii* (*Lagarostrobos franklinii*;
- ²⁰ Huon pine)). In contrast, the latest Paleocene is characterized by lower abundances of these taxa (~ 14%), but exhibits higher percentages of *Cyathidites* spp. (~ 18%) and the constant presence of *Myricipites harrisii* (probably Casuarinaceae; sheoak). Remarkably, both the middle and late Paleocene are characterized by the presence of *Arecipites* spp. (Arecaceae; palms) and *Cycadopites* spp. (Cycadales).

25 3.1.2 Middle/late Paleocene transition interval (59.5–59.0 Ma)

The sporomorph assemblages of the middle/late Paleocene transition interval are represented by 10 samples. They exhibit higher percentages of *Araucariacites*





spp. (Araucariaceae; ~19%), Nothofagidites spp. (Nothofagus; ~13%), and Ceratosporites spp. (Lycopodiaceae, Selaginellaceae; $\sim 6\%$) than the assemblages from the under- and overlying strata of middle respectively late Paleocene age (Fig. 2). Other abundant taxa are, in the order of decreasing abundances, *Cyathidites* spp. (Cyatheaceae), Dilwynites granulatus (Wollemia/Agathis) and Gleicheniidites spp. 5 (Gleicheniaceae). The percentages of podocarpaceous types (*Podocarpidites* spp. (Podocarpaceae) and Phyllocladidites mawsonii (Lagarostrobos franklinii)) are lower $(\sim 10\%)$ than in assemblages from the under- and overlying middle and late Paleocene. Notably, the assemblages from the middle/late Paleocene transition interval are devoid of Arecipites spp. (Arecaceae; palms) and Cycadopites spp. (Cycadales) pollen.

3.1.3 PETM and early Eocene interval (55.6–54.2 Ma)

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Assemblages from the PETM are documented in 2 samples with counts > 100 individuals. They are characterized by high percentages (up to 39%) of Dilwynites granulatus (Wollemia/Agathis). Other abundant taxa are, in the order of decreasing abundances,

- Cyathidites spp. (Cyatheaceae), Araucariacites spp. (Araucariaceae), and Gleicheni-15 idites spp. (Gleicheniaceae). The assemblages of the PETM interval are further characterized by the presence of *Malvacipollis* spp. (Euphorbiaceae, probably Eumalvoideae) and Myricipites harrisii (probably Casuarinaceae). Findings of Spinizonocolpites prominatus (Nypa palm) are restricted to the PETM and the earliest Eocene (Fig. 2).
- Sporomorph assemblages of the early Eocene interval are documented in 8 sam-20 ples. They show in general very high percentages (mean: 20%) of *Cyathidites* spp. (probably Cyatheaceae; Fig. 2). Other abundant taxa are, in the order of decreasing abundances, Dilwynites granulatus (Wollemia/Agathis), Araucariacites spp. (Araucariaceae), Podocarpidites spp. (Podocarpaceae), and Phyllocladidites mawsonii (Lagarostrobos franklinii). The early Eocene interval is also characterized by relatively high 25
- percentages of Proteacidites spp. (~7%) and the constant presence of Malvacipollis spp. (Euphorbiaceae, probably Eumalvoideae), Myricipites harrisii (probably Casuarinaceae), Arecipites spp. (Arecaceae; palms), and Cycadopites spp. (Cycadales).





3.2 Constraints on sporomorph source region

Several lines of evidence suggest that eastern Tasmania was the main source of the sporomorphs encountered in the Middle Paleocene to Lower Eocene of ODP Site 1172. The distance of this site to the paleo-shoreline of eastern Tasmania during the Pale-

- ⁵ ocene early Eocene was on the order of ~ 100 km, whereas the minimum distance to George V Land (Antarctica) amounted to ~ 500 km (Fig. 1). A Tasmanian source is further suggested based on the distribution pattern of reworked Permian and Triassic sporomorphs. The assemblages from ODP Site 1172 are characterized by the constant presence of elements reworked from Permian and Triassic strata (e.g., *Cannanoropollis*)
- spp., Protohaploxypinus spp., Alisporites spp.; see Supplement, Plate II); the percentages of reworked sporomorphs reach up to 16% of the total assemblages. A similar input of reworked Permian and Triassic material is known for sporomorph assemblages from Paleocene–Eocene strata along the Australo-Antarctic Gulf (Otway Basin; Harris, 1965) and from the Eocene of the Wilkes Land margin (Contreras et al., 2013). In
- the Tasmania region, reworked Permian and Triassic sporomorphs are recorded in the Paleocene–Eocene of the Bass Basin (Partridge et al., 2003) as well as in Eocene strata off western Tasmania and on the South Tasman Rise (Truswell, 1997). This pattern is consistent with the fact that sporomorph-bearing sediments of Permian and Triassic age occur in several regions of Tasmania (e.g., Playford, 1965; Truswell, 1978;
- ²⁰ Calver et al., 1984). In contrast, reworked sporomorphs on the continental shelf off George V Land (Antarctica; Fig. 1) comprise only taxa with Cretaceous and Cenozoic ages (Truswell, 1983). A similar picture emerges for the Cenozoic of the Gippsland Basin where Permian and Triassic sporomorphs occur only sporadically and in low numbers (see reports in Department of Primary Industries, 1999).
- ²⁵ A further constraint on the source of the sporomorphs at ODP Site 1172 comes from the DCA-based comparison of the floristic composition of the Site 1172 record with other coeval Paleocene–Eocene sporomorph records from Southeast Australia (Bass and Gippsland Basins, Southeast Highlands) and New Zealand (Middle Waipara





and Otaio River sections) (Fig. 3b; see Fig. 1 and Table 1 for site locations and details on records). It suggests marked differences in floristic compositions between most records, a result that can be further corroborated for the Bass and Gippsland Basins based on carbon-isotope data from both basins. The Paleocene–Eocene successions

- of these basins exhibit distinctly different bulk carbon isotope values, which is interpreted to reflect the signal from different plant communities contributing to the organic carbon input into the basins during that time (Boreham et al., 2003). However, despite the floristic differences between the records analysed, there is a remarkable similarity in DCA scores between the records from ODP Site 1172 and the Bass Basin (Fig. 3b),
- ¹⁰ which is located mainly on the shelf off northern Tasmania, but also extends into Northeast Tasmania (Moore et al., 1984). This similarity further corroborates the scenario of a Tasmanian source for the ODP Site 1172 sporomorph assemblages.

3.3 Sporomorph-based paleoclimate estimates

The paleoclimate estimates derived from the sporomorph assemblages from ODP Site 1172 are presented in Fig. 4 and summarized in Table 3. For the middle Paleocene (60.7–59.5 Ma), all reconstructed temperature parameters yield relatively cool values (MAT ≃12 °C; CMMT ≃7 °C; WMMT ≃18 °C). A pronounced further cooling is documented in all temperature parameters for the middle/late Paleocene transition interval (59.5–59.0 Ma). The reconstructed values represent the lowest temperatures of the active recent temperature parameters for the middle/late Paleocene transition in-

- the entire record; typical values are ~ 10°C for MAT, ~ 5°C for CMMT and ~ 16°C for WMMT (Fig. 4). Markedly higher temperatures prevailed during the late Paleocene (59.0–55.6 Ma) and early Eocene (55.6–54.2 Ma), with estimates for that interval being on the order of ~ 15°C for MAT, ~ 8°C for CMMT and ~ 20°C for WMMT. The highest temperatures of the entire study interval are recorded for the Paleocene/Eocene transition interval (including the DETM). MAT CMMT and WMMT reached values of _ 22°C
- tion interval (including the PETM); MAT, CMMT, and WMMT reached values of \sim 22 °C,
 - \sim 18 °C and \sim 26 °C, respectively, during that time (Fig. 4).

The sporomorph-based MAP estimates yield high, near-constant values throughout the sequence studied, with an average value of $\sim 180 \,\text{cm}\,\text{yr}^{-1}$ (Fig. 4).





4 Interpretation

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4.1 Floristic and climatic evolution

Based on our qualitative and quantitative results from the sporomorph record from ODP Site 1172, three main vegetation types prevailed on Tasmania from the middle Pale ocene to early Eocene. These vegetation types (i) exhibit different floristic compositions based on the DCA results (Figs. 2 and 3a), (ii) show similar diversities based on the rarefaction results (Fig. 2), and (iii) represent specific climatic conditions based on our sporomorph-derived paleoclimatic reconstructions (Fig. 4). The floristic characteristics and climatic requirements of all three vegetation groups are discussed in the following sections.

4.1.1 Middle Paleocene (60.7–59.5 Ma) and late Paleocene (59.0–55.6 Ma) intervals

During both time intervals, the flora of Tasmania was characterized by gymnospermrich forests dominated by podocarps; Araucariaceae, ferns, protealeans and *Nothofa*-

- ¹⁵ gus were further important components of the vegetation. Arecaceae (palms) and Cycadales were present during both time intervals. Although the middle Paleocene and late Paleocene forests exhibited a strongly similar composition, important differences existed. Podocarps reached a markedly stronger dominance during the middle Paleocene than during the late Paleocene, whereas the late Paleocene was characterized
- ²⁰ by higher abundances of ferns (mainly Cyatheaceae) and the presence of Casuarinaceae (Fig. 2).

With regard to modern analogues, these forests bear resemblance to the extant warm-temperate, Podocarpaceae-dominated forests of Southeast Australia and New Zealand (see, e.g., Kershaw, 1988). Based on the structure of extant temperate forests from southern latitudes (Kershaw, 1988; Enright and Hill, 1995; Veblen et al., 1996; Reid et al., 1999), the overstorey of Tasmanian forests during the middle and late





Paleocene was dominated by Podocarpaceae and, to a somewhat lesser extent, Araucariaceae and *Nothofagus*. The understorey, in turn, was likely dominated by ferns, with Cyatheaceae being particularly abundant during the late Paleocene. Considering the habitats of their extant representatives, Arecaceae (palms), Cycadales, Proteaceae, and Casuarinaceae formed parts of both the over- and understorey (compare, Johnson

and Casuarinaceae formed parts of both the over- and understorey (compare Johnson and Wilson, 1993; Hill, 1994; Morley, 2000; Jones, 2002).

Climatically, the presence of tree ferns (i.e., Cyatheaceae), Arecaceae and Cycadales implies mild climates with no or merely rare frost events. Owing to physiological constraints (including manoxylic wood, large, unprotected buds, soft, water-rich

tissues, and a near-absence of frost-"hardening" mechanisms), all these plants are unable to cope with sustained freezing (Sakai and Larcher, 1987; Wing and Greenwood, 1993); today, they only occur in settings with CMMT ≥ 5.5 °C (Greenwood and Wing, 1995; Utescher and Mosbrugger, 2013).

4.1.2 Middle/late Paleocene transition interval (59.5-~59.0 Ma)

- ¹⁵ Across the middle/late Paleocene boundary, the warm-temperate forests characterizing the vegetation on Tasmania during most of the middle and late Paleocene as described above were transiently replaced by cool-temperate forests dominated by *Nothofagus* (mainly *N. fusca* type) and Araucariaceae. Ferns, podocarps and protealeans were further prominent components of this vegetation. Palms (Arecaceae) and Cycadales, as
- ²⁰ they occurred both during the preceding part of the middle (60.7–59.5 Ma) and the subsequent part of the late Paleocene (59.0–55.6 Ma; see above), were absent (Fig. 2). In light of the age control and the temporal resolution of our record, these cool-temperate forests prevailed on Tasmania for ~ 0.5 Ma (based on the duration of elevated *Nothofagus* percentages) respectively ~ 0.3 Ma (based on the absence of tropical to subtrop-
- ical indicators such as palms and Cycadales). The 1.3 Myr-long hiatus precludes us from firmly determining the complete duration of the cold interval in the mid-Paleocene (Fig. 2), however from marine records from New Zealand we deduce a duration of 0.5 Ma (Waipara section and ODP Site 1121; Hollis et al., 2014).





Today, vegetation dominated by *Nothofagus* (*N. fusca* type) is typical of cooltemperate forests from southern Australia and New Zealand (Kershaw, 1988). Considering the structure of such forests (e.g., McGlone et al., 1996; Reid et al., 1999), the vegetation thriving during the middle/late Paleocene transition interval was character-

- ⁵ ized by a canopy dominated by *Nothofagus* and open understories dominated by ferns (mainly Cyatheaceae and the parent plants of *Ceratosporites* spp. (Selaginellaceae, Lycopodiaceae)). Araucariaceae were also present. Because extant members of Araucariaceae are tall trees generally confined to the lower mid-latitudes (Kershaw and Wagstaff, 2001), their presence appears at first sight incompatible with an occurrence
- of cool-temperate forests. However, members of the genus Araucaria also thrive in cold temperate forests in mountainous areas of Chile (Veblen, 1982) and can withstand frost events as cold as -15°C (Prentice et al., 1992), which supports our observation that Araucariaceae were a component of the cool-temperate forests of Tasmania across the middle/late Paleocene transition.
- Based on the overall floristic evidence, Tasmania witnessed a transient period of cooler conditions lasting from ~ 59.5 to ~ 59.0. In light of the frost sensitivity of Arecaceae, Cycadales and Cyatheaceae (compare Sect. 4.1.1), the decline or total absence of these taxa across the middle/late Paleocene transition (Fig. 2) suggests harsher winters (with particularly frequent and/or cold frost events) during that time.
- ²⁰ Such lower temperatures are corroborated by the comparison with coeval TEX^L₈₆- and TEX^H₈₆-based SST data from ODP Site 1172 (Bijl et al., 2009, 2013b; Hollis et al., 2014). For the interval from ~ 59.4 to ~ 59.0 Ma, they show the lowest values of the entire Paleocene–Eocene SST record (Fig. 4, Table 3).

4.1.3 PETM and early Eocene interval (55.6–54.2 Ma)

During the early Eocene, the composition of the forests on Tasmania was distinctly different from that of the temperate forests thriving during the Paleocene (Figs. 2 and 3a); the underlying floristic turnover coincides with the onset of the PETM (see DCA)





Axis 1 sample scores in Fig. 2). Based on our sporomorph data, the early Eocene vegetation was dominated by ferns and different angiosperms (mainly Proteaceae, Casuarinaceae and Euphorbiaceae/Eumalvoideae). Remarkably, taxa that were common during the Paleocene (e.g., *Ceratosporites* spp. (Lycopodiaceae, Selaginellaceae) and

- Gambierina spp. (extinct clade)) declined dramatically in abundance or disappeared completely during that time (Fig. 2). Because extant Selaginellaceae and Lycopodiaceae are cosmopolitan families (Jermy, 1990; Øllgaard, 1990), it is difficult to connect the disappearance of these taxa on Tasmania during the earliest Eocene with specific ecological and climatic conditions.
- ¹⁰ The coexistence of frost-tolerant (e.g., Araucariaceae, Podocarpaceae) and thermophilous taxa (e.g., Casuarinaceae, Arecaceae (palms)) suggests the presence of paratropical forests *sensu* Morley (2000). In particular, the occurrence of *Nypa* from the PETM onward into the early Eocene suggests the presence of tropical mangrove vegetation along the coast of Tasmania. A similar vegetation, also containing ther-
- ¹⁵ mophilous taxa such as *Nypa* and *Gymnostoma* (Casuarinaceae), is documented in early Eocene macrofloras from western Tasmania (Pole, 2007; Carpenter et al., 2012). Hence, forests on Tasmania during the early Eocene consisted temporarily of at least two vegetation associations: (i) a mangrove association characterized by *Nypa*, which is only recognized during the PETM and the earliest Eocene, and (ii) a paratropical
- association characterized by the coexistence of frost-tolerant taxa (i.e., Araucariaceae, Podocarpaceae and *Nothofagus*) and thermophilous elements such as palms and Casuarinaceae.

Considering the ecology of the nearest living relatives of the plants represented by the encountered sporomorphs, the overstorey vegetation during the early Eocene comprised taxa such as Araucariaceae, Podocarpaceae and *Nothofagus*, whereas the understorey was probably dominated by ferns (mainly Cyatheaceae). Members of the Proteaceae, Casuarinaceae, Cycadales, Arecaceae (palms), and Euphorbiaceae/Eumalvoideae may have been both components of the over- and the understorey (Johnson and Wilson, 1993; Hill, 1994; Morley, 2000; Jones, 2002).





The number of sporomorph species registered at ODP Site 1172 remained relatively constant from the middle Paleocene to the early Eocene (Fig. 2). This observation is in contrast to Southeast Australia, where sporomorph assemblages from non-marine and marginal marine settings (Partridge, 1976) exhibit a considerably higher diversity during

- the early Eocene than during the Paleocene (Macphail et al., 1994; see Sect. 4.2.2. below). The reasons behind this discrepancy may be sought in the particularly high sea level during the early Eocene as it is recorded regionally based on sedimentological and paleontological data from ODP Site 1172 (Exon et al., 2004a) and globally (Miller et al., 2005; Cramer et al., 2011). Owing to the selective nature of marine sporomorph
 transport as a function of transport distance (e.g., Moss et al., 2005), the higher sea
- level during the early Eocene than during the Paleocene potentially caused a diversity decrease of the sporomorph assemblages at ODP Site 1172 (see also below).

Our data suggest that the floristic change connected to the PETM is similar to that registered for the early Eocene. However, unravelling the exact anatomy of vegeta-

- tion change across the PETM at ODP Site 1172 is difficult due to the low sporomorph yields in the respective sediments at that site. In addition, the interpretation of the available data is hampered by the sea-level rise during the PETM (Sluijs et al., 2011); the transgression-induced change in depositional setting towards more distal conditions may have caused a bias in the composition and diversity patterns of sporomorph
- ²⁰ assemblages, with the resulting assemblages being skewed towards a dominance of easily transported sporomorphs (compare Traverse, 1994, 2008). In light of this bias, the high abundances (up to 39%; Fig. 2) of *Dilwynites granulatus (Wollemia/Agathis)* and the remarkably low diversities (Fig. 2) in the PETM samples from ODP Site 1172 likely represent a change in depositional setting rather than a true paleoecological
- signal. This interpretation is supported by the higher abundances of *Dilwynites granulatus* (~ 35 %) in early Paleogene sediments deposited in distal environments of the Bass Basin when compared to the markedly lower abundances (~ 10 %) in coeval sediments from nearshore settings in the same basin (see reports in Partridge et al., 2003). Nevertheless, our data show that the environmental perturbations connected



to the PETM had a profound impact on the vegetation on Tasmania. They lead to the extirpation of various ferns (e.g., *Perinomonoletes* spp., *Ceratosporites* spp. (Lycopodiaceae, Selaginellaceae)) and angiosperms (e.g., *Gambierina rudata, Nothofagidites* sp. 1 (*Nothofagus*)), and the appearance of new angiosperms mainly within the Proteaceae family (e.g., *Proteacidites grandis*).

4.2 Integration with other terrestrial vegetation records and temperature estimates from the southern high latitudes

Our results from ODP Site 1172 yield a ~ 6.5 Ma-long vegetation record for the Tasman sector of the SW Pacific region spanning from the middle Paleocene to the early

- Eocene. To obtain insights into the potential regional differentiation of terrestrial ecosystems and climates in the high southern latitudes during that time, we have integrated our data from ODP Site 1172 with the available information on terrestrial ecosystems and temperatures from other parts of the SW Pacific domain. Our integration is augmented by newly generated temperature estimates for previously published sporo-
- ¹⁵ morph records from Southeast Australia and New Zealand (see Tables 1 and 3 for further information on records evaluated and results). Because other continuous, stratigraphically well-calibrated vegetation records across the middle/late Paleocene transition interval are not yet available for the SW Pacific region, we focus our comparison on the middle Paleocene, late Paleocene and early Eocene.

20 4.2.1 Middle and late Paleocene

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Paleobotanical records for the middle and late Paleocene are well known from Southeast Australia (e.g., Bass Basin – Macphail et al., 1994; Blevin, 2003, Gippsland Basin – Stover and Partridge, 1973; Macphail et al., 1994; Department of Primary Industries, 1999). In summary, the middle and late Paleocene vegetation in this region consisted predominantly of warm temperate forests that were characterized by podocarps.

²⁵ sisted predominantly of warm temperate forests that were characterized by podocarps, Araucariaceae and ferns; angiosperms were represented mainly by Proteaceae (e.g.,





Taylor et al., 1990; Macphail et al., 1994; Greenwood et al., 2003; Greenwood and Christophel, 2005). Similar warm temperate forests dominated by podocarps and Araucariaceae, and with a strong contribution of Proteaceae, thrived on New Zealand (Mildenhall, 1980; Raine et al., 2009).

- Based on our results, Podocarpaceae together with Araucariaceae, Cyatheaceae and Proteaceae were also the prevailing group of plants during the middle and late Paleocene on Tasmania (Fig. 2; compare also Sect. 4.1.1). However, important floristic differences existed between Southeast Australia, New Zealand and Tasmania (Fig. 3b); they are mainly based on the restriction of certain sporomorph taxa to specific regions
- (e.g., Liliacidites spp., Cibotiidites tuberculiformis New Zealand; Ilexpollenites spp. Southeast Australia [Bass and Gippsland Basins, Southeast Highlands]; Tripunctisporis maastrichtiensis – New Zealand, Bass Basin and Tasmania). Despite the differences in floristic composition, the remarkable dominance of Podocarpaceae and Araucariaceae in Southeast Australia, Tasmania and New Zealand suggests that warm-temperate
- ¹⁵ forests dominated by gymnosperms were the prevalent vegetation type in the Southwest Pacific region during the middle and late Paleocene. Moreover, thermophilous taxa such as palms, Olacaceae (*Anacolosa*) and Cupanieae first appeared and/or increased significantly in abundance and diversity during the latest Paleocene in Southeast Australia (Kemp, 1978; Macphail et al., 1994). On New Zealand, typical tropical
- taxa (e.g., Cupanieae, Austrobuxus (Euphorbiaceae), Nypa) also begin to appear during the latest Paleocene prior to the PETM (Crouch and Brinkhuis, 2005; Raine et al., 2009). Hence, the arrival of thermophilous elements indicates the onset of warmer conditions in the Southwest Pacific region during the latest Paleocene. A scenario of warm conditions is further corroborated by MAT estimates reaching ~ 18 °C as derived from latest Paleocene macrofloras in Southeast Australia (Greenwood et al., 2003).

Considering our sporomorph-based climate estimates for Southeast Australia and Tasmania (Table 3), temperatures were higher during the late Paleocene than during the middle Paleocene. Hence, the overall climatic and vegetation signal suggests that





terrestrial settings across the Southwest Pacific region consistently experienced a pronounced warming during the late Paleocene.

4.2.2 PETM and early Eocene

The effects of the PETM on terrestrial ecosystems in the high southern latitudes are ⁵ yet poorly constrained. Available records from Southeast Australia (Bass and Gippsland Basins) covering the PETM and the earliest Eocene suggest the widespread presence of *Nypa* during that time (Partridge, 1976). Climatically, this indicates a MAT > 21.7 °C (Utescher and Mosbrugger, 2013). For the South Island of New Zealand, sporomorph data from nearshore marine sediments document the development of *Nypa* mangrove swamps and the appearance of pollen from the thermophilous subfamily Cupanieae connected to the PETM interval (Handley et al., 2011); moreover,

- the PETM is characterized by a percentage increase of fern spores as well as of Euphorbiaceae/Eumalvoideae and Myrtaceae pollen at the expense of gymnosperm pollen percentages (Handley et al., 2011). Similarly, sporomorph data for the North
- ¹⁵ Island of New Zealand as available from the Tawanui section show the presence of *Nypa* pollen connected to the PETM (Crouch and Visscher, 2003). However, besides a marked increase of *Dilwynites granulatus* (*Wollemia/Agathis*) pollen, no other significant changes in floristic composition occur (Crouch and Visscher, 2003). Considering that the PETM sediments of the Tawanui section are part of a transgressive systems tract (Crouch and Prinkhvia, 2005). Chaile at al. 2000) the birth chundrases of *Dil*
- tract (Crouch and Brinkhuis, 2005; Sluijs et al., 2008), the high abundances of *Dilwynites granulatus (Wollemia/Agathis)* represent a taphonomic rather than a paleoecological signal as suggested for the sporomorph record of the PETM from ODP Site 1172 (see Sect. 4.1.3.).

With regard to the early Eocene, the majority of vegetation records in the South-²⁵ west Pacific region come from Southeast Australia; they suggest that by early Eocene times the warm temperate, conifer-dominated forests of the late Paleocene had been replaced by more diverse, meso- to megathermal angiosperm forests (Macphail et al., 1994; Greenwood et al., 2003; Greenwood and Christophel, 2005). Although





sporomorph percentages are extremely variable within the available records from Southeast Australia, Araucariaceae, Casuarinaceae, Euphorbiaceae/Eumalvoideae, Proteaceae, and ferns are generally the dominant taxa; typical tropical elements such as *Nypa*, *Anacolosa* and Cupanieae are also recorded (Kemp, 1978; Macphail et al.,

⁵ 1994). This trend in vegetation development during the early Eocene is also documented for Tasmania and it is strongly connected to the PETM event based on the sporomorph record from ODP Site 1172 (see Sect. 4.1.3).

On New Zealand, early Eocene sporomorph assemblages exhibit a mixed Paleocene-Eocene character, with a continued high abundance of conifer pollen

- (Crouch and Visscher, 2003; Raine et al., 2009). However, thermophilous taxa such as Cupanieae, Casuarinaceae and Euphorbiaceae (*Austrobuxus*) are constantly present in these records (Pocknall, 1990; Raine et al., 2009), and Casuarinaceae pollen abruptly started to dominate the sporomorph assemblages from ~ 54.5 Ma onwards (Raine et al., 2009). On the Wilkes Land margin (Antarctica), paratropical vegetation has been recorded during the early Eocene (53.9–51.9) with the notable presence of
- thermophilous taxa such as Arecaceae (palms) and Bombacoideae (Pross et al., 2012; Contreras et al., 2013).

Related to temperature conditions, early Eocene macrofloras from Southeast Australia suggest a MAT of \sim 19 °C from \sim 56 to \sim 53 Ma (Greenwood et al., 2003), which

- ²⁰ is very similar to our MAT estimates for coeval sporomorph records (~ 55.8–54.3 Ma; lower Malvacipollis diversus zone of Partridge, 2006; Table 1) from the Bass and Gipp-sland Basins (~ 18 °C; Table 3). On the Wilkes Land margin, climatic estimates for the early Eocene (53.9–51.9) suggest MAT of ~ 16 °C for the lowland regions (Pross et al., 2012). Although occasionally MAT values as high as ~ 23 °C are recorded during the
- earliest Eocene at ODP Site 1172, the mean MAT for this time interval is ~ 14 °C (Fig. 4, Table 3). This is markedly lower than those from Southeast Australia, and even lower than those from the Wilkes Land margin. Considering that our sporomorph-derived climate data from ODP Site 1172 mainly reflect climate conditions along the coast of eastern Tasmania (compare discussion on sporomorph source region in Sect. 3.2), these





relatively low values may suggest that the eastern part of Tasmania was influenced by the relatively cool Tasman Current (Fig. 1). However, this argument is not supported by the TEX₈₆-derived SSTs from ODP Site 1172 for the early Eocene, which are much higher (mean: 23 °C – TEX^L₈₆, 28 °C – TEX^H₈₆; Bijl et al., 2013b). Terrestrial, macroflorally derived temperatures on the order of 24 °C from western Tasmania (Carpenter et al., 2012) suggest that this region was significantly warmer than the eastern part of Tasmania. Alternatively, another potential explanation for this discrepancy is that the sea-level rise during the early Eocene biased the composition of the sporomorph assemblages at ODP Site 1172 (compare Sect. 4.1.3) towards a dominance of easily transported and/or particularly abundant sporomorphs indicative of cool conditions at the expense of rarer sporomorphs indicative of warmer conditions. This scenario is supported by the fact that many thermophilous plants from the Lower Eocene of the Southwest Pacific region (e.g., Arecaceae (palms), Cupanieae, *Ilex, Nypa*) are mainly insect-pollinated (Bush and Rivera, 1998; Barfod et al., 2011). Hence, these taxa reach only low abun-

dances in pollen spectra when compared to wind-pollinated taxa, and they are not likely to be transported over larger distances before they settle (Jackson, 1994).

Despite of the potential bias on the early Eocene sporomorph assemblages at Site 1172, the supraregional replacement of temperate forests by paratropical forests during the early Eocene on Southeast Australia, New Zealand and Tasmania and the widespread occurrence of *Nypa* palms during the PETM on the same regions consistently indicate a pronounced reorganization of the vegetation during the early Eocene in the high southern latitudes connected to the PETM.

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4.3 Integration with other precipitation records from the southern high latitudes

Based on our paleoclimatic results from the sporomorph record of ODP Site 1172,
 MAP on Tasmania was nearly constant (~ 180 cm yr⁻¹) from the middle Paleocene to the early Eocene (Fig. 4). These values are comparable to the present-day precipitation received by rainforests in western Tasmania at ~ 42° S (185 cm yr⁻¹, Corinna; Bureau of Metereology, 2013), along the west coast of the South Island of New Zealand



(212 cm yr⁻¹; Westport; NIWA, 2013), northern Australia (180 cm yr⁻¹, Darwin Botanic Gardens; Bureau of Metereology, 2013) and on the east coast of New Caledonia (197 cm yr⁻¹, Puoébo; Pesin et al., 1995). The high-precipitation regime on Tasmania during the early Paleogene as deduced from the sporomorphs is further corrobo-

rated by the dominance of the dinocyst genus *Senegalinium* at ODP Site 1172 during this time interval (Sluijs et al., 2011); this genus is characterized by many fresh-water tolerant species (Sluijs et al., 2005; Brinkhuis et al., 2006). Hence, the observed *Senegalinium* dominance, which is best explained by substantial freshwater input, is well compatible with high precipitation on Tasmania and a resulting strong freshwater influx
 on the Tasmanian continental shelf during the early Paleogene.

Because of the general lack of precipitation data from other sites at high southern latitudes we can mainly compare our estimates from Tasmania with values deduced from other coeval paleobotanical records. Based on our results from sporomorph assemblages from Southeast Australia (Table 3), this region experienced similarly high precipitation (MAP mean: ~ 200 cmyr⁻¹) during the early Paleogene; this is consistent with MAP mean estimates (186–240 cmyr⁻¹) as derived from macrofloral records from the Upper Paleocene to Lower Eocene of the same region (Greenwood et al., 2003). On the Wilkes Land margin (Antarctica), high precipitation values (MAP mean: ~ 132 cmyr⁻¹) are also suggested for the early Eocene (Pross et al., 2012). These high precipitation values (> 100 cmyr⁻¹) have been corroborated recently by alkaline major element geochemistry for the same Eocene sediments from Antarctica (Passchier et al., 2013).

Based on the overall precipitation data, Tasmania and the Australia-Antarctic region experienced high rainfall conditions during the early Paleogene comparable with

²⁵ present-day rainforests from southern latitudes. This lends support to modeling studies that include high atmospheric humidity as an important warming mechanism for the higher latitudes (e.g., Abbot et al., 2009).





4.4 Comparison with marine temperature evolution

Based on our sporomorph data (as evidenced in the Axis 1 sample scores of the DCA results, which represent the variation in floristic composition along the studied interval; Fig. 4) and the TEX_{86}^{L} and TEX_{86}^{H} data of Bijl et al. (2009, 2013b) and Hollis et al. (2014),

- there is a strong correlation between the vegetation composition on eastern Tasmania and SST at ODP Site 1172 (Fig. 4). This connection is clearly borne out by our results from the state space model, where there is a very strong correlation of 0.997 when DCA Axis 1 sample scores are compared with TEX^L₈₆ (95% highest posterior density: 0.633–0.999 based on Eq. (4); compare Sect. 2.5) and 0.978 when DCA Axis 1 sample
- scores are compared with TEX^H₈₆ (95 % highest posterior density: 0.879–0.997 based on Eq. (4); compare Sect. 2.5). Hence, the strong correlation between the temperature variability derived from TEX₈₆ and the floristic composition recorded at ODP Site 1172 demonstrates the impact of temperature on the vegetation dynamics in the Southwest Pacific region during the early Paleogene.
- A close coupling between the temperature evolution in the marine and the terrestrial realms is also evident through the comparison of our sporomorph-based temperature estimates (notably WMMTs) with the TEX₈₆-derived SSTs (Fig. 4 and Table 3); it is only during the early Eocene that the pronounced warming trend recorded by TEX^L₈₆ and TEX^H₈₆ is not clearly reflected in the sporomorph-based temperature estimates, likely due to the sea-level increase during the early Eocene (compare Sects. 4.1.3 and 4.2.2).

The terrestrial, sporomorph-derived MATs are markedly cooler than the SSTs derived from TEX^L₈₆ and TEX^H₈₆ (Fig. 4), which based on the traditional perception of the TEX₈₆ proxy are supposed to represent surface-water MAT (e.g., Schouten et al., 2002). At the same time, the TEX₈₆-derived SSTs are closely related to the sporomorph-derived WMMTs (Fig. 4, Table 3). These observations suggest that TEX₈₆based temperatures are biased towards summer conditions when applied to early Paleogene records from the high southern latitudes. Such a seasonal bias has also been





suggested for other early Paleogene records from the Southwest Pacific Ocean based on the comparison of TEX₈₆-derived SSTs with δ^{18} O and Mg/Ca values, and with SSTs derived from climate model simulations (Hollis et al., 2012).

5 Conclusions

- ⁵ The middle Paleocene to early Eocene vegetation on Tasmania as reconstructed from the sporomorph record of ODP Site 1172 was characterized by three different forest types that thrived in high-precipitation regimes under different temperature conditions. These forest types were: (i) warm-temperate rainforests dominated by Podocarpaceae during the middle and late Paleocene; (ii) cool-temperate rainforests dominated by *Nothofagus* and Araucariaceae that transiently prevailed across the middle/late Pa-
- leocene transition interval (iii) paratropical rainforests dominated by Cyatheaceae during the early Eocene with the remarkable presence of the mangrove palm *Nypa* during the PETM and the earliest Eocene. The comparison with other, previously published floral records from the Southwest Pacific region (including Southeast Australia
- and New Zealand) further supports the validity of our data for Tasmania. It shows that temperate forests were replaced by paratropical forests during the early Eocene throughout the Southwest Pacific region. This reorganisation in vegetation composition included an increase in fern (mainly Cyatheaceae) and angiosperm abundances (e.g., Proteaceae, Euphorbiaceae/Eumalvoideae, Casuarinaceae) at the expense of gymnosperms (mainly podocarps).

The integration of terrestrial (i.e., floristic) and previously published marine (i.e., TEX₈₆-based SST) climate information from ODP Site 1172 shows that the surfacewater cooling of ~ 3 °C across the middle/late Paleocene transition interval (~ 59.5 to ~ 59.0 Ma) was paralleled by a transient demise of frost-sensitive plants (i.e., palms and cycads) and the establishment of cool-temperate forests dominated by *Nothofagus* and

²⁵ cycads) and the establishment of cool-temperate forests dominated by *Nothofagus* and Araucariaceae on Tasmania. This suggests that cooler conditions (and notably harsher winters with strong and/or frequent frosts) prevailed on Tasmania during that time.





In light of the statistically robust connection between the floristic composition of the sporomorph record from ODP Site 1172 and the previously published TEX₈₆-based SST record from the same site, the vegetation dynamics on Tasmania during the middle Paleocene to early Eocene were mainly driven by temperature; precipitation remained high (with a MAP mean of $\sim 180 \,\mathrm{cm\,yr}^{-1}$) throughout that time. Based on the comparison of our sporomorph-derived temperatures with the TEX₈₆-based SSTs, we conclude that TEX_{86}^{L} - and TEX_{86}^{H} -derived temperatures for the high southern latitudes of the early Paleogene are likely biased towards summer conditions.

Supplementary material related to this article is available online at http://www.clim-past-discuss.net/10/291/2014/cpd-10-291-2014-supplement.zip.

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Table 1. Sporomorph datasets evaluated in this study from the Southeast Australia (Bass Basin, Gippsland Basin, Southeast Highlands) and New Zealand (Middle Waipara and Otaio River sections). Asterisks indicate data derived from palynological reports in Department of Primary Industries (1999); Plus signs denote data derived from the Appendix C of Blevin (2003). Biozones and ages are based on Stover and Evans (1973), Stover and Partridge (1973) and Partridge (2006).

| Section | Region | Age | Pollen Biozone | Authors |
|--------------------------------------|---|--|---------------------------------------|----------------------------------|
| Southern Monaro sections | highlands of southeastern Australia | 60–58 Ma | Lygiestepollenites balmei | Taylor et al. (1990) |
| * Burong-1 | onshore Gippsland Basin | late Paleocene | Upper L. balmei | Partridge and Macphail (1997) |
| * Kingfish-8 | offshore Gippsland Basin | early Eocene | Lower Malvacipollis diversus | Partridge (1992) |
| + Konkon-1 | Bass Basin | middle to late Paleocene, early Eocene | L. balmei and Lower M. diversus | Partridge (2003) |
| Otaio and Middle Waipara sections | New Zealand | 58–56 Ma | | Pancost et al. (2014 |
| + Poonboon-1 | Bass Basin | late Paleocene, early Eocene | Upper L. balmei, Lower M. diversus | Partridge (2003) |
| * Roundhead-1 | offshore Gippsland Basin | middle to late Paleocene, early Eocene | L. balmei, Lower M. diversus | Partridge (1989) |
| * Sweetlips-1 | offshore Gippsland Basin | middle to late Paleocene | L. balmei | Partridge (1989) |
| * Turrum-4 | offshore Gippsland Basin | middle to late Paleocene | L. balmei | Partridge (1993) |
| * Turrum-4 | offshore Gippsland Basin | middle to late Paleocene | L. balmei | Partridge (1993) |



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Table 2. List of fossil sporomorph taxa from the Middle Paleocene to Lower Eocene of ODP Site 1172 with known botanical affinities and literature source, the nearest living relative (NLR) used in the climate reconstruction, and database where climate profiles of the NLRs are derived from. Database (1) = PALAEOFLORA (Utescher and Mosbrugger, 2013), Database (2) = Pross et al. (2012). Taxa used in the climatic evaluation are printed in boldface.

| Fossil taxon | Botanical affinity | Source | NLR used for climate analysis | database |
|---|--|----------------------------|-------------------------------------|----------|
| Araucariacites spp. | Araucariaceae | Raine et al. (2008) | Araucariaceae | 1, 2 |
| Arecipites spp. | Arecaceae | Nichols et al. (1973) | Arecaceae | 1 |
| Baculatisporites spp. | Osmundaceae | Raine et al. (2008) | Osmundaceae | 1 |
| Banksieaeidites arcuatus | Proteaceae (<i>Banksia,</i>) Dryandra, Musgravea | Raine et al. (2008) | | |
| Beaupreaidites cf. diversiformis | Proteaceae (Beauprea) | Raine et al. (2008) | | |
| Caryophillidites sp. | Caryophyllaceae | Raine et al. (2008) | | |
| Ceratosporites spp. | Lycopodiaceae, Selaginellaceae | Raine et al. (2008) | | |
| Crassoretitriletes cf. vanraadshooveni | Lygodium | Germeraad et al. (1968) | Lygodium | 1 |
| Cyathidites spp. | Probably Cyatheaceae | Mohr (2001) | Cyatheaceae | 1 |
| Cycadopites spp. | Cycadales | Raine et al. (2008) | Cycadales: Bowenia, Lepidozamia, | 1 |
| | Dedeestreeses | | Macrozannia | |
| australiensis | <i>Dacrycarpaceae</i> (<i>Dacrycarpus</i>) <i>dacrydioides</i> | Raine et al. (2008) | Dacrycarpus | I |
| Dacrydiumites florinii | Podocarpaceae (<i>Dacrydium</i>) | Raine et al. (2008) | Dacrydium | 1 |
| <i>Dacrydiumites</i> spp. | Podocarpaceae (<i>Dacrydium</i>) | Raine et al. (2008) | Dacrydium | 1 |
| Dilwynites granulatus | Araucariaceae (Wollemia/Agathis) | Macphail et al. (2013) | | |
| Dilwynites tuberculatus | Araucariaceae (Wollemia/Agathis) | Macphail et al. (2013) | | |
| Ephedripites sp. | Ephedra | | | |



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Table 2. Continued.

| Fossil taxon | Botanical affinity | Source | NLR used for climate analysis | database |
|---|--|---------------------|--|----------|
| Gleicheniidites senonicus | Gleicheniaceae | Raine et al. (2008) | Dicranopteris, Diplopterygium, Gleichenia, Sticherus | 1 |
| <i>Gleicheniidites</i> spp. | Gleicheniaceae | Raine et al. (2008) | Dicranopteris, Diplopterygium, Gleichenia, Sticherus | 1 |
| Intratriporopollenites cf. notabilis | Sterculioideae?, Bombacoideae?, Tilioideae? | Raine et al. (2008) | | |
| Malvacipollis diversus | Euphorbiaceae (<i>Austrobuxus,</i> <i>Dissiliaria,</i> <i>Petalostigma</i>); Eumalvoideae? | Raine et al. (2008) | Euphorbiaceae | 1 |
| <i>Microalatidites</i> spp. | Podocarpaceae (cf. <i>Phyllocladus</i>) | Raine et al. (2008) | Phyllocladus | 1 |
| Microcachrydites antarcticus | Podocarpaceae (<i>Microstrobos,</i> <i>Microcachrys tetragona</i>) | Raine et al. (2008) | | |
| Myricipites harrisii | Casuarinaceae, possibly also Myricaceae | Raine et al. (2008) | Casuarinaceae (all Australian species) | 1 |
| Myrtaceidites spp. | Myrtaceae | Raine et al. (2008) | Myrtaceae | 1 |
| Nothofagidites | Nothofagaceae | Truswell and | N. cunninghamii, | 1 |
| asperus complex | (<i>Nothofagus</i> subg. <i>Lophozonia</i>) | Macphail (2009) | N. moorei | |
| Nothofagidites | Nothofagaceae | Truswell and | Nothofagaceae | 2 |
| brachyspinulosus | (<i>Nothofagus</i> subg. | Macphail (2009) | (<i>Nothofagus</i> subg. | |
| complex | Fuscospora) | | Fuscospora) | |
| Nothofagidites | Nothofagaceae | Truswell and | all subgenera (including | 2 |
| emarcidus complex | (Nothofagus) | Macphail (2009) | Brassospora) | |
| (including N. endurus) | | | | |
| Nothofagidites | Nothofagaceae | Raine et al. (2008) | Nothofagaceae | 2 |
| flemingii | (Nothofagus subg. | | (<i>Nothofagus</i> subg. | |
| complex | Nothofagus) | | Fuscospora) | |



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Table 2. Continued.

| Fossil taxon | Botanical affinity | Source | NLR used for climate analysis | database |
|---|---|---------------------|---|----------|
| Nothofagidites lachlaniae complex | Nothofagaceae (<i>Nothofagus</i> subg. <i>Fuscospora</i>) | Raine et al. (2008) | Nothofagaceae (<i>Nothofagus</i> subg. <i>Euscospora</i>) | 2 |
| Nothofagidites sp.1 | Nothofagaceae (Nothofagus) | | all subgenera (including Brassospora) | 2 |
| Nothofagidites | Nothofagaceae | | all subgenera (including | 2 |
| spp. undifferentiated | (Nothofagus) | | Brassospora) | |
| Osmundacidites spp. | Osmundaceae | Raine et al. (2008) | Osmundaceae | 1 |
| Parvisaccites catastus | Podocarpaceae | Raine et al. (2008) | | |
| | (Halocarpus) | | | _ |
| Phyllocladidites | Lagarostrobos | Raine et al. (2008) | Lagarostrobos | 2 |
| mawsonii | | | - / | |
| Podocarpidites | Podocarpaceae | Raine et al. (2008) | Podocarpus | 1 |
| ellipticus | (Podocarpus) | | | |
| Podocarpidites | Podacarpaceae | Raine et al. (2008) | | |
| exiguus Proto o cidito o | Dretesses | | | 0 |
| Proteacidites | | Raine et al. (2008) | Adenantnos | 2 |
| adenantholdes | (Adenantnos) | Daina at al (0000) | Videmelum | 0 |
| Proleaciones | (Yulamalum agaidentale | Raine et al. (2008) | Aylometum | 2 |
| annularis | or Lambertia) | | (all Australian species) | |
| Proteacididites cf. | Proteaceae | Raine et al. (2008) | | |
| amolosexinus | (<i>Knightia excelsa</i> type) | | | |
| Proteacidites cf. | Proteaceae | Raine et al. (2008) | Adenanthos | 2 |
| adenanthoides | (Adenanthos) | | | |
| Proteacidites parvus | Proteaceae | Raine et al. (2008) | Bellendena montana | 2 |
| | (Bellendena montana type) | | | |
| Proteacidites | Proteaceae | Raine et al. (2008) | Gevuina, Hicksbeachia | 2 |
| reticuloscabratus | (Gevuina/Hicksbeachia type) | | | - |
| Proteacidites | Proteaceae | Raine et al. (2008) | Symphyonema, | 2 |
| symphyonemoides/ | (Symphyonema, | | Petrophile | |
| P. pseudomoides | Carnarvonia) | | | |
| Pseudowinterapoilis sp. | | Haine et al. (2008) | | |
| Hellifletes CI. | Lycopodiaceae | Haine et al. (2008) | | |
| rosewoodensis | (Lycopoaium) | | | |



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Table 2. Continued.

| Fossil taxon | Botanical affinity | Source | NLR used for climate analysis | database |
|------------------------------------|---|---------------------------------|-------------------------------|----------|
| Retitriletes facetus | Lycopodicaceae (<i>Lycopodium</i>) | Raine et al. (2008) | | |
| Rubinella cf. major | Leptolepis? | Raine et al. (2008) | | |
| Spinizonocolpites prominatus | Nypa (Arecaceae) | Muller (1968) | Nypa | 1 |
| Stereisporites sp. | Sphagnaceae | Truswell and Macphail (2009) | | |
| Triporoletes cf. reticulatus | cf. Riccia beyrichiana | Raine et al. (2008) | | |
| Troporopollenites ambiguus | Proteaceae (<i>Telopea truncata,</i> <i>Oreocallis pinnata</i>) | Raine et al. (2008) | | |
| Tripunctisporites maastrichtiensis | Sphagnaceae? | | | |

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Table 3. Sporomorph-based climate estimates for Paleocene to early Eocene records from Southeast Australia (Bass Basin, Gippsland Basin, Southeast Highlands), New Zealand (Middle Waipara and Otaio River sections) and ODP Site 1172. SST values given for each stratigraphic interval represent average values of the data of Bijl et al. (2009, 2013b) and Hollis et al. (2014), with (H) and (L) denoting $\text{TEX}_{86}^{\text{H}}$ respectively $\text{TEX}_{86}^{\text{L}}$. Sporomorph-derived climate estimates are based on the methodology of Greenwood et al. (2005) and are presented with the average and the standard deviation. MAT = Mean Annual Temperature, CMMT = Coldest Month Mean Temperature, WMMT = Warmest Month Mean Temperature, MAP = Mean Annual Precipitation.

| Age | Location | SST (°C) | MAT (°C) | CMMT (°C) | WMMT (°C) | MAP ($cm yr^{-1}$ |
|--|---|----------------------------------|--|--|--|--|
| early Eocene | ODP Site 1172 | 23.4 ± 1.7 (L) 27.8 ± 2.5 (H) | 14±3 | 8.3±3 | 19.5 ± 2.5 | 182 ± 13 |
| | Bass Basin Gippsland Basin Otaio section | | 19 ± 2.6 17.2 ± 1.8 18.9 | 11.2 ± 2.2 10.2 ± 2.7 11.8 | 23.4 ± 1.6 22 ± 0.9 24.1 | 200 ± 8 200 ± 6 125 |
| late Paleocene | ODP Site 1172 | 22 ± 1.2 (L) 25.5 ± 1.1 (H) | 14.5 ± 2.1 | 7.8 ± 1 | 20.2 ± 1.9 | 184 ± 17 |
| | Bass Basin Gippsland Basin Southern Morano sections Middle Waipara section | 14 ± 1.6 (L) 19.3 ± 1.7 (H) | $16.2 \pm 1.2 \\ 16.2 \pm 1.8 \\ 15.5 \pm 1.4 \\ 14.1 \pm 2.7$ | 8.8 ± 0.9 8.9 ± 1 8.2 ± 1 8.9 ± 2 | 21.5 ± 0.4 20.4 ± 2 20.1 ± 1.7 17.9 ± 2.6 | 197 ± 10 201 ± 6 198 ± 12 179 ± 7 |
| middle/late Paleocene transition | ODP Site 1172 | 16.5 ± 1.5 (L) 21.2 ± 1.1 (H) | 9.9±2.2 | 5.4 ± 1.1 | 16.2±2 | 185 ± 10 |
| middle Paleocene | ODP Site 1172 | 18.1 ± 1.9 (L) 23.5 ± 1 (H) | 11.8 ± 2.5 | 6.7 ± 1.5 | 18.1 ± 2.5 | 186 ± 10 |
| | Bass Basin Gippsland Basin | | 14.8 ± 1.5 14.2 ± 1.4 | 7.4 ± 0.8 7.7 ± 1.4 | 20.3 ± 0.5 18.6 ± 1.6 | 200 ± 3 198 ± 5 |
| | | | | | | |







Fig. 1. Map of the Southwest Pacific Ocean showing the early Eocene (~ 53 Ma) continental configuration, illustrating modern continents (gray), areas shallower than 300 m (blue) and locations of ODP Site 1172 and localities listed in Table 1. SM = Southern Monaro Sections (Southeast Highlands), B = Burong-1, K = Kingfish-8, Ko = Konkon-1, MW-OS = Middle Waipara and Otaio River sections, P = Poonboon-1, R = Roundhead-1, S = Sweetlips-1, T = Turrum-4, ETP = East Tasman Plateau. Modified after Cande and Stock (2004) and Sluijs et al. (2011).





Fig. 2. Relative abundances of selected sporomorph taxa ([A] angiosperms, [G] gymnosperms, [P] pteridophytes) representative of the middle Paleocene to early Eocene assemblages from ODP Site 1172. DCA Axis 1 scores represent the fluctuations in floristic composition between samples. Relative abundances and DCA results are based on samples with counts \geq 100 individuals only. Numbers of sporomorph species are rarefied at 100 individuals. The intervals corresponding to the middle/late Paleocene transition and PETM are marked by horizontal blue and red bars, respectively. Age model and dinocyst zonation after Bijl et al. (2013b).







Fig. 3. Comparison of the floristic composition based on the DCA sample scores for (a) middle Paleocene to early Eocene sporomorph assemblages from ODP Site 1172; (b) Paleocene/early Eocene sporomorph assemblages from Southeast Australia (Bass Basin, Gippsland Basin, Southeast Highlands), New Zealand (Middle Waipara and Otaio River sections) and ODP Site 1172. Results are based on samples with counts > 100 individuals.





Fig. 4. Comparison of the sporomorph-derived climate estimates, SST values based on TEX_{86}^{L} and TEX_{86}^{H} , and DCA Axis 1 sample scores from the Middle Paleocene to Lower Eocene of ODP Site 1172. Sporomorph-derived climate estimates are based on the methodology of Greenwood et al. (2005). Error bars represent the minimum and maximum estimates determined using that method. SST data are from Bijl et al. (2009, 2013b) and Hollis et al. (2014). Sporomorph-derived climate estimates and DCA results are based on samples with counts \geq 100 individuals only. The intervals corresponding to the middle/late Paleocene transition and the PETM are marked by horizontal blue and red bars, respectively. Age model and dinocyst zonation after Bijl et al. (2013b).



