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# Eocene to Oligocene vegetation and climate in the Tasmanian Gateway region were controlled by changes in ocean currents and $p\text{CO}_2$

Michael Amoo<sup>1</sup>, Ulrich Salzmann<sup>1</sup>, Matthew J. Pound<sup>1</sup>, Nick Thompson<sup>1</sup>, and Peter K. Bijl<sup>2</sup>

<sup>1</sup>Department of Geography and Environmental Sciences, Northumbria University, Newcastle upon Tyne, UK

<sup>2</sup>Marine Palynology and Palaeoceanography, Utrecht University, Princetonlaan 8A, Utrecht, the Netherlands

**Correspondence:** Michael Amoo (michael.amoo@northumbria.ac.uk)

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**Abstract.** Considered one of the most significant climate reorganizations of the Cenozoic period, the Eocene–Oligocene Transition (EOT; ca. 34.44–33.65) is characterized by global cooling and the first major glacial advance on Antarctica. In the southern high latitudes, the EOT cooling is primarily recorded in the marine realm, and its extent and effect on the terrestrial climate and vegetation are poorly documented. Here, we present new, well-dated, continuous, high-resolution palynological (sporomorph) data and quantitative sporomorph-based climate estimates recovered from the East Tasman Plateau (ODP Site 1172) to reconstruct climate and vegetation dynamics from the late Eocene (37.97 Ma) to the early Oligocene (33.06 Ma). Our results indicate three major climate transitions and four vegetation communities occupying Tasmania under different precipitation and temperature regimes: (i) a warm-temperate *Nothofagus*–Podocarpaceae-dominated rainforest with paratropical elements from 37.97 to 37.52 Ma; (ii) a cool-temperate *Nothofagus*-dominated rainforest with secondary Podocarpaceae rapidly expanding and taking over regions previously occupied by the warmer taxa between 37.306 and 35.60 Ma; (iii) fluctuation between warm-temperate–paratropical taxa and cool temperate forest from 35.50 to 34.49 Ma, followed by a cool phase across the EOT (34.30–33.82 Ma); and (iv) a post-EOT (earliest Oligocene) recovery characterized by a warm-temperate forest association from 33.55 to 33.06 Ma. Coincident with changes in the stratification of water masses and sequestration of carbon from surface water in the Southern Ocean, our sporomorph-based temperature estimates between 37.52 and 35.60 Ma (phase ii) showed 2–3 °C terrestrial cooling. The

unusual fluctuation between warm and cold temperate forest between 35.50 to 34.59 Ma is suggested to be linked to the initial deepening of the Tasmanian Gateway, allowing eastern Tasmania to come under the influence of warm water associated with the proto-Leeuwin Current (PLC). Further to the above, our terrestrial data show the mean annual temperature declining by about 2 °C across the EOT before recovering in the earliest Oligocene. This phenomenon is synchronous with regional and global cooling during the EOT and linked to declining  $p\text{CO}_2$ . However, the earliest Oligocene climate rebound along eastern Tasmania is linked to a transient recovery of atmospheric  $p\text{CO}_2$  and sustained deepening of the Tasmanian Gateway, promoting PLC throughflow. The three main climate transitional events across the studied interval (late Eocene–earliest Oligocene) in the Tasmanian Gateway region suggest that changes in ocean circulation due to accelerated deepening of the Tasmanian Gateway may not have been solely responsible for the changes in terrestrial climate and vegetation dynamics; a series of regional and global events, including a change in the stratification of water masses, sequestration of carbon from surface waters, and changes in  $p\text{CO}_2$ , may have also played vital roles.

## 1 Introduction

Palynological reconstruction demonstrates a high sensitivity of global vegetation to past changes in climate, leading to major shifts in biome distribution (Pound and Salzmann, 2017). The Eocene–Oligocene Transition (EOT; 34.44–33.65 Ma; Katz et al., 2008; Hutchinson et al., 2021) is one of

the most important climate transitions of the Cenozoic and is characterized by a shift from largely ice-free greenhouse conditions to an icehouse climate, involving the development of the Antarctic cryosphere and global cooling (Liu et al., 2009; Pearson et al., 2009; Pagani et al., 2011; Hutchinson et al., 2021).

Tectonic opening of the southern gateways (Kennett, 1977) and a large and sharp drop in global atmospheric CO<sub>2</sub> (DeConto and Pollard, 2003; Huber et al., 2004; Zachos et al., 2008; Goldner et al., 2014; Ladant et al., 2014) have been proposed as possible drivers for this climate transition. The opening of the Australian–Antarctic Seaway (Tasmanian Gateway) and the Drake Passage led to the strengthening of the Antarctic Circumpolar Current (ACC), which thermally isolated Antarctica (Kennett, 1977). However, marine geology, micropalaeontology, and model simulation showed a potential time lag between the onset of the ACC and palaeogeographic changes, hence challenging global climate change at the EOT driven by Southern Hemisphere tectonics (Huber et al., 2004; Stickley et al., 2004; Goldner et al., 2014).

Although southern gateway opening and deepening have failed to fully explain Antarctic cooling at the EOT, the oceanographic changes following gateway opening and deepening have been reported to climatically impact Southern Ocean surface waters regionally (Stickley et al., 2004; Sijp et al., 2011; Houben et al., 2019; López-Quirós et al., 2021; Thompson et al., 2022). However, the extent and effect of the opening and deepening of the Tasmanian Gateway and its associated oceanographic changes on the coeval terrestrial climate and vegetation are not readily known. The lack of continuous and well-dated EOT terrestrial records place considerable limitations on the detailed temporal and spatial reconstruction of vegetation and climate. These challenges are further compounded by the fact that the few late Eocene and early Oligocene terrestrial palynoflora records indicate a rather heterogeneous vegetation response at the EOT (Pound and Salzmann, 2017). For example, in southeastern Australia, the late Eocene to early Oligocene vegetation records indicate a shift from a warm-temperate to a cool-temperate rainforest (Korasidis et al., 2019; Lauretano et al., 2021), whereas in New Zealand, a warm humid rainforest persisted (Pocknall, 1989; Homes et al., 2015; Prebble et al., 2021). East Antarctica (Prydz Bay) saw the collapse of tall woody vegetation and their replacement by impoverished, taiga-like vegetation with dwarfed trees before the EOT during the late Eocene (Macphail and Truswell, 2004; Truswell and Macphail, 2009; Tibbett et al., 2021), whereas a major vegetation change did not take place across the Drake Passage region until the early Oligocene, where there is a distinct expansion of gymnosperms and cryptogams, indicating glacial expansion (Thompson et al., 2022).

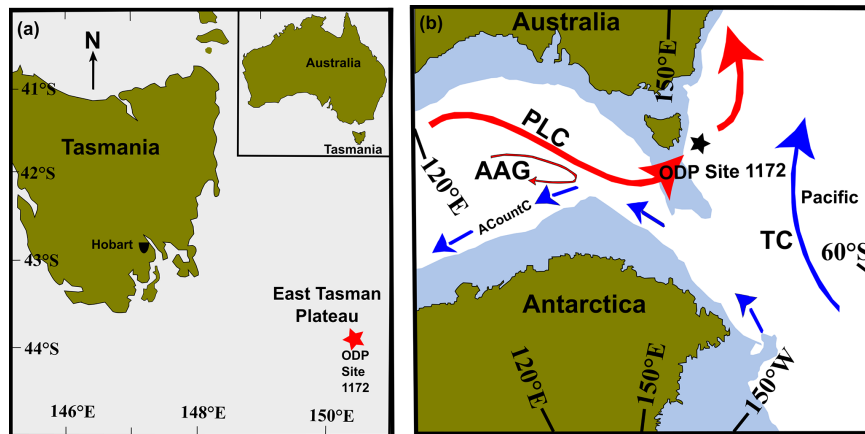
To further our understanding of the timing and potential drivers of southern high-latitude terrestrial environment change at the EOT, this study presents a new sporomorph record recovered from ODP Site 1172 (Fig. 1) on the East

Tasman Plateau (ETP) spanning the late Eocene (37.97 Ma) to earliest Oligocene (33.06 Ma). The proximity of our study site to the Tasmanian Gateway places it in an excellent geographical position to identify potential climate or tectonic impacts on terrestrial vegetation of the Australo-Antarctica region. To further investigate potential links between the terrestrial and marine realm, we also compare our pollen-based quantitative climate estimates with newly published TEX<sub>86</sub>-based sea-surface temperature (SST) and mean annual air temperature (MAAT<sub>soil</sub>) reconstructions from the same site (Bijl et al., 2021). Our study reveals a significant terrestrial cooling  $\sim 3$  Myr prior to the EOT, and a warming in the earliest Oligocene that is most likely controlled by transient rebound of atmospheric *p*CO<sub>2</sub> and sustained deepening of the Tasmanian Gateway.

## 2 Materials and methods

### 2.1 Tectonic evolution and depositional setting

Continental breakup and seafloor spreading between Australia and the continental blocks of Lord Howe Rise, Campbell Plateau, and New Zealand (LCNZ) started in the Late Cretaceous ( $\sim 75$  Ma; Cande and Stock, 2004). Northward movement of Australia was propagated by rifting leading to the formation of the Tasman Sea and the separation of northeastern Australia in the Paleocene ( $\sim 60$  Ma; Gaina et al., 1999). The series of tectonic events paved the way for major ocean currents to flow along the coasts of eastern Australia and Tasmania, the ETP, and the South Tasman Rise (STR; Exon et al., 2004a). However, the Tasman promontory remained and separated the Australo-Antarctic Gulf (AAG) from the Pacific Ocean until the late Eocene ( $\sim 35.5$  Ma; Stickley et al., 2004). Our study site (ODP Site 1172 on the ETP; Fig. 1) is located on one of the four continental blocks sampled during ODP Leg 189 (Exon et al., 2004b)  $\sim 170$  km southeast of Tasmania (43°57.6' S, 149°55.7' E; Fig. 1a; Shipboard Scientific Party, 2001) at water depths of  $\sim 2620$  m (Exon et al., 2004a) and is enclosed by an 1800 m high seamount (Royer and Rollet, 1997). Prior to the Tasman Sea break-up in the Late Cretaceous (95 Ma), the ETP (which presently forms an oval platform) was part of Tasmania and the STR (Royer and Rollet, 1997; Exon et al., 2004b) and subsided slowly until the late Eocene. Bathymetric studies indicate that the ETP is connected to the east coast of Tasmania by the East Tasman Saddle (Royer and Rollet, 1997), which gives no indication of a deep basin in between (Hill and Exon, 2004). A dredging exercise confirmed the continental origin of the plateau (Exon et al., 1997). However, the age of the guyot/seamount (dated as 36 Ma; Lanyon et al., 1993) disqualifies the ETP itself from being the potential source of the terrestrial organic matter (Bijl et al., 2021). In addition, common Permo-Triassic reworked elements in our late Eocene–early Oligocene sporomorph assemblage likely indicate an eastern Tasmanian sporomorph



**Figure 1.** (a) Location of the East Tasman Plateau (ODP Site 1172; red star) and present-day Tasmania (Quilty, 2001). The Tasmania landmass is shown in green and the submerged ODP Site 1172 in grey, at a water depth of  $\sim 2620$  m. (b) Early Oligocene palaeogeography and palaeoceanography of the Tasmanian Gateway. ODP Site 1172 is marked by a five-pointed black star. Surface currents are modified after reconstructions by Stickley et al. (2004). TC: Tasman current, PLC: proto-Leeuwin current, ACountC: Antarctic Counter Current, AAG: Australo-Antarctic Gulf. Solid red arrows indicate warmer ocean currents from the AAG, and solid blue arrows indicate cooler ocean currents. Arrow size indicates the relative strength of the current. Figure is modified after Hoem et al. (2021).

source, in line with the Permian–Triassic upper Parmeener Group that contains terrestrial deposits and presently makes up the surface lithology across east Tasmania. A previous Paleocene–Eocene sporomorph assemblage from the ETP (ODP Site 1172) further supports an eastern Tasmanian terrestrial palynomorph source (Contreras et al., 2014).

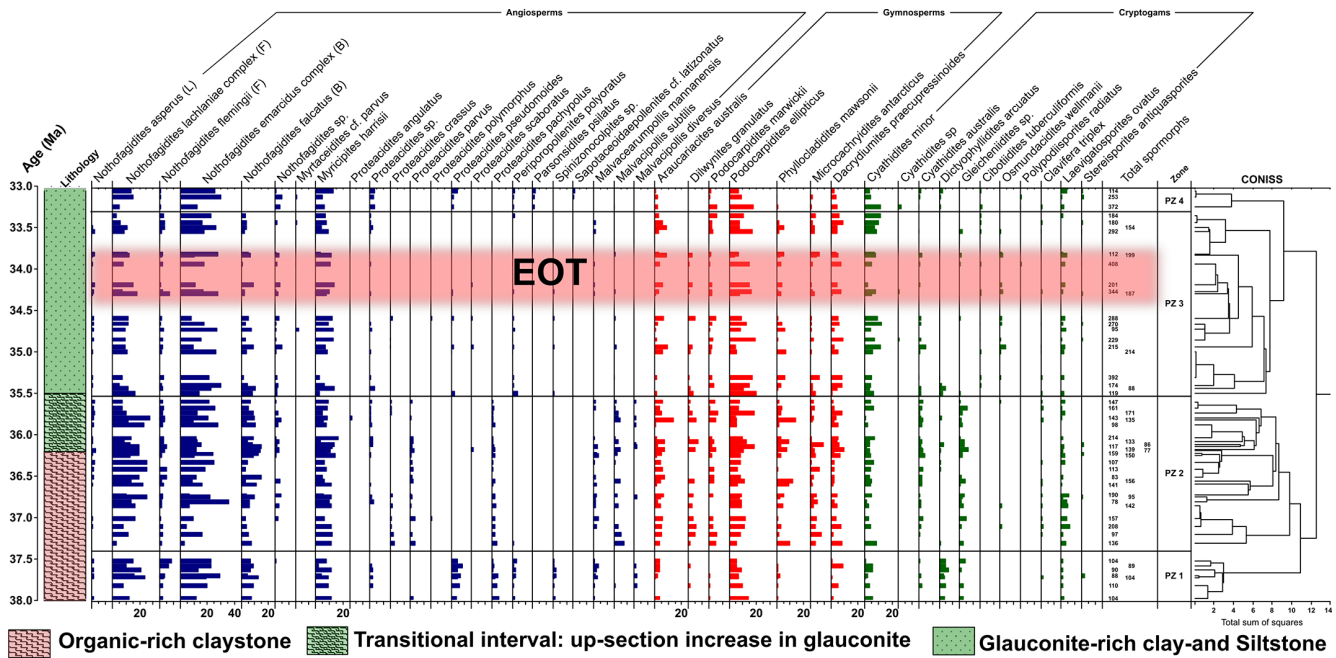
Lithologically, the marine sedimentary record is divided into three units: (i) shallow-marine, organic-rich middle Eocene to lower upper Eocene clay; (ii) a highly condensed middle upper Eocene to lowermost Oligocene glauconite-rich, shallow-marine silty sandstone; and (iii) lower Oligocene siliceous-rich carbonate ooze (Stickley et al., 2004; Exon et al., 2001). Holes A and D of ODP Site 1172 on the East Tasman Plateau both yielded EOT records and have been analysed for their pollen and spore contents. The age model relies on magnetostratigraphy (which has a particularly clear signal in the late Eocene; Stickley et al., 2004; Fuller and Touchard, 2004) and biostratigraphy (dinoflagellate cysts, nannoplankton, and diatoms; Stickley et al., 2004; Bijl et al., 2013), as presented in Houben et al. (2019) and Bijl et al. (2021).

## 2.2 Study material

A total of 66 samples from the late Eocene to the earliest Oligocene of ODP Site 1172 (37.97–33.06 Ma) were analysed for terrestrial palynomorphs to reconstruct the palaeovegetation and palaeoclimate. Raw pollen data, including data on non-pollen palynomorphs (NPPs) and reworked sporomorphs, are available from the Zenodo data repository (Amoo et al., 2021). These samples were prepared at the Laboratory of Palaeobotany and Palynology, Utrecht University, following standard palynological pro-

cessing techniques (Bijl et al., 2013). Sample processing involved treatment with 30 % HCl and 38 % HF and sieving the residues through a  $15\ \mu\text{m}$  nylon mesh (Pross, 2001). The residues were mounted onto microscope slides with glycerine gel used as the mounting medium. When analysing marine sediments such as those used in this study, sieving is a standard technique that is required to remove unwanted organic/inorganic matter and to increase the pollen concentration. To reduce the potential risk of losing small pollen grains, we regularly monitored our residues sieved at 10 and  $15\ \mu\text{m}$  mesh size. We found no evidence of a selective loss of smaller pollen grains such as *Myrtaceidites* and *Sapotaceoidaepollenites* cf. *latizonatus*. Similar to pollen records recovered from lakes (diameter  $> 200$  m) and estuaries in Australia, our marine sporomorph record is likely to be biased towards abundant taxa in the regional vegetation, whereas sporomorphs recovered from coal, lignite, peat, and backswamp deposits are more likely to reflect local flora, with higher diversity and occasional high values of underrepresented taxa (Macphail et al., 1994).

Leica DM 500 and DM 2000 LED microscopes were used to analyse two slides for each sample at  $\times 400$  or  $\times 1000$  magnification. Where possible, 300 fossil spores and pollen grains (excluding reworked sporomorphs) were analysed for each sample, followed by further scanning of the entire microscope slide to record rare taxa. Aside from nine samples with counts below 50 grains, overall pollen preservation and counts were generally good. Reworked sporomorphs were identified based on the thermal maturation (colour) of their outer coat (exine) and occurrence outside their known stratigraphic range. Non-pollen palynomorphs were recorded but not added to the total pollen counts. Sporomorph percentages were calculated based on the total sum of pollen and



**Figure 2.** Sporomorph assemblages and relative abundances of major sporomorph taxa (angiosperms, gymnosperms, cryptogams) recovered from the late Eocene–early Oligocene of ODP Site 1172. Angiosperm relative abundances are marked by blue bars, gymnosperm relative abundances by red bars, and cryptogam relative abundances by green bars. In the angiosperm group, *Nothofagidites* is further divided into subgenera. These are *Brassospora* (B), *Fuscospora* (F), and *Lophozonia* (L) types. CONISS ordination constrains our late Eocene–early Oligocene sporomorph assemblages into four distinct pollen zones (PZ 1–PZ 4) or vegetation and climate phases. The age model is after Houben et al. (2019) and Bijl et al. (2021).

spores excluding reworked grains, and were plotted using Tilia version 2.6.1 (Fig. 2; Grimm, 1990). Using the Edwards and Cavalli-Sforza chord distance, we applied a stratigraphically constrained incremental sum-of-squares cluster analysis (CONISS; Grimm, 1987) to determine pollen assemblage zones (PZ; Fig. 2). Sporomorph identification and botanical affinities (used for nearest living relative identification of fossil spores and pollen) were established using Macphail and Cantrill (2006), Macphail (2007), Truswell and Macphail (2009), Daly et al. (2011), Kumaran et al. (2011), Raine et al. (2011), Bowman et al. (2014), and Macphail and Hill (2018).

### 2.3 Bioclimatic analysis

The nearest living relative (NLR) approach was used to estimate and reconstruct the mean annual temperature (MAT), warm mean month temperature (WMMT), cold mean month temperature (CMMT), and mean annual precipitation (MAP). The bioclimatic analysis used in this study involved all pollen and spore taxa that could be related to an NLR and are listed in Table 1. The NLR is a uniformitarian approach based on the assumption that climate tolerance of extant taxa can be extended into the past. However, factors such as misidentification of fossil taxa and/or their NLRs, unresolved differences in climate tolerance between fossil

taxa and their NLRs, potentially incomplete climate tolerance of NLRs, and potential weakening of the connection between fossil taxa and NLRs through deep time may pose some concerns and need to be considered prior to the application of NLR-based climate reconstructions (Mosbrugger and Utescher, 1997; Mosbrugger, 1999; Pross et al., 2000; Utescher et al., 2000, 2014). Generally, these uncertainties and issues with the NLR approach increase when analysing plant remains or samples from deep-time geological records (Poole et al., 2005). To test the validity of our NLR-based climate estimates, we compare them to previously published independent botanical or geochemical proxies in the southern high latitudes spanning the late Eocene to early Oligocene (e.g. Colwyn and Hren, 2019; Houben et al., 2019; Korasidis et al., 2019; Bijl et al., 2021; Lauretano et al., 2021; Tibbett et al., 2021). Overall, these are generally in agreement and provide a certain level of confidence in the utility of the NLR-based climate estimates.

The NLR analysis in this study is combined with the probability density function (PDF). The PDF works by statistically constraining the most likely climate co-occurrence envelope for an assemblage (Harbert and Nixon, 2015; Hollis et al., 2019). Bioclimatic analysis was performed by using the *dismo* package in R (Hijmans et al., 2017) to cross-plot the modern distribution of the NLR from the Global Biodiversity Information Facility (GBIF; GBIF, 2021) with gridding from



**Table 1.** List of sporomorph taxa from the late Eocene to early Oligocene of ODP Site 1172, accompanied by botanical affinities, literature sources, nearest living relatives (NLRs) selected for climatic reconstruction, and inferred climate ranges from Macphail (2007).

Fossil taxon	Botanical affinity	Source	Selected NLR for climate analysis	Inferred climate range (Macphail, 2007)
<b>Gymnosperms</b>				
<i>Arauciacites australis</i>	Araucariaceae	Raine et al. (2011)	Araucariaceae	Lower to upper ?mesotherm
<i>Dilwynites granulatus</i>	Araucariaceae	Raine et al. (2011)	Araucariaceae	Lower to upper ?mesotherm
<i>Dacrydiomites preacupressinoides</i>	Podocarpaceae	Raine et al. (2011)	<i>Dacrydium cupressinum</i>	
<i>Podocarpidites ellipticus</i>	Podocarpaceae	Raine et al. (2011)	Podocarpaceae	Microtherm to ?megatherm
<i>Podocarpidites</i> spp.	Podocarpaceae	Truswell and Macphail (2009)	Podocarpaceae	Microtherm to ?megatherm
<i>Dacrycarpites australiensis</i>	Podocarpaceae	Truswell and Macphail (2009)	Podocarpaceae	Upper microtherm to lower mesotherm
<i>Podocarpidites marwickii</i>	Podocarpaceae	Raine et al. (2011)	Podocarpaceae	Microtherm to ?megatherm
<i>Phyllocladidites mawsonii</i>	<i>Lagarostrobos</i>	Raine et al. (2011)	<i>Lagarostrobos</i>	Upper microtherm to lower mesotherm
<i>Phyllocladidites reticulascaccatus</i>	Podocarpaceae	Raine et al. (2011)	Podocarpaceae	
<i>Microacachrydites antarcticus</i>	Podocarpaceae	Raine et al. (2011)	<i>Microcachrys</i>	Upper microtherm to lower mesotherm
<i>Taxodiaceaeopollenites hiatus</i>	Cupressaceae	Raine et al. (2011)	Cupressaceae	
<i>Microalatiidites</i> sp.	Podocarpaceae	Raine et al. (2011)	Podocarpaceae	Upper microtherm to lower mesotherm
<b>Angiosperms</b>				
<i>Malvacipollis subtilis</i>	Euphorbiaceae	Raine et al. (2011)	Euphorbiaceae	
<i>Myricipites harrisi</i>	Casuarinaceae	Raine et al. (2011), Macphail (2007)	<i>Gymnostoma</i>	Lower mesotherm to megatherm
<i>Nothofagidites flemingii</i>	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Raine et al. (2011)	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites</i> spp.	<i>Nothofagus</i>	Raine et al. (2011)	<i>Nothofagus</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites emarcidus</i> complex	<i>Nothofagus</i> subg. <i>Brassospora</i>	Truswell and Macphail (2009)	<i>Nothofagus</i> subg. <i>Brassospora</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites falcatus</i>	<i>Nothofagus</i> subg. <i>Brassospora</i>	Raine et al. (2011)	<i>Nothofagus</i> subg. <i>Brassospora</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites lachlaniae</i> complex	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Raine et al. (2011)	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites waipawaensis</i>	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Raine et al. (2011)	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Upper microtherm to lower mesotherm
<i>Nothofagidites asperus</i>	<i>Nothofagus</i> subg. <i>Lophozonia</i>	Truswell and Macphail (2009)	<i>Nothofagus</i> subg. <i>Lophozonia</i>	Upper microtherm to lower mesotherm

Table 1. Continued.

Fossil taxon	Botanical affinity	Source	Selected NLR for climate analysis	Inferred climate range (Maephail, 2007)
<i>Nothofagidites senectus</i>	<i>Nothofagus</i>	Raine et al. (2011)	<i>Nothofagus</i>	
<i>Nothofagidites brachyspinulosus</i>	<i>Nothofagus</i> subg. <i>Fuscospora</i>	Raine et al. (2011)	<i>Nothofagus</i> subg. <i>Fuscospora</i>	
<i>Protacidites crassus</i>	Proteaceae	Raine et al. (2011)	Proteaceae	Lower to upper mesotherm
<i>Protacidites pachypolus</i>	Proteaceae	Maephail and Hill (2018)	Proteaceae	Lower to upper mesotherm
<i>Protacidites pseudomoides</i>	Proteaceae	Raine et al. (2011)	<i>Carruvonia</i>	Lower to upper mesotherm
<i>Protacidites leightoni</i>	Proteaceae	Truswell and Maephail (2009)	Proteaceae	Lower to upper mesotherm
<i>Protacidites reticulatus</i>	Proteaceae	Truswell and Maephail (2009), Raine et al. (2011)	Proteaceae	Lower to upper mesotherm
<i>Protacidites scaboratus</i>	Proteaceae	Raine et al. (2011)	Proteaceae	Lower to upper mesotherm
<i>Protacidites similis</i>	Proteaceae	Raine et al. (2011)	Proteaceae	Lower to upper mesotherm
<i>Protacidites parvus</i>	Proteaceae	Bowman et al. (2014), Raine et al. (2011)	<i>Bellendena</i>	Lower to upper mesotherm
<i>Periporopollenites polyoratus</i>	Caryophyllaceae, Trimeniaceae	Raine et al. (2011)	Caryophyllaceae	
<i>Parsonsites psilatus</i>	<i>Parsonsia</i>	Raine et al. (2011)	<i>Parsonsia</i>	
<i>Spinizonocolpites</i> sp.	Areaceae	Raine et al. (2011), Kumaran et al. (2011)	Areaceae	Upper mesotherm to megatherm
<i>Tricolpites triobolatus</i>	Scrophulariaceae, Convolvulaceae	Raine et al. (2011)	<i>Hebe</i>	Lower to upper mesotherm
<i>Myrtacidites</i> cf. <i>parvus</i>	Myrtaceae	Raine et al. (2011)	Myrtaceae	
<i>Malvacearumpollis mannemansis</i>	Malvaceae	Raine et al. (2011)	Malvaceae	
<i>Nupharipollis mortonensis</i>	Araceae, Nymphaeaceae	Raine et al. (2011)	<i>Nuphar</i>	Upper mesotherm to megatherm
<i>Sapotaceoidapollenites</i> cf. <i>lantzianus</i>	Sapotaceae	Raine et al. (2011)	Sapotaceae	
Cryptogams				
<i>Cyathidites australis</i>	Cyatheaceae	Raine et al. (2011), Maephail et al. (1994)	Cyatheaceae	Upper microtherm to lower mesotherm
<i>Cyathidites minor</i>	Cyatheaceae	Raine et al. (2011)	Cyatheaceae	Upper microtherm to lower mesotherm

Table 1. Continued.

Fossil taxon	Botanical affinity	Source	Selected NLR for climate analysis	Inferred climate range (Macphail, 2007)
<i>Cyatidites</i> sp.	Cyatheaceae	Raine et al. (2011)	Cyatheaceae	Upper microtherm to lower mesotherm
<i>Laevigatosporites ovatus</i>	Blechnaceae	Raine et al. (2011), Truswell and Macphail (2009)	Blechnaceae	
<i>Osmundacidites wellmanii</i>	Osmundaceae	Raine et al. (2011)	<i>Todea</i>	
<i>Osmundacidites</i> sp.	Osmundaceae	Raine et al. (2011)	Osmundaceae	
<i>Baculatisporites comaumensis</i>	Osmundaceae, Hymenophyllaceae	Raine et al. (2011), Truswell and Macphail (2009), Macphail and Cantrill (2006)	<i>Hymenophyllum</i>	
<i>Gleicheniidites senonicus</i>	Gleicheniaceae	Truswell and Macphail (2009), Raine et al. (2011)	Gleicheniaceae	
<i>Gleicheniidites</i> spp.	Gleicheniaceae	Truswell and Macphail (2009)	Gleicheniaceae	
<i>Dictyophyllidites arcuatus</i>	Gleicheniaceae	Raine et al. (2011)	Gleicheniaceae	
<i>Kuylisporites waterbolkkii</i>	Cyatheaceae	Raine et al. (2011)	Cyatheaceae	Upper microtherm to lower mesotherm
<i>Clavifera rudis</i>	Gleicheniaceae	Raine et al. (2011)	Gleicheniaceae	
<i>Clavifera triplex</i>	Gleicheniaceae	Raine et al. (2011), Truswell and Macphail (2009)	Gleicheniaceae	
<i>Laevigatosporites major</i>	Blechnaceae	Truswell and Macphail (2009), Raine et al. (2011), Macphail and Hill (2018)	Blechnaceae	
<i>Stereisporites antiquasporites</i>	Sphagnaceae	Truswell and Macphail (2009)	<i>Sphagnum</i>	± microtherm
<i>Ceratospores equalis</i>	Selaginellaceae	Raine et al. (2011)	<i>Selaginella</i>	
<i>Cibotiidites tuberculiformis</i>	Schizaeaceae	Raine et al. (2011), Daly et al. (2011)	Schizaeaceae	
<i>Polypodiisporites radiatus</i>	Polypodiaceae	Raine et al. (2011)	Polypodiaceae	
<i>Retriletes austroclavatifidites</i>	Lycopodiaceae	Raine et al. (2011)	<i>Lycopodium</i>	



the WorldCLIM climate surface (Fick and Hijmans, 2017). The datasets are then filtered to remove multiple entries per climate grid cell, plants whose botanical affinities are vague, doubtful, or redundant, and occurrences termed exotic (e.g. garden plants). Filtering was performed to avoid bias in the probability function that would likely lead to results leaning towards a particular location (Reichgelt et al., 2018). To test the robustness of the dataset, bootstrapping was applied, which was followed by the calculation of the likelihood of a taxon that occurs at a specific climate variable using the mean and standard deviation of the modern range of each taxon (Kühl et al., 2002; Willard et al., 2019). For a more detailed explanation of this method, see Willard et al. (2019) and Klages et al. (2020).

## 2.4 Quantitative and statistical analyses

Diversity indices (rarefaction, Shannon diversity index, equitability) were generated using the PAST statistical software (Hammer et al., 2001) with sample counts of  $\geq 75$  individuals. Rarefaction is an interpolation technique that is used to compare taxonomic diversity in samples of different sizes (Birks and Line, 1992; Birks et al., 2016). Rarefaction analyses using sample counts of  $\geq 75$  and  $\geq 100$  showed similar diversity trends. We, however, settled on counts with  $\geq 75$  individual grains because they offered an added advantage of filling in the gaps that would have been created if only samples with counts of  $\geq 100$  grains were used, thereby increasing the resolution of the studied section. The Shannon diversity index ( $H$ ) is a measure of diversity that considers the number of individuals as well as the number of taxa and the evenness of the species present (Shannon, 1948).  $H$  ranges from 0 for vegetation communities with a single taxon to higher values where taxa are evenly distributed (Legendre and Legendre, 2012). Equitability ( $J$ ), on the other hand, measures the level of abundance and how the species are distributed in an assemblage. Low  $J$  values indicate the dominance of a few species in the population (Hayek and Buzas, 2010).

Pollen zones (PZ) were defined following stratigraphically constrained analysis (CONISS; Grimm, 1987) in Tilia (version 2.6.1) using the total sum of squares with chord distance square-root transformation (Cavalli-Sforza and Edwards, 1967). In addition, we used detrended correspondence analysis (DCA; Hill and Gauch, 1980) sample scores to measure the sample-to-sample variance. DCA sample scores were generated using the *vegan* package (Oksanen et al., 2019) of the R statistical software (R Core Team, 2019).

## 3 Results

### 3.1 Palynological results from ODP Site 1172

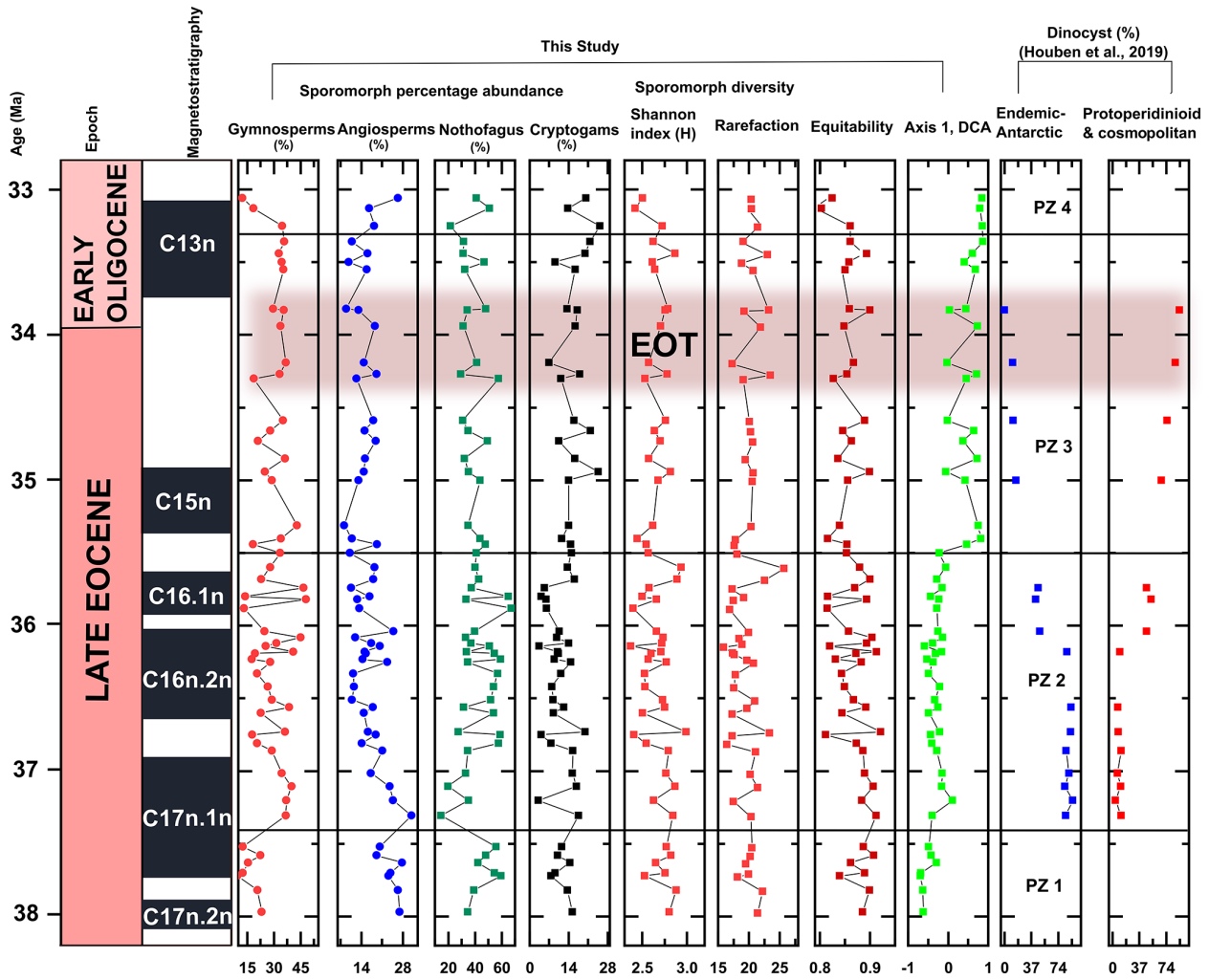
The late Eocene–early Oligocene samples from the East Tasman Plateau (ODP Site 1172) yielded moderately to well-preserved sporomorphs. Eighty-one (81) individual sporomorph taxa were recorded from the 57 productive samples across the studied section. The sporomorph record is dominated by *Nothofagidites* spp., which makes up between 38 % to 48 % of all non-reworked sporomorphs across the studied interval (Fig. 2). *Podocarpidites* spp., *Myricipites harrisii*, *Cyathidites* spp., *Phyllocladidites mawsonii*, and *Araucariacites australis* form significant components of the palynoflora and occur with varying frequency (Fig. 2).

Based on results from rarefaction, the average diversity for the entire studied section was  $20.1 \pm 1.74$  taxa per sample of 75 individuals. The sporomorph record based on CONISS is grouped into four pollen zones (PZ; Fig. 2), PZ 1 (early late Eocene; 37.97–37.52 Ma), PZ 2 (late Eocene–latest Eocene; 37.30–35.60 Ma), PZ 3 (latest Eocene–earliest Oligocene 35.50–33.36 Ma), and PZ 4 (earliest Oligocene; 33.25–33.06 Ma). All four zones consist of characteristic palynoflora assemblages that are described below. Taxa names in brackets refer to the NLRs.

#### 3.1.1 Pollen zone 1 (37.97–37.52 Ma; 7 samples)

Pollen zone 1 is dominated by *Nothofagidites* spp. (*Nothofagus*), which accounts for  $\sim 48\%$  of all non-reworked palynomorphs. Taxa belonging to the *Brassospora* ( $\sim 28\%$ ) subgenus of *Nothofagus* make up the most abundant component, followed by *Fuscospora* (19%) and *Lophozonia* (1%), respectively. Other angiosperms (non-*Nothofagus*) on average account for 24% of all sporomorphs. These are represented mainly, in order of decreasing occurrence, by *Myricipites harrisii* (*Gymnostoma*), *Proteacidites pseudomoides* (*Carnarvonina*), *Proteacidites* spp., *Spinizonocolpites* spp. (Arecaceae), *Malvacearumpollis mannanensis* (Malvaceae), and *Malvacipollis* spp. (Euphorbiaceae). The abundance of gymnosperms is generally low throughout PZ 1 and accounts for about 16% of all non-reworked palynomorphs. These are also represented mainly, in order of decreasing occurrence, by *Podocarpidites* spp. (Podocarpaceae), *Phyllocladidites mawsonii* (*Lagarostrobos*), *Dacrydiumites praecupressinoides* (*Dacrydium*), and *Araucariacites australis* (Araucariaceae). Ferns and mosses account for about 12% of the total sporomorphs and are represented by *Cyathidites* spp. (Cyatheaceae), *Dictyophyllidites* sp. (Gleicheniaceae), *Gleicheniidites* sp. (Gleicheniaceae), *Laevigatosporites* spp. (Blechnaceae), and *Stereisporites* sp. (*Sphagnum*).

Quantitatively, the sporomorph diversity for this zone, based on rarefied values, is  $19.65 \pm 1.32$  species per sample of 75 individuals. With respect to the diversity indices, the yields for Shannon diversity ( $H$ ) are between 2.33 and 2.69,



**Figure 3.** Sporomorph percentage abundances, diversity, and detrended correspondence analysis (DCA) results for ODP Site 1172. Percentage abundances for the major groups (gymnosperms, other angiosperms, *Nothofagus*, and cryptogams) are presented for all samples with pollen counts  $\geq 75$  grains. The DCA results are derived from the sample scores of axis 1 (which measures the sample-to-sample variance) and show four distinct compositional groupings, as observed with CONISS, for the late Eocene–early Oligocene Site 1172 samples. Diversity is calculated based on Sander’s rarefaction analysis with samples rarefied at 75 grains/individuals. Relative percentage abundances of Antarctic-endemic and protoperidinioid dinoflagellate cysts, the magnetostratigraphy, and the age model are after Houben et al. (2019).

**Table 2.** Summary of quantitative species diversity and DCA axis 1 sample scores between the late Eocene and early Oligocene from ODP Site 1172.

Analysis	Pollen zone 1		Pollen zone 2		Pollen zone 3		Pollen zone 4	
	Mean	(SD)	Mean	(SD)	Mean	(SD)	Mean	(SD)
Rarefaction (75 individuals)	19.65	1.32	19.44	2.49	20.15	1.79	21.16	1.37
Rarefaction (100 individuals)	22.29	1.87	21.78	2.85	22.52	2.31	23.75	1.32
Shannon index ( <i>H</i> )	2.57	1.12	2.56	0.22	2.58	0.12	2.54	0.15
Equitability ( <i>J</i> )	0.85	0.02	0.86	0.04	0.85	0.02	0.83	0.03
DCA (axis 1 sample scores)	−0.55	0.15	−0.29	0.15	0.44	0.33	0.83	0.03

averaging at  $2.57 \pm 1.12$ . Equitability ( $J$ ) scores are set between 0.81 and 0.88, with an average of  $0.85 \pm 0.02$  (Fig. 3; Table 2).

### 3.1.2 Pollen zone 2 (37.30–35.60 Ma; 27 samples)

PZ 2 sees the decline of *Nothofagidites* spp. to about 42 %. The *Brassospora* type remains the dominant *Nothofagus* subgenus, but with a substantial decline in abundance from about 28 % in PZ 1 to 22 % in PZ 2. The *Fuscospora* and *Lophozonia* subgenera, however, accounted for 19 % and 1 %, respectively (Fig. 2). In comparison to PZ 1, other angiosperms (non-*Nothofagus*) see a decline from about 24 % to 17 %. In order of decreasing abundance, the most significant taxa among non-*Nothofagus* angiosperms are *Myricipites harrisii* (*Gymnostoma*), *Proteacidites* spp. (Proteaceae), *Malvacearumpollis mannanensis* (Malvaceae), and *Malvacipollis* spp. (Euphorbiaceae). A sharp decline in *Proteacidites pseudomoides* (*Carnarvon*) is coupled with the disappearance of *Spinizonocolpites* spp. (Arecaceae). Gymnosperms, on the other hand, almost double in relative abundance from about 16 % in PZ 1 to over 29 % in PZ 2. In order of decreasing abundance, gymnosperm taxa are dominated by *Podocarpus* spp. *Araucariacites australis* (Araucariaceae), *Dacrydiumites praecupressinoides* (*Dacrydium*), and *Phyllocladidites mawsonii* (*Lagarostrobos*). *Microcachryidites antarcticus* (*Microcachrys*) is a taxon that first appears in this zone and forms an important component ( $\sim 11$  %) of the gymnospermous assemblage. In addition to the above, cryptogams decline slightly in this zone, accounting for roughly 10 % of the total sporomorphs. The main members of this group are *Cyathidites* spp. (Cyatheaceae), *Gleichenioidites* (*Gleicheniaceae*), and *Laevigatosporites* spp. (Blechnaceae).

This zone has lower diversity than PZ 1. Based on rarefied values, the average diversity for PZ 2 is  $19.44 \pm 2.49$  species per sample of 75 individuals. The results for the Shannon diversity index ( $H$ ) are between 2.15 and 2.97, averaging at  $2.56 \pm 0.22$ . Equitability is set between 0.78 and 0.93, with an average of  $0.86 \pm 0.04$  (Fig. 3; Table 2).

### 3.1.3 Pollen zone 3 (35.50–33.36 Ma; 20 samples)

Zone 3 shows a further decline in *Nothofagidites* spp. to  $\sim 38$  %. However, the *Brassospora*-type *Nothofagus* sees a slight increase in abundance, while the *Fuscospora*-type *Nothofagus* declines sharply from the peak of 19 % observed in PZ 2 to 12 %. The *Lophozonia* type remains stable ( $\sim 1$  %). Other angiosperms (non-*Nothofagus*) see a slight decline and account for  $\sim 14$  % of all non-reworked sporomorphs. These are represented mainly by *Myricipites harrisii* (*Gymnostoma*) and *Proteacidites* spp. (Proteaceae), *Malvacipollis* spp. (Euphorbiaceae), and *Malvacearumpollis mannanensis* (Malvaceae). Another important observation for this interval is the re-appearance of *Spinizonocolpites* spp. (Arecaceae) and *Proteacidites pseudomoides* (*Carnar-*

*von*). However, in contrast to PZ 1, *Spinizonocolpites* spp. are not consistently present. Gymnosperms increase slightly in this zone, accounting for  $\sim 31$  %. The gymnosperms remain dominant with *Podocarpidites* spp. (Podocarpaceae). However, other important taxa such as *Araucariacites australis* (Araucariaceae), *Phyllocladidites mawsonii* (*Lagarostrobos*), and *Microcachryidites* (*Microcachrys*) decline. *Dacrydiumites praecupressinoides* (*Dacrydium*) reaches its peak abundance in this zone. Cryptogams significantly increase in abundance and are represented, in order of abundance, by *Cyathidites* spp. (Cyatheaceae), *Laevigatosporites* spp. (Blechnaceae), *Osmundacidites* (Osmundaceae), *Polypodiisporites radiatus* (Polypodiaceae), and *Clavifera* spp. (Gleicheniaceae).

Based on rarefied values, the average diversity for this PZ is  $20.15 \pm 1.79$  species per sample. The results for Shannon diversity ( $H$ ) are between 2.37 and 2.80, averaging at  $2.58 \pm 0.12$ . Equitability ( $J$ ) is set between 0.81–0.91, averaging at  $0.85 \pm 0.02$  (Fig. 3; Table 2).

### 3.1.4 Pollen zone 4 (33.25–33.06 Ma; 3 samples)

The percentage abundances of *Nothofagidites* spp. (*Nothofagus*), including *Brassospora* ( $\sim 23$  %), *Fuscospora* (12 %), and *Lophozonia* types, remain unchanged, whereas other angiosperm percentages increase substantially from 14 % in PZ 3 to  $\sim 20$  %. In order of decreasing abundance, these are represented by *Myricipites harrisii* (*Gymnostoma*) and *Proteacidites pseudomoides* (*Carnarvon*). PZ 4 also sees the emergence of new angiosperms such as *Sapotaceoidae-pollenites* cf. *latizonatus* (Sapotaceae) and *Parsonsidites psilatus* (*Parsonsia*). Gymnosperms, however, see a sharp decline in this interval, accounting for about 21 % of the total sporomorph taxa, with *Podocarpidites* spp. (Podocarpaceae) and *Dacrydium praecupressinoides* (*Dacrydium*) as the main components. *Microcachryidites antarcticus* (*Microcachrys*), *Araucariacites australis* (Araucariaceae), and *Phyllocladidites mawsonii* (*Lagarostrobos*) show significant declines, whereas cryptogams increase to  $\sim 20$  %. The cryptogams are represented, in order of decreasing abundance, by *Cyathidites* spp. (Cyatheaceae), *Laevigatosporites* spp. (Blechnaceae), *Dictyophyllidites* sp. (Gleicheniaceae), and *Cibotidites tuberculiformis* (Schizaeaceae).

Average diversity ( $21.16 \pm 1.37$  species per sample) is slightly higher than in PZ 3. The results for Shannon diversity ( $H$ ) are between 2.42 and 2.72, averaging at  $2.54 \pm 0.15$ . Equitability ( $J$ ) is set between 0.80–0.87, averaging at  $0.83 \pm 0.03$  (Fig. 3; Table 2).

## 4 Discussion

### 4.1 Vegetation composition and altitudinal zonation

Throughout the studied section, abundant *Nothofagidites* spp. and common *Podocarpidites* spp. *Myricipites har-*

*risii* and *Phyllocladidites mawsonii* indicate the presence of *Nothofagus*-dominated temperate rainforest (Truswell and Macphail, 2009; Bowman et al., 2014), which likely grew across lowland and mid-altitude elevations in eastern Tasmania. The occurrence of *Araucariacites australis*, *Microcachrydites antarcticus*, and *Proteacidites parvus* may also suggest that a component of the sporomorph assemblage reflects higher altitudes with more open forest conditions (Macphail, 1999; Kershaw and Wagstaff, 2001). In addition, pollen taxa belonging to *Arecaceae*, *Gymnostoma*, and *Carnarvonnia* indicate the existence of a paratropical vegetation community that grew in sheltered lowland and coastal areas (Huurdemans et al., 2021). The paratropical rainforest likely occupied lowlands and coastal areas, while temperate rainforest likely grew at higher elevations, similar to vegetation communities that prevailed on Wilkes Land and Tasmania during the early to mid-Eocene (Pross et al., 2012; Contreras et al., 2013, 2014). The existence of different vegetation communities with NLRs that currently grow at different temperatures and elevations suggests that vegetation across eastern Tasmania were subject to climatic gradients related to differences in elevation and/or distance to the coastline. This is supported by reports of a topographic divide between sites facing the cool Tasman Current (Gippsland basin, eastern Tasmania) and the westerly located south Australian basins (Holdgate et al., 2017), which may have served as the locations for higher-altitude temperate forest taxa. The following sub-sections further describe each of these vegetation communities in detail.

#### 4.1.1 Lowland to mid-altitude *Nothofagus*–*Podocarpus* rainforest

Abundant *Nothofagidites* spp. and common *Podocarpidites* spp., *Myricipites harrisii*, *Phyllocladidites mawsonii*, and *Cyatheaceae* give an indication of a lowland to mid-altitude *Nothofagus*–*Podocarpus*-dominated rainforest thriving under high-precipitation regimes ( $\text{MAP} > 1300 \text{ mm yr}^{-1}$ ) in Tasmania during the late Eocene to the earliest Oligocene. The main canopy is primarily made up of *Nothofagidites* spp. (*Nothofagus*/southern beech) and podocarps (*Dacrydi-umites*, *Podocarpidites*, *Dacrycarpites*) with rare Cupressaceae trees. Southern beech forests can either occur as pure stands or a mixed forest, making the definition and recognition of regional or local forest types from fossil pollen and spores challenging. Today, pure beech stands in New Zealand are mostly montane to subalpine, and lowland mixed beech forests are associated with diverse broadleaf angiosperms and canopy-emergent gymnosperms (Ogden et al., 1996). Following Dettmann et al. (1990), we categorize *Nothofagidites* pollen taxa into the subgenera *Brassospora*, *Fuscospora*, and *Lophozonia*. Extant *Fuscospora* and *Lophozonia* types thrive in cool-temperate conditions in Tasmania, southeastern Australia, New Zealand, and southern South America (Hill, 1994, 2017; Veblen et al., 1996; Read et

al., 2005), while the *Brassospora* type are today restricted to warm-temperate–subtropical conditions in New Guinea and New Caledonia (Hill and Dettmann, 1996; Veblen et al., 1996). These *Brassospora*-type *Nothofagus* grow today at lower to mid-altitudes that receive high and consistent rainfall, but also in montane and subalpine areas (typically above 500 m), pointing to their wide ecological and climate tolerance ( $\text{MAT: } 10.6 \text{ to } 23.5 \text{ }^\circ\text{C}$ ; Read et al., 2005).

*Myricipites harrisii* (Casuarinaceae) has two potential NLRs, *Casuarina/Allocasuarina* and *Gymnostoma*. *Casuarina/Allocasuarina* have xeromorphic features indicating adaptation to an arid climate with frequent fires (Hill, 2017; Lee et al., 2016; Hill et al., 2020). We selected the rainforest clade *Gymnostoma* as the most likely NLR of our fossil taxon *Myricipites harrisii*, based on the subtropical affinities of the associated palynoflora. This is also supported by Paleogene vegetation reconstruction for southeastern Australia based on macrofossil remains, which indicates rainforest communities (Christophel et al., 1987; Macphail et al., 1994; Hill, 2017), with *Gymnostoma* being common from the Paleocene to Oligocene and later being replaced by *Casuarina/Allocasuarina* (sclerophyll taxa) in the Miocene (Evi et al., 1995; Boland et al., 2006; Holdgate et al., 2017; Hill et al., 2020).

*Dacrydium cupressinum* is suggested as the most likely NLR of *Dacrydi-umites praecupressinoides* (rimu; Raine et al., 2011). Today, *Dacrydium cupressinum* occurs as a minor component in the kauri forest of Northland, New Zealand and as emergent taxa commonly associated with *Agathis australis* (Araucariaceae) and *Podocarpus totara* (Farjon, 2010). The NLR of *Phyllocladidites mawsonii*, *Lagarostrobos franklinii* (Tasmanian Huon pine; Raine et al., 2011), is very abundant at Site 1172. *Lagarostrobos* are evergreen cool-temperate riparian trees that grow in Tasmania close to riverbanks (Farjon, 2010; Hill, 1994, 2017). Apart from forming groves that mark stream courses at low altitudes (Hill and Macphail, 1983; Farjon, 2010), they may also be found away from water courses on wet hillsides in temperate forest (Farjon, 2010; Bowman et al., 2014). *Lagarostrobos* is one of the most common gymnosperms at ODP Site 1172, and its percentage abundance is similar to those recovered from wells offshore Gippsland Basin (the Gropper-1, Mullet-1, and Bluebone-1 wells; Partridge, 2006). *Lagarostrobos* occurs with even higher percentages in the Middle *Nothofagidites asperus* Zone of the terrestrial record of the Gippsland Basin, where it appears to be overrepresented (Holdgate et al., 2017).

The two possible NLR relatives for *Proteacidites pseudo-moides* are *Carnarvonnia* and *Lomatia*. *Carnarvonnia* thrives in warm-temperate to paratropical areas such as wet northeastern Australia (Mabberley, 1997; Cooper and Cooper, 2004) and grows into large trees (Hyland, 1995). *Lomatia* grows as shrubs and small trees in remnant gallery warm-temperate rainforests, for example along creek lines on sandstones in Northern Sydney (Bowman et al., 2014; Myer-



scough et al., 2007). *Carnarvonia* was selected as the likely NLR relative because it significantly increases in intervals where warmth-loving taxa such as Arecaceae, *Brassospora*-type *Nothofagus*, Gleicheniaceae, and Cyatheaceae thrive.

#### 4.1.2 High-altitude temperate rainforest and shrubland

Components of the palynoflora that reflect higher-altitude and more open vegetation on soils with low fertility are *Araucariacites australis*, *Proteacidites parvus*, and *Microcachryidites antarcticus* (Kershaw and Wagstaff, 2001; Macphail et al., 1999). Today, Araucariaceae trees grow in cool-temperate forests in Chile and Argentina and extend to the tree line (Veblen et al., 1996; Sanguinetti and Kitzberger, 2008). In the Andes, trees belonging to Araucariaceae are found at altitudes of 600–800 m a.s.l., receive high amounts of annual rainfall (2000–3000 mm yr<sup>-1</sup>), and experience hot and dry spells in summer (Farjon, 2010). Araucariaceae build pure stands at higher altitudes or mixed Valdivian rainforest at lower altitudes (Farjon, 2010). Increases in araucarian sporomorph taxa between 37.30–35.60 Ma in Tasmania give an indication of a dense, emergent cover of Araucariaceae thriving in relatively dry environments (Kershaw and Wagstaff, 2001). *Microcachrys* (Raine et al., 2011), the nearest living relative of *Microcachryidites antarcticus*, is a creeping shrub that grows in alpine/subalpine areas and is today restricted to western Tasmania under boreal to cool-temperate conditions (Truswell and Macphail, 2009; Biffin et al., 2012; Carpenter et al., 2011). Therefore, increases in this Tasmanian endemic alpine shrub (*Microcachrys*) from 37.30 to 35.60 Ma, together with *Bellendena* (a low-growing protea shrub; NLR of *Proteacidites parvus*), and Araucariaceae (emergent canopy), suggest that the vegetation that thrived at higher altitudes in Tasmania preferred cool-temperate conditions.

#### 4.2 Subtropical vegetation and early late Eocene cooling from 37.97 to 35.60 Ma

Throughout PZ 1 (37.97–37.52 Ma), abundant *Nothofagus* (especially the *N. brassii* type) with secondary Podocarpaceae and *Gymnostoma* and minor Arecaceae, *Carnarvonia*, and cryptogams suggest the presence of a temperate *Nothofagus*-dominated rainforest with subtropical elements. Sporomorph-based climate estimates indicate these forests grew under a MAT of 14.2–15.1 °C and a MAP of 1467–1681 mm yr<sup>-1</sup> (Fig. 4). The vegetation-based summer temperature reconstructions of ca. 18.5 °C closely corroborate the brGDGT-biomarker reconstructions from the same site (Bijl et al., 2021), supporting the notion of a potential seasonal bias of this palaeothermometer (Contreras et al., 2014; Naafs et al., 2017). The warmth-loving taxa formed the main lowland forest components occupying sheltered areas and lowland subtropical coastal zones (Dowe, 2010; Car-

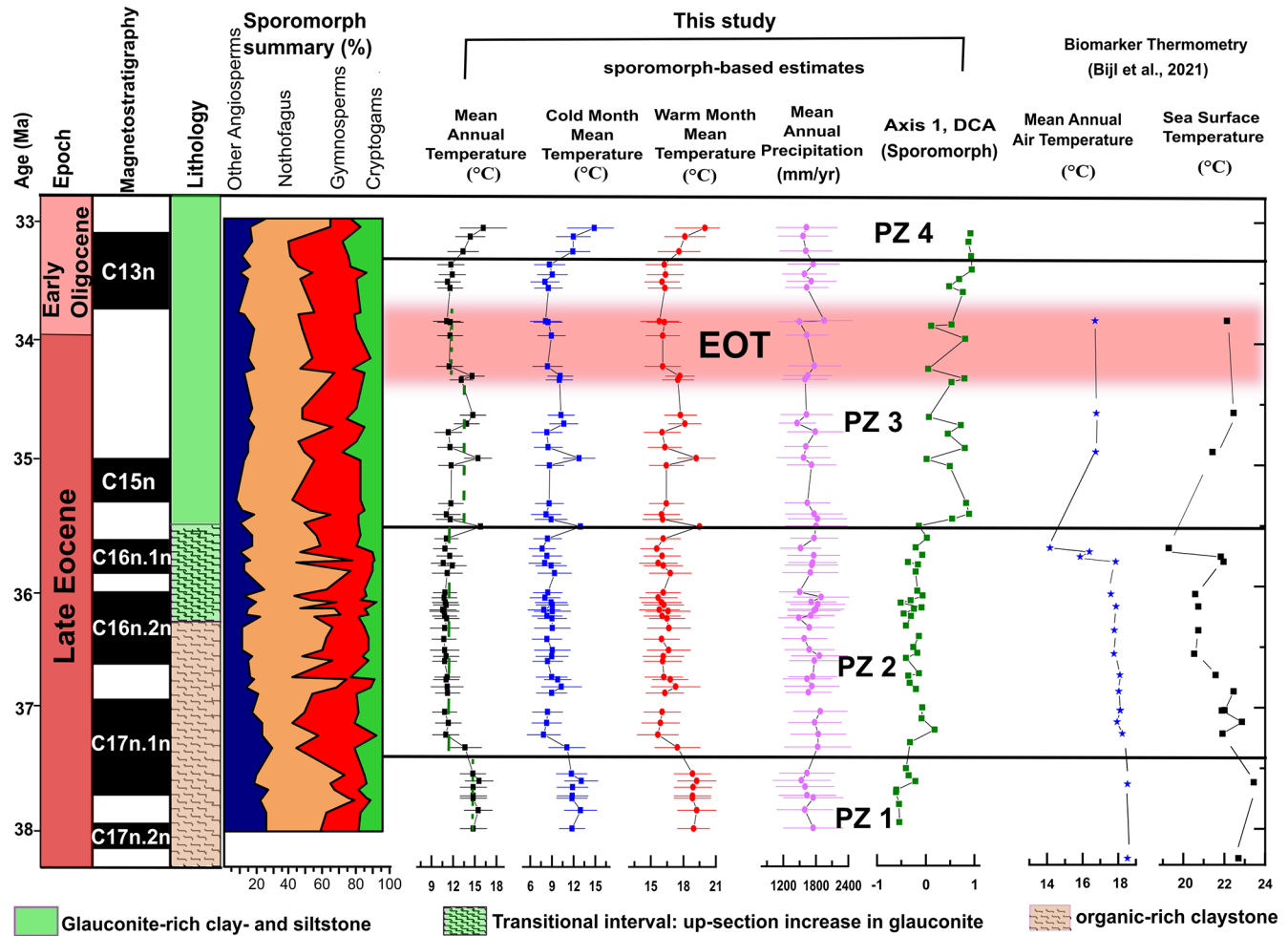
penter et al., 2012; Tripathi and Srivastava, 2012; Verma et al., 2020)

and swamps (Kershaw, 1988). Sporomorph-based temperature estimates yield a cold month mean temperature (CMMT) of well above freezing (11.2–12.5 °C; Fig. 4). The decline and, to a large extent, the absence of cool-temperate taxa, coupled with persistent warm-temperate (12–17 °C; Emanuel et al., 1985) to subtropical (17–24 °C; Emanuel et al., 1985) taxa, further point to the expansion of warm-temperate–paratropical rainforest up into the mid-altitudes and uplands.

The *Nothofagus*-dominated rainforest continued into PZ 2 (37.30–35.60 Ma). However, at ~37.30 Ma, a distinct environmental change occurred, leading to a drop in, and in some instances the demise of, warm-temperate and subtropical taxa (*Nothofagus* subgenus *Brassospora*, *Carnarvonia*, Arecaceae; Fig. 2). This vegetation change continued throughout PZ 2, along with a concomitant rise in the relative abundances of *Lagarostrobos* and *Microcachrys* and a decline in diversity (Table 2) ~3 Ma prior to the EOT. The increased occurrence of microthermal taxa points to a cool-temperate (southern beech)-dominated rainforest with secondary Podocarpaceae expanding into lowland regions previously occupied by mesothermal taxa. The late Eocene cool-temperate *Nothofagus*–Podocarpaceae-dominated rainforest has been suggested to resemble the modern Valdivian rainforest of Chile (Veblen, 1982; Cantrill and Poole, 2012; Bowman et al., 2014) and the cool-temperate *Nothofagus*-dominated rainforest with riparian *Lagarostrobos* restricted to river gullies in Victoria, Australia (Read and Hill, 1985) or on fertile soils in lowland Tasmania (Read and Hill, 1985; Macphail, 2007; Francis et al., 2008).

This interpretation is reflected in our sporomorph-based MAT estimates indicating a 2–3 °C decline in MAT (Fig. 4). Our findings also corroborate previous late Eocene studies throughout Australia indicating an increase in *Nothofagus* subgenus *Fuscospora* along with a substantial decline in *Brassospora*-type *Nothofagus* and the demise of most Proteaceae, Arecaceae, and most Australian angiosperms (Kemp, 1978; Kershaw, 1988; Christophel and Greenwood, 1989; Truswell, 1993; Martin, 1994, 2006; Macphail et al., 1994; Partridge and Dettmann, 2003; Korasidis et al., 2019). In line with the vegetation change, biomarker-based reconstruction from Site 1172 also indicates that SSTs decline by ca. 2–3 °C starting around 37.5 Ma (Fig. 4). However, the cooling indicated by both independent proxies is not reflected in the lipid-biomarker-based terrestrial MAT estimates, and the reason for this disparate trend remains unknown.

The transition from a warm-temperate rainforest with paratropical elements to cool-temperate forests in the Tasmanian Gateway region also matches an early late Eocene (37.3 Ma) cooling in the Southern Ocean (Kerguelen Plateau) ~3 Myr prior to the EOT (Villa et al., 2008, 2014; Scher et al., 2014). The 2–3 °C sporomorph-based MAT (100–200 kyr) cooling around 37.3 Ma coincides with a regional



**Figure 4.** Comparison of our sporomorph-based climate estimates,  $MAAT_{soil}$  values based on MBT\*5me,  $TEX_{86}$ -based SSTs, and sample scores for DCA axis 1 from the late Eocene to early Oligocene of ODP Site 1172. Sporomorph-based estimates are based on the use of the nearest living relative (NLR) and probability density function (PDF). The range of each climate estimate represents the mathematical error and not the real range, and may have been a result of uncertainties associated with the use of the NLR approach. Broken green lines indicate average temperatures for sporomorph-based MATs. Biomarker thermometry data are from Bijl et al. (2021). The ~ 790 kyr interval corresponding to the EOT (34.44–33.65 Ma; Hutchinson et al., 2021) is indicated by the horizontal pink bar. The age model is after Houben et al. (2019).

transient (~ 140 kyr) cooling event at ODP Site 738 (Kerguelen Plateau) known as the Priabonian Oxygen Maximum (PrOM; Scher et al., 2014). The PrOM event, placed within magnetochron C17n.1n of the late Eocene, points to the temporary growth of ice sheets on East Antarctica based on a positive excursion of benthic  $\delta^{18}O$  (Scher et al., 2014). However, on the Kerguelen Plateau, differences in neodymium isotopic composition ( $\epsilon_{Nd}$ ) between bottom waters and terrigenous sediments point to changes in sediment provenance as opposed to changes in the reorganization of ocean currents (Scher et al., 2014). The transient 2–3 °C sporomorph-based MAT cooling phase is followed by a period with a sustained cooler climate from 37.2 to 35.6 Ma (Fig. 4). This sustained cooler climate may have caused the climate threshold of the frost-sensitive (subtropical) taxa to be exceeded,

hence their continued decline and demise. In the marine realm, Antarctic-endemic dinoflagellate cysts (e.g. *Deflandrea antarctica*, *Vozzhennikovia* spp., and *Enneadocysta dictyostila*) become dominant at Site 1172 (Fig. 3; Stickley et al., 2004; Houben et al., 2019). The dominant Antarctic-endemic dinocysts in addition to general sea surface circulation models (Huber et al., 2004) point to the East Tasman Plateau and east Tasmania being bathed by relatively cool Antarctic-derived surface waters (Houben et al., 2019), which is consistent with  $TEX_{86}$ -based sea surface temperature records (~ 3–4 °C cooling; Houben et al., 2019; Bijl et al., 2021). Regionally, this sustained cool-temperate terrestrial MAT matches with oligotrophic conditions associated with low nutrients, stratification of the water mass, and increased efficiency of the ocean’s biological pump, which



favoured cooling as a result of carbon being sequestered from surface water in the Southern Ocean (Villa et al., 2008, 2014).

Close to the top of PZ 2 (35.7 Ma; Fig. 4), branched GDGT (glycerol dialkyl glycerol tetraether)-based MATs and SSTs show strong and rapid cooling, which is not mirrored by the pollen-based climate estimates. However, strong fluctuations in gymnosperms and an increase in cryptogams (Figs. 2 and 3) and diversity towards the top of PZ 2 indicate increasing environmental disturbance, which might be linked to the recorded change in lipid biomarker-based MATs. The rapid cooling most likely created gaps within the canopy, triggering an expansion of cryptogams. The divergence between the different proxy signals could be related to their different origins and transport mechanisms. Whereas the lipid biomarkers are strongly controlled by the depositional settings, including the river run-off, and the tectonic and geographic evolution (Bijl et al., 2021), the terrestrial palynological signal mainly consists of wind-dispersed pollen and spores. The long distance between the study site (ODP Site 1172) and mainland Tasmania (more than 100 km) in the Eocene makes a major influence of river/water-transported sporomorphs unlikely.

#### 4.3 Warm- and cold-temperate terrestrial climate fluctuation from 35.50–34.59 Ma

PZ 3 (35.50–33.36 Ma) is characterized by a major shift in sporomorph assemblages, represented by an increase in *Podocarpus*, declines in *Lagarostrobos*, *Microcachrys*, *Araucariaceae*, and *Fusca*-type *Nothofagus*, along with the re-emergence of subtropical and warm-temperate taxa. The peak in tree ferns, especially *Cyatheaceae*, indicate a period of disturbance (Vajda et al., 2001) within this interval of vegetation shift. However, we are not able to attribute the disturbance within this period (35.50–34.59 Ma) to an increase in fire frequencies as there is an absence of charcoal particles within our records. Sporomorph-based temperature reconstructions indicate several fluctuations between warm and cool climate phases with MATs between 10.6 and 15.3 °C (Fig. 4). In the regional Australo-Antarctic area, a similar phase of warming and cooling is observed in the late Eocene (35.8–34.7 Ma) climate records of Prydz Bay (Passchier et al., 2017; Tibbett et al., 2021) and southern Australia (Benbow et al., 1995). Again, pollen-based WMMTs at Site 1172 closely match lipid-biomarker-derived MATs (Fig. 4). In comparison, our sporomorph-based warm and cool climate fluctuation phase between 35.50 to 34.59 Ma is recorded as a recovery phase in the lipid-biomarker-based MAT reconstruction. The fluctuations of pollen-based temperature estimates may be at least partly caused by the proxy method, which relies on presence–absence data. However, a more detailed proxy comparison is hampered by the relatively low resolution of the lipid biomarker in PZ 3.

Expansions and restrictions of cool-temperate and warm-temperate forests, which indicate cooling and warming phases, are consistent with previous late Eocene geochemical, sedimentological, and palynological studies reporting an increase in sea surface temperature (TEX<sub>86</sub>-based SST; Houben et al., 2019; Bijl et al., 2021), widespread deposition of glauconite (Stickley et al., 2004), and increases in cosmopolitan and protoperidinioid dinocysts (Fig. 3; Stickley et al., 2004; Houben et al., 2019; Bijl et al., 2021). Though the glauconitic unit is interpreted as marking deepening and current inception due to a widening of the Tasmanian Gateway (Stickley et al., 2004), a more recent counterargument links the deposition of the greensand to atmospherically forced invigorated circulation in the Southern Ocean, which helped to prepare Antarctica for rapid expansion of ice (Houben et al., 2019) and a further circulation change ~2 Ma later (at the EOT). However, in addressing the deposition of greensands along the south Australian margin, ocean model studies (Baatsen et al., 2016) point to a further expansion of the eastward throughflow into the southwest Pacific Ocean. Our sporomorph-based MAT consequently showed an average 2 °C rise in temperature between 35.50 and 34.59 Ma, coinciding with earlier reports of the initial deepening of the Tasmanian Gateway (Stickley et al., 2004). This is further supported by the common appearance of low-latitude cosmopolitan dinoflagellate cyst taxa that, rather than being supplied by the East Australian Current, are reported to have been sourced from the throughflow associated with the eastward proto-Leeuwin Current (Huber et al., 2004; Stickley et al., 2004; Houben et al., 2019). These events, coupled with an ~2 °C recovery in SSTs (TEX<sub>86</sub>-based; Houben et al., 2019; Bijl et al., 2021) between 35.7 and 34.59 Ma, most likely point to an influence of warm surface waters associated with the Australo-Antarctic Gulf (AAG) at ODP Site 1172 (Houben et al., 2019), which is reported to have been close to land (eastern Tasmania; Stickley et al., 2004) at that time, thereby affecting the terrestrial climate and vegetation.

#### 4.4 EOT cooling and climate rebound in the earliest Oligocene from 34.30 to 33.06 Ma

At the onset of the EOT, our sporomorph record provides evidence for a return to a sustained cooler period spanning 34.30 to 33.82 Ma in Tasmania. This EOT cool phase coincides with the demise of *Spinizonocolpites* sp. (Arecaceae), a drop in *Cyatheaceae* and *Gleicheniaceae*, and slight increases in *Microcachrys* and *Lagarostrobos*. The palynoflora assemblage during the EOT is further characterized by a drop in overall angiosperm (non-*Nothofagus*) diversity, with gymnosperms and *Nothofagus* (*Brassospora* type) being common and co-dominating. Previous studies in southeast Australia (e.g. Macphail et al., 1994; Benbow et al., 1995; Holdgate et al., 2017; Lauretano et al., 2021) record a contemporaneous drop in angiosperm diversity and the final demise of *Arecaceae* (thermophilous elements) at the end of the Eocene

(Pole and Macphail, 1996; Martin, 2006). Quantitatively, our sporomorph-based MAT reconstruction records an  $\sim 2^\circ\text{C}$  decline across the EOT (Fig. 4), which coincides with  $\sim 2.4$  and  $5^\circ\text{C}$  cooling steps in southeastern Australia (MBT'5me-based MAATsoil; Laurentano et al., 2021) and East Antarctica (Prydz Bay; MBT'5me-based MAATsoil; Tibbett et al., 2021), respectively. This cooling in our terrestrial record further matches the principal geochemical signature of EOT in the marine realm: the  $\sim +1.5\%$  excursion of the oxygen isotope ratio of deep-sea benthic foraminifera (Zachos et al., 1996; Coxall et al., 2005; Pälike et al., 2006; De Vleeschouwer et al., 2017; Fig. 5) associated with global cooling (Zanazzi et al., 2007; Pearson et al., 2009; Pagani et al., 2011; Hutchinson et al., 2021; Tibbett et al., 2021). This cooling at the EOT has been linked to a global decline in atmospheric  $p\text{CO}_2$  (Pearson et al., 2009; Laurentano et al., 2021).

Between  $\sim 33.25$  and  $33.06$  Ma (PZ 4), the sporomorph-based climate estimates indicate a warming, with MATs between  $12.7$  and  $15.3^\circ\text{C}$  (Fig. 4). In addition, the presence of warmth-loving taxa, notably Sapotaceae, *Parsonsia* (Silkpod), and Polypodiaceae (subtropical epiphytes), further indicate a warming phase. The pollen flora resembles Oligocene warm-temperate *brassii* southern-beech-dominated forests of Karamu in the Waikato Coal Measures of New Zealand (Pocknall, 1985). The increase in *brassii*-type *Nothofagus* coupled with the appearance of Sapotaceae and subtropical epiphytes suggests that, at least locally on lowlands, eastern Tasmania was warm enough to accommodate warm-temperate vegetation in the earliest Oligocene. Previous earliest Oligocene studies in southeast Australia (Korasidis et al., 2019) show the presence of a cool-temperate rainforest community. The palynoflora on east Antarctica (Askin, 2000; Askin and Raine, 2000; Prebble et al., 2006; Tibbett et al., 2021) and northeast Tasmania (Hill and Macphail, 1983) suggest an early Oligocene cold-temperate *Nothofagus* (subgenus *Lophozonia* or *Fuscospora*)–Podocarpaceae vegetation. These palynoflora of northern Tasmania and east Antarctica are, however, reported to have most likely been made up of prostrate deciduous dwarf trees or small-stature closed southern beech/podocarp refugia with a vegetation community that likely struggled to survive (Askin, 2000; Askin and Raine, 2000; Prebble et al., 2006; Francis et al., 2008; Tibbett et al., 2021). However, rather than a regional scrub (e.g. in Antarctica), the slight increase in angiosperms (other than *Nothofagus*) and cryptogams point to a local warm-temperate forest growing along eastern Tasmania in the earliest Oligocene. Today, temperate forests in New Zealand and Tasmania host a diverse range of cryptogams, as compared to scrub communities that do not offer other taxa to thrive under the low, closed canopies (Prebble et al., 2006).

The terrestrial cooling observed across the EOT followed by a rapid recovery in the earliest Oligocene matches a partial return to warmer temperatures seen in previously reported

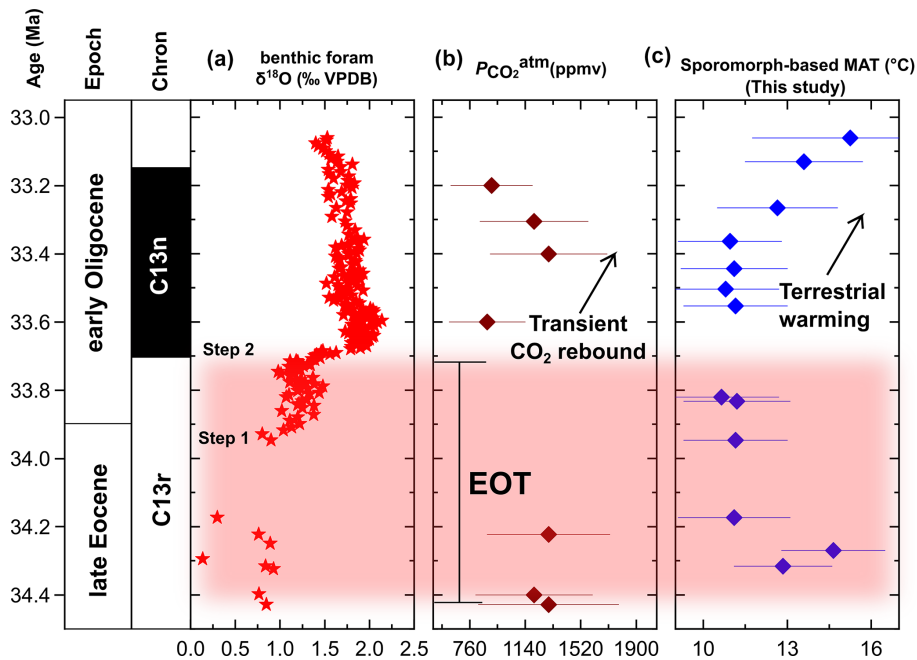
terrestrial (Colwyn and Hren, 2019; Laurentano et al., 2021) and marine studies (Katz et al., 2008; Lear et al., 2008; Liu et al., 2009; Houben et al., 2012). The synchronicity between terrestrial and marine records suggests that, in addition to localized events (sustained Tasmanian Gateway deepening and widening; Stickley et al., 2004), the EOT and earliest Oligocene ETP record may also be responding to a much wider regional or global event. The most common explanation for the global cooling (Zanazzi et al., 2007; Pagani et al., 2011; Hutchinson et al., 2021; Tibbett et al., 2021) across the EOT and the transient warming in the earliest Oligocene is the decline in concentration of atmospheric carbon dioxide ( $p\text{CO}_2$ ) and its recovery or rebound in the earliest Oligocene, respectively (Pearson et al., 2009; Heureux and Rickaby, 2015; Anagnostou et al., 2016; Fig. 5). Our results suggest that the warming, or at least the lack of sustained cooling following the EOT in eastern Tasmania, may be related to a combination of the  $p\text{CO}_2$  recovery (Pearson et al., 2009) and sustained Tasmanian Gateway deepening and widening (Stickley et al., 2004; Houben et al., 2019), allowing the influx of more warm surface waters from the AAG into the southwest Pacific, thereby affecting the terrestrial climate and vegetation along eastern Tasmania.

## 5 Conclusions

The late Eocene–early Oligocene vegetation reconstructed from terrestrial palynomorphs recovered from ODP Site 1172 (East Tasman Plateau) is characterized by three major climate transitions.

The early late Eocene sporomorph record suggests a distinct  $2\text{--}3^\circ\text{C}$  terrestrial cooling at  $37.30$  Ma coupled with a transition from a warm-temperate *Nothofagus*–Podocarpaceae-dominated rainforest with paratropical elements to a cool-temperate *Nothofagus*-dominated rainforest with secondary Podocarpaceae. This terrestrial cooling at  $37.30$  Ma and sustained cool climate from  $37.2\text{--}35.60$  Ma coincides with a long-term SST decline from  $\sim 23$  to  $19^\circ\text{C}$  at ODP Site 1172, a regional transient cooling event (PrOM) at ODP Site 738 (Kerguelen Plateau; Scher et al., 2014), and a relatively long-term regional Southern Ocean cooling due to carbon being sequestered from surface water (Villa et al., 2008, 2014).

The expansion and restriction of cool- and warm-temperate forests from  $35.5$  to  $34.49$  Ma were followed by a period of cooling across the EOT ( $34.30\text{--}33.82$  Ma). The terrestrial climate fluctuation in this zone is consistent with latest-Eocene geochemical, sedimentological, and palynological studies that report an increase in SST, a recovery in MBT'5me-based MAATsoil (biomarker thermometry), widespread deposition of glauconite, and the common occurrence of low-latitude cosmopolitan and protoperidinioid dinocysts. These are interpreted as being linked to the initial deepening of the Tasmanian Gateway, paving the way for



**Figure 5.** Comparison of the sporomorph-based MAT in the Tasmanian Gateway region across the EOT and earliest Oligocene to regional and global marine EOT and earliest Oligocene records. **(a)** Marine benthic foraminiferal calcite  $\delta^{18}\text{O}$  record from ODP Site 1218 (Pälike et al., 2006). **(b)** Marine  $\delta^{11}\text{B}$ -derived atmospheric  $p\text{CO}_2$  record (Anagnostou et al., 2016). **(c)** Terrestrial temperature change across the EOT and earliest Oligocene based on our sporomorph-based MATs from ODP Site 1172.

the warm water associated with the PLC to affect both the terrestrial and the marine climate in this region.

The post-EOT (earliest Oligocene) recovery was characterized by a warm-temperate forest association from 33.55 to 33.06 Ma. This earliest Oligocene recovery in Tasmanian terrestrial temperatures following prior cooling across the EOT coincides with a rebound of atmospheric  $p\text{CO}_2$  at the earliest Oligocene glacial maximum (EOGM; Pearson et al., 2009) coupled with ice sheet expansion in Antarctica (Galeotti et al., 2016) and sustained deepening of the Tasmanian Gateway (Stickley et al., 2004).

Our study shows that, against a backdrop of global cooling in the late Eocene (a sustained decline in  $p\text{CO}_2$ ), a series of regional events in the marine realm, including a change in the stratification of water masses, sequestration of carbon from surface water, and changes in the ocean circulation due to Tasmanian-Gateway-accelerated deepening, may have had a knock-on effect in driving terrestrial climate and vegetation change in the Tasmanian Gateway region.

**Data availability.** All data are available for download from the Zenodo data repository at <https://doi.org/10.5281/zenodo.5924930> (Amoo et al., 2021).

**Author contributions.** MA and US conceived, designed, and led this study. PKB supplied the palynological samples for this study

and provided the biomarker thermometry data. MA and US undertook palynological analyses. MA interpreted palynological data and performed sporomorph-based bioclimatic analyses. MJP and NT provided guidance and expertise with pollen-based palaeoenvironmental reconstruction. MA prepared the manuscript with contributions from US, PKB, MJP, and NT.

**Competing interests.** The contact author has declared that neither they nor their co-authors have any competing interests.

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