# Expansion and diversification of high-latitude radiolarian assemblages in the late Eocene linked to a cooling event

# 3 in the Southwest Pacific

4

# K. M. Pascher<sup>1,2</sup>, C. J. Hollis<sup>1</sup>, S. M. Bohaty<sup>3</sup>, G. Cortese<sup>1</sup>, R. M. McKay<sup>2</sup>, H. Seebeck<sup>1</sup>, N. Suzuki<sup>4</sup> and K. Chiba<sup>4</sup>

- 7 [1] GNS Science, P O Box 30368, Lower Hutt 5040, New Zealand
- 8 [2] Victoria University Wellington, Antarctic Research Centre, P O Box 600, Wellington
- 9 6140, New Zealand
- 10 [3] Ocean and Earth Science, University of Southampton, National Oceanography Centre
- 11 Southampton, European Way, Southampton SO14 3ZH, United Kingdom
- 12 [4] Institute of Geology and Paleontology, Graduate School of Science, Tohoku University,
- 13 Sendai City, 980-8578, Japan
- 14 Correspondence to: K. M. Pascher (k.pascher@gns.cri.nz)
- 15

## 16 Abstract

The long-term cooling trend from middle to late Eocene was punctuated by several large-17 18 scale climate perturbations that culminated in a shift to "icehouse" climates at the Eocene-19 Oligocene transition. We present radiolarian microfossil assemblage and foraminiferal 20 oxygen and carbon stable isotope data from Deep Sea Drilling Project (DSDP) sites 277, 280, 21 281, 283 and Ocean Drilling Project (ODP) Site 1172 to identify significant oceanographic changes in the Southwest Pacific through this climate transition (~40-30 Ma). We find that 22 23 the Middle Eocene Climatic Optimum at ~40 Ma, which is truncated but identified by a negative shift in foraminiferal  $\delta^{18}$ O values at Site 277, is associated with a small increase in 24 25 radiolarian taxa with low-latitude affinities (5% of total fauna). In the early late Eocene at 26 ~37 Ma, a positive oxygen isotope shift at Site 277 is correlated to the Priabonian Oxygen Isotope Maximum (PrOM). Radiolarian abundance, diversity, and preservation increase 27 within this cooling event at Site 277 at the same time as diatom abundance. A negative  $\delta^{18}$ O 28 29 excursion above the PrOM is correlated with a late Eocene warming event (~36.4 Ma).

1 Radiolarian abundance and diversity decline within this event and taxa with low-latitude 2 affinities reappear. Apart from this short-lived warming event, the PrOM and latest Eocene 3 radiolarian assemblages are characterised by abundant high-latitude taxa. High-latitude taxa 4 are also abundant during the late Eocene and early Oligocene (~38-30 Ma) at DSDP sites 5 280, 281, 283 and 1172 and are associated with very high diatom abundance. We therefore infer a northward expansion of high-latitude radiolarian taxa onto the Campbell Plateau in the 6 7 latest Eocene. In the early Oligocene there is an overall decrease in radiolarian abundance 8 and diversity at Site 277, and diatoms are scarce. These data indicate that, once the Antarctic 9 Circumpolar Current was established in the early Oligocene (~30 Ma), a frontal system 10 similar to the present day developed, with nutrient-depleted Subantarctic waters bathing the 11 area around DSDP Site 277, resulting in a more restricted siliceous plankton assemblage.

# 12 **1** Introduction

13 The long-term evolution of climate through the early-to-mid Paleogene (56–34 Ma) has been 14 established from geochemical proxies and paleontological data. The primary proxy record, stable oxygen isotope ( $\delta^{18}$ O) values of benthic foraminifera, shows a trend from an early 15 16 Cenozoic greenhouse climate to an icehouse climate with an abrupt positive shift in benthic  $\delta^{18}$ O values of ~1.2–1.5‰ in the earliest Oligocene (~34 Ma) (Shackleton and Kennett, 1975; 17 Diester-Haass et al., 1996; Zachos et al., 2001). After a prolonged period of maximum 18 19 warmth during the Early Eocene Climatic Optimum centred around 53-51 Ma, long-term 20 cooling was interrupted by the Middle Eocene Climatic Optimum (MECO), a ~500 kyr 21 period of warmth peaking  $\sim 40$  Ma that has been linked to an increase in atmospheric pCO<sub>2</sub> 22 (Bohaty and Zachos, 2003; Bohaty et al., 2009; Bijl et al., 2010). Lipid biomarker-based climate proxies suggest the Southwest Pacific sea surface temperatures were tropical during 23 24 the MECO (28°C) and continued to be warm throughout the late Eocene (24–26°C), cooling 25 only slightly across the Eocene-Oligocene transition (EOT, ~22°C) (Liu et al., 2009; Bijl et 26 al., 2010).

The warm conditions of the Eocene indicated by geochemical proxies are generally consistent with fossil-based reconstructions of Southern Ocean circulation developed from high-latitude drill cores (Kennett, 1977; Nelson and Cooke, 2001; Kennett and Exon, 2004; Houben et al., 2013), in which subtropical waters are interpreted to have extended close to the Antarctic margin until the late Eocene. However, the latest generation of ocean circulation and climate

1 modelling simulations fail to reproduce the degree of high-latitude warmth indicated for the Eocene by these new proxies (Hollis et al., 2012; Lunt et al., 2012). Even under hyper-2 3 greenhouse conditions, the models produce a cyclonic gyre that blocks subtropical waters 4 from penetrating southward beyond 45°S (Huber and Sloan, 2001; Huber et al., 2004). High-5 latitude warmth also conflicts with increasing evidence for ephemeral Antarctic glaciation during the latest Eocene from both fossil and geochemical proxies (Lazarus and Caulet, 1993; 6 Scher et al., 2014; Barron et al., 2015). Following the MECO event, benthic  $\delta^{18}$ O values 7 8 increased to their maximum Eocene values of ~2.3‰ at about 37.3 Ma during a short-lived 9 cooling episode in the early late Eocene, referred to as the Priabonian Oxygen Isotope 10 Maximum (PrOM) (Scher et al., 2014). Further climate oscillations are reported for the late Eocene (Vonhof et al., 2000; Pälike et al., 2001; Bohaty and Zachos, 2003; Villa et al., 2008; 11 12 Westerhold et al., 2014) prior to the expansion of Antarctic ice that defines the EOT. A 13 negative  $\delta^{18}$ O excursion reported at ODP sites 689 (Maud Rise), 738, 744, and 748 14 (Kerguelen Plateau) (Diester-Haass and Zahn, 1996; Bohaty and Zachos, 2003; Villa et al., 15 2008; Villa et al., 2014) has been interpreted to be a short-lived warming event in the late Eocene (~36.4 Ma). 16

17 Identifying the initial timing and establishment of a high-latitude fauna in the Southern Ocean helps to constrain the development of the Southern Ocean frontal systems and, in turn, heat 18 19 transfer between low and high latitudes. Kennett (1978) provided the first summary on the 20 biogeographic development of planktic biota in the circum-polar Southern Ocean throughout 21 the Cenozoic. He inferred that the development of distinct polar plankton assemblages was 22 related to the evolution of the Antarctic Circumpolar Current (ACC) and the Antarctic Polar 23 Front (AAPF). This change was linked by Kennett (1978) to Southern Ocean circulation 24 changes associated with the opening of Drake Passage and Tasmanian Gateway in the late 25 Eocene-early Oligocene and implicated as the main causal mechanisms for Antarctic 26 glaciation. Subsequent deep-sea drilling campaigns have provided additional data on regional 27 changes in Southern Ocean plankton, which were integrated by Lazarus and Caulet (1993) into a set of circum-polar maps across specific time intervals. Moreover, these authors also 28 29 carried out the first synthesis of radiolarian biogeography for the region and found a pattern 30 of increasing endemism in the Southern Ocean across the EOT. Nelson and Cooke (2001) 31 undertook a comprehensive review of previous work and presented an updated synthesis on 32 the oceanic front development in the Southwest Pacific during the Cenozoic. According to

1 these authors, a proto-Subtropical Front was established in the late Eocene (ca. 35 Ma) and an 2 AAPF in the early Oligocene. A more detailed study of radiolarian biogeographic patterns 3 and trends in the Southwest Pacific was done by Lazarus et al. (2008), who found increased 4 endemism in the radiolarian fauna in the late Eocene (ca. 35 Ma). Further radiolarian studies 5 from the Atlantic sector of the Southern Ocean were performed by Funakawa and Nishi (2008), who recorded the first expansion of an Antarctic assemblage significantly earlier 6 7 (38.5 Ma). They identified several faunal turnover events in the Antarctic assemblage from 8 the late middle Eocene to late Oligocene and linked these events to migrations of the AAPF. 9 Latest research suggests that the ACC was not developed until ~30 Ma, together with the 10 establishment of an AAPF (Scher et al. 2015), when the Tasmanian gateway aligned with the 11 westerly wind flow (Hill et al., 2013). From the middle to late Eocene, a westward Antarctic 12 Slope Current is inferred to have flowed across the gateway, driven by the polar easterlies 13 (Bijl et al. 2013; Scher et al. 2015).

14 In this paper, we document variation in radiolarian assemblages and foraminiferal oxygen 15 and carbon stable isotopes from middle Eocene to early Oligocene (~40 to 30 Ma) at DSDP 16 Site 277 and relate these variations to radiolarian assemblage changes at DSDP sites 280, 17 281, 283 and ODP Site 1172. DSDP Site 277 provides a unique record of pelagic sedimentation in the Southwest Pacific during late Paleocene to Oligocene times and the first 18 Eocene foraminiferal  $\delta^{18}$ O record was generated from this site (Shackleton and Kennett, 19 20 1975). Although Lazarus et al.'s (2008) study of radiolarian assemblages included all above 21 mentioned DSDP sites, this new work includes a more thorough taxonomic review of the 22 radiolarian assemblages at these sites and integrates the radiolarian assemblage trends with 23 new stable isotope data for Site 277. Our results help to identify the extent to which tropical 24 or warm-subtropical conditions prevailed during the middle and late Eocene, refine the timing and nature of the development of a distinctive Southern Ocean radiolarian fauna and 25 26 discuss implications for the paleoceanography of the Southwest Pacific from the middle 27 Eocene to early Oligocene.

# 28 2 Study sites

Deep Sea Drilling Project (DSDP) sites 277, 280, 281 and 283 were drilled during DSDP Leg
29 (Kennett et al., 1975) (Figure 1). The main focus of our study is Site 277, which is located
on the western margin of the Campbell Plateau (52°13.43'S; 166°11.48'E) at a water depth of

1 1214 m. Forty-six cores were drilled with a maximum penetration of 472.5 meters below sea 2 floor (mbsf), but with total length of 434.5 m of cored section and only 59.6% recovery. Poor 3 recovery was due to 9.5 m coring runs being conducted every 19 m (i.e. alternate drilling and 4 coring at 9.5 intervals) between 301.5 and 368.0 mbsf. Below 10 mbsf, a Paleogene sequence spanning from the late Paleocene to middle Oligocene was recovered (Kennett et al., 1975). 5 We studied Cores 277-35R (349.2 mbsf) to 277-15R (134.5 mbsf) that cover a middle 6 7 Eocene-to-lower Oligocene interval. The sediment at Site 277 (paleolatitude ~55°S) 8 throughout the succession is highly calcareous indicating a depositional environment well 9 above the lysocline, with a paleodepth estimated at around 1500 m (Kennett et al., 1975; 10 Hollis et al., 1997).

11 Four additional sites were included in our study in order to acquire a regional picture of 12 radiolarian assemblage change and biogeography during the middle Eocene to early Oligocene (Figure 1). DSDP Site 280 comprises two holes (48°57.44'S; 147°14.08'E) 13 14 located ~100 km south of the South Tasman Rise and drilled at a water depth of 4176 m. We 15 collected radiolarian assemblage data from Hole 280A, which consists of a 201 m cored 16 section that includes a 97.2 m middle Eocene-to-middle Oligocene interval. The studied 17 interval spans Core 280A-7R (123.4 mbsf) to Core 280A-5R (92.54 mbsf). DSDP Site 281 on the South Tasman Rise (47°59.84'S; 147°45.85'E), drilled at a water depth of 1591 m, 18 19 encompasses two holes (281 and 281A). We examined Hole 281 which was cored to 169 20 mbsf and recovered a 105.6 m (62.5% recovery) upper Eocene-to-Pleistocene section. The 21 studied interval covers Core 281-16R (149 mbsf) to Core 281-14R (122.5 mbsf). DSDP Site 22 283 lies in the Central Tasman Sea (43°54.6'S; 154°16.96'E) at a water depth of 4729 m and 23 also comprises two holes (283 and 283A). We examined Hole 283 which was drilled to 156 24 mbsf (39% recovery) and recovered a Paleocene-to-Pleistocene section that contains an upper Eocene-to-possible Miocene hiatus. Samples from Core 283-8R (192.25 mbsf) to Core 283-25 26 5R (87.75 mbsf) were studied from this site. ODP Site 1172 is situated west of the East 27 Tasman Plateau (43°57.58'S; 149°55.69'E) in a water depth of 2622 m and was drilled during ODP Leg 189 (Exon et al., 2004). It comprises four holes (1172A, 1172B, 1172C and 28 29 1172D). The examined samples were from Section 1172A-39X-1 to Section 48X-CC 30 (354.625–450.55 mbsf), spanning a middle Eocene-to-lower Oligocene interval, and from 31 Section 1172D-2R-2 to Section 1172D-2R-3 (355.225-356.875), covering a lower Oligocene 32 interval.

# **3 Material and methods**

#### 2 **3.1 Sample preparation and analysis**

3 This study is based on 33 sediment samples from DSDP Site 277 (~350 to 135 mbsf) 4 spanning a middle Eocene-to-lower Oligocene interval (17 reported by Hollis et al. (1997) 5 and 16 new samples), 6 samples from DSDP Site 283 (new, all from the DSDP/ODP Micropaleontology Reference Centre (MRC)), 7 from DSDP Site 281 (3 from the 6 7 DSDP/ODP MRC, 4 new) and 4 from DSDP Site 280 (new). Due to incomplete core 8 recovery in all study sections, the sampling resolution of our study is variable ( $\sim 0.5$  to  $\sim 30$  m 9 sample spacing). To obtain a consistent taxonomic identification across all sites, all samples 10 previously reported from DSDP sites 277, 280, 281 and 283 were re-examined and re-11 counted as part of this study. The Supplementary files include taxonomic notes for all 12 radiolarian species recorded in this study, plates of selected species, and radiolarian 13 distribution charts and sample information for DSDP sites 277, 280, 281 and 283 14 (Supplementary Tables 1–5). Radiolarian census data of 41 samples from ODP Site 1172, covering a middle Eocene-to-lower Oligocene interval, are provided in the Supplementary 15 16 Table 6. The radiolarian taxonomy, sample preparation and analysis methodology were 17 published in Suzuki et al. (2009).

18 For strewn slide preparation, 1–10 g of sample material was broken into ~5 mm-diameter 19 chips and acidified with 10% HCl to dissolve carbonate. Samples were then washed through 20 a 63- $\mu$ m sieve, and the >63- $\mu$ m residue was cleaned by gently heating in a 1:1 solution of 21 10% hydrogen peroxide and sodium hexametaphosphate ((NaPO<sub>3</sub>)<sub>6</sub>). The residue was washed 22 though a 63-µm sieve and dried. Dependent on the volume of the processed residue and the 23 abundance of radiolarians, 1-5 strewn slides were prepared for each sample. If the 24 radiolarians were sparse, specimens were individually picked from the dried residue under a 25 stereo microscope. For strewn slides, a known portion of dried residue was evenly distributed 26 on a pre-glued coverslip, which was inverted and placed gently on a glass slide with a thin 27 coating of Canada Balsam. The slide was placed on a hot plate until the balsam was fixed.

Strewn slides were examined using a Zeiss transmitted light microscope fitted with a Zeiss AxioCam ERc5s digital camera. Radiolarian census data were derived along vertical slide traverses under transmitted light following the method of Hollis (2006). For samples with sparse radiolarians (<300 specimens per slide), all radiolarians on the prepared slide(s) were 1 counted. For richer samples, all specimens were counted until a total number of ~300 2 specimens was achieved. The proportion of the slide examined to this point was determined 3 and the abundance of common taxa (>15 observed specimens) estimated for the rest of the 4 slide. The remaining portion was then examined and rare taxa (<15 specimen observed in 5 initial count) recorded. All intact tests were assigned to a counting group that range from 6 undifferentiated order (e.g. Nassellaria undet.) and family (e.g. Actinommidae undet.) to 7 species and subspecies. This approach allows for an accurate estimate of the abundance of 8 individual species, but does result in overall diversity being underestimated.

9 Radiolarian abundance was calculated using the following equation:

10 
$$(X_R \times X_S \times \frac{1}{X_P}) \div A_{Sed}$$
 (1)

11 With  $X_R$  being the total number of radiolarians per slide,  $X_S$  the number of slides made of a 12 known portion  $X_P$  of the dried material,  $A_{Sed}$  is the initial amount of dried sediment.

13 Additional data derived for each sample assemblages includes taxic richness, the Fisher a 14 Diversity index and the Simpson index of Evenness. The latter two indices were calculated 15 using the PAST software version 3.07 (Hammer et al., 2001). The Fisher  $\alpha$  index is a general 16 guide to diversity, calculated from the number of taxa and the total number of individuals. 17 The Simpson index of Evenness determines the degree to which assemblages are dominated 18 by individual taxa and ranges from 0 to 1. Since taxic richness is correlated to preservation 19 and is also dependent on the sample size, we performed an individual rarefaction analysis for Site 277 samples with PAST (Supplementary Table 2). This allows the comparison of 20 21 taxonomic diversity in samples of different sizes. We used 100, 200, 300 and 500 counts as 22 sample sizes, respectively, to calculate taxic richness. Additionally, we derived a range-23 through taxic richness after subsampling for Site 277 with R version 3.1.3 (www.r-24 project.org) (Supplementary Table 2). We chose sample sizes of 100 and 300, respectively, 25 both with a subsampling of 1000. This approach shows if a diversity drop in the middle of a 26 series is a true diversity drop or a temporary absence due to preservation. The 27 diatom/radiolarian (D/R) ratio was calculated using the counts of diatoms and radiolarians of 28 one examined slide. In case of very rare diatoms, all specimens were counted on a slide, 29 otherwise several transverses were counted for diatoms and the total number estimated for the 30 whole slide. Although this method is not an accurate measure of total diatom abundance as 31 most pelagic diatoms are smaller than the 63-µm screen used in this study, it serves to

identify the order of magnitude in changes in diatom abundance that allows us to identify
 significant diatom event horizons.

#### 3 **3.2** Radiolarian biogeographic affinities

4 The assignment of biogeographic affinities to radiolarian species, subspecies and informally 5 defined morphotypes encountered in our study is based on a comprehensive literature review. 6 We focussed on published records of these taxa or their close relatives from the Southwest 7 Pacific and Southern Ocean (e.g. Petrushevskaya, 1975; Takemura and Ling, 1997; 8 Sanfilippo and Caulet, 1998; Hollis, 2002; Funakawa and Nishi, 2005; Hollis et al., 2005; 9 Hollis, 2006; Funakawa et al., 2006; Funakawa and Nishi, 2008; Kamikuri et al., 2012). This 10 literature review was complemented with radiolarian occurrence data from the NSB (Neptune Sandbox Berlin) Database (Lazarus, 1994; Spencer-Cervato, 1999). Unfortunately, this 11 12 database lacks many Paleogene radiolarians, and, for those that are present, occurrences need 13 to be cross-checked with the DSDP/ODP reports. The first step was to assess the 14 paleolatitude of each site for the interval of radiolarian ranges. We used 15 www.paleolatitude.org (van Hindsbergen et al., 2015) to extract paleolatitude information in 16 intervals of 10 Ma for the past 60 Ma and created the mean value for each site for an age 17 range (Supplementary Table 11). We listed radiolarian taxa and their range and abundance at 18 high-latitude (>45°N/S), mid-latitude (25–45°N/S) and low-latitude sites (0–25°N/S) and 19 observed that presence/absence data are not always a reliable guide to biogeographic affinity 20 (Supplementary Table 12). For instance, Lithomelissa ehrenbergi (Buetschli 1882) was 21 described from Barbados, which may indicate that this species has a tropical or cosmopolitan 22 ecology. However, the species is far more abundant at high-latitude sites, and only rarely 23 recorded at low-latitude sites. Moreover, Haeckel (1887) found recent L. ehrenbergi from 24 deep-water samples at low latitudes. Therefore, we interpret this species as a cold-water 25 indicator, commonly found in high-latitude samples and sometimes found in deep-water 26 samples in low latitudes. The biogeographic affinities of Amphicraspedum murrayanum and 27 A. prolixum group also warrant some discussion. These taxa are widely reported in early and 28 middle Eocene sediments but occur in greater abundance in the Southwest Pacific at times of 29 global warmth (Hollis, 2006). Liu et al. (2011) suggested that these taxa were not valid 30 indicators of high-latitude warming because they are found in the Paleocene in the North 31 Atlantic. However, their assumption that Southwest Pacific and North Atlantic Ocean 32 conditions would have been similar in the Paleogene is not supported by ocean circulation

1 models (Huber et al., 2003, 2004). These models indicate that oceanic conditions for the 2 North Atlantic and the Southwest Pacific were substantially different in the early Paleogene: 3 the North Atlantic was bathed in warm currents of  $\sim 25^{\circ}$ C moving northwards (Huber et al., 4 2003), while the Southwest Pacific was influenced by a strong cyclonic gyre preventing 5 warm waters from penetrating southwards, except during times of extreme global warmth (Huber et al., 2004; Hollis et al. 2012). Thus, the occurrence of warm-water indicators 6 7 throughout the Paleocene-Eocene interval in the mid-latitude North Atlantic is consistent 8 with both the global circulation model results and our interpretation of influxes of 9 Amphicraspedum as being indicative of warming.

10 Tectonic reconstructions of the Australia-Antarctica-Pacific plate circuit were undertaken in 11 GPlates version 1.5 (Boyden et al. 2011) using finite poles of rotation for the relative motions 12 between: Australia-East Antarctica from Cande and Stock (2004) (0-38.13 Ma); East Antarctica-West Antarctica from Granot et al. (2013) (30.94-40.13 Ma); and West 13 Antarctica-Pacific from Croon et al. (2008) (0-47.54 Ma). Relative motions of the Australia-14 15 Antarctica-Pacific plate circuit were tied to the Australian paleomagnetic apparent polar 16 wander path of Torsvik et al. (2012) to provide an estimate of paleolatitude appropriate for 17 paleoclimate studies (van Hinsbergen et al., 2015). The 2000 m isobath from the GEBCO 18 bathymetric grid (www.gebco.net) was used to approximate continental boundaries. The 19 continental/oceanic boundaries of Bird (2003) are also shown (dashed lines in Figure 1 and 8) 20 for regions where extension has significantly thinned continental crust. Each DSDP and ODP 21 study site was assigned to the appropriate plate for reconstruction.

22 The overlap of the North and South Islands of New Zealand in these reconstructions is a 23 consequence of the finite poles of rotation determined from the Adare Trough by Granot et al. 24 (2013), which constrain the motion of East and West Antarctica between 40 and 30 Ma. 25 These new poles result in a poor fit (significant overlap) of continental crust between the two 26 islands that is not supported by geological data. The discrepancy between geological and 27 paleomagnetic data could be reconciled with the use of seafloor spreading data from the 28 Emerald basin (e.g. Keller, 2003), which describes Australia-Pacific relative motions 29 (Sutherland, 1995) between 40 to 30 Ma, and the Adare Trough. However, our sites lie south of New Zealand and so we make no attempt to resolve this issue here. 30

#### **3.3 Stable isotope analysis**

Stable oxygen ( $\delta^{18}$ O) and carbon ( $\delta^{13}$ C) isotope measurements of foraminiferal samples from 2 Site 277 were conducted in the stable isotope laboratories at the University of Southampton 3 4 (UoS) and University of California Santa Cruz (UCSC). Sample analyses included bulk 5 carbonate, benthic foraminifera (Cibicidoides spp.), and the planktic foraminifera Subbotina spp. (thermocline) from 332.62–159.88 mbsf and *Globigerinatheka index* (mixed layer) from 6 7 332.62–188.58 mbsf (its last occurrence). In total, 169 samples spanning the middle Eocene-8 to-lower Oligocene interval of DSDP Hole 277 were measured (Supplementary Tables 7–10). 9 Stable isotope analyses at the UoS were performed on a Europa GEO 20-20 dual-inlet mass 10 spectrometer with CAPS preparation oven maintained at 70°C and analyses at UCSC were 11 performed on a VG Prism dual-inlet mass spectrometer coupled to carousel preparation 12 device with common acid bath maintained at 90°C. All values are reported relative to the 13 Vienna Pee Dee Belemnite (VPDB) standard. In both labs, analytical precision, based on replicate analyses of in-house marble standards and NBS-19 averaged ~0.07‰ (1 $\sigma$ ) for  $\delta^{13}$ C 14 and ~0.08‰ (1 $\sigma$ ) for  $\delta^{18}$ O. 15

# 16 **4 Results**

#### 17 **4.1 Site 277 biostratigraphy and stable isotope stratigraphy**

18 Broad age control for DSDP Site 277 is based on the biostratigraphic synthesis of Hollis et al. 19 (1997) who correlated the succession to Southern Hemisphere (SH) radiolarian Zones RP6 to 20 RP15. In this study we confirm the location of the base of RP12(SH) (Lowest Occurrence 21 [LO] of Lophocyrtis longiventer) at 371.2-349.2 mbsf, the base of RP14(SH) (LO of 22 Eucyrtidium spinosum, 38 Ma) at 264.5-254.5 mbsf, the base of RP15(SH) (LO of 23 Eucyrtidium antiquum) at 197.8–186.5 mbsf, and the base of upper Zone RP15(SH) at 143.9– 24 134.5 mbsf (Lowest Common Occurrence [LCO] of Axoprunum? irregularis). We revise the 25 base of Zone RP13(SH) to 313.5–312.7 mbsf (LO of Zealithapium mitra) (Figure 2). Further 26 refinement of the age control for Site 277 is possible through application of several additional 27 bioevents, which help to correlate the discontinuous stable isotope record of this site to those 28 from other Southern Ocean sites (Figure 2). The base of the local New Zealand stage Kaiatan 29 is defined by the Highest Occurrence [HO] of Acarinina primitiva (Morgans 2009) occurring 30 at 280-273 mbsf based on Jenkins (1975) (39.1 Ma; Raine et al., 2015). We set the base of

1 the Kaiatan at 276.5 mbsf to allow for the correlation between isotope records (Figure 2). The 2 base of the local Whaingaroan Stage (latest Eocene, 34.6 Ma; Raine et al., 2015) is identified 3 by the HO of *Globigerinatheka index*; this event was identified at 189.6 mbsf by Jenkins 4 (1975) but in the course of preparing foraminifera for stable isotope analysis we have 5 determined that the event occurs slightly higher at 188.58-187.5 mbsf. The base of 6 nannofossil zone NP17 (HO of Chiasmolithus solitus, 40.4 Ma; Gradstein et al., 2012) is 7 placed at 312.5–301.5 mbsf (Edwards and Perch-Nielsen, 1975). The LCO of Chiasmolithus 8 oamaruensis, 37.32 Ma (Gradstein et al., 2012), defines the base of NP18 at 244.5-240.6 9 mbsf (Edwards and Perch-Nielsen, 1975). The base of NP19-20 is defined by the LO of 10 Isthmolithus recurvus, 36.97 Ma (Gradstein et al., 2012) at 226.58-225.5 mbsf (Edwards and 11 Perch-Nielsen, 1975). Within NP19-20, the HO of Cribrocentrum reticulatum is found at 12 206.5-201.1 mbsf (Edwards and Perch-Nielsen, 1975), estimated at 36.44 Ma (Raine et al., 13 2015). The base of NP21-22 (HO of Discoaster saipanensis) is placed at 191.6–190.1 mbsf 14 (Edwards and Perch-Nielsen, 1975) and is dated at 34.44 Ma (Gradstein et al., 2012). As D. 15 saipanensis is a warm-water taxon, its disappearance is likely to have occurred earlier at high 16 latitudes. The Eocene-Oligocene boundary is approximated by the HO of G. index at DSDP 17 Site 277. More precise location is complicated by incomplete recovery and the highly 18 disturbed nature of Cores 277-19R, 20R, and 21R.

19 Although the recovery gaps in the Site 277 stable isotope record preclude detailed correlation, 20 the broad trends and major events such as the MECO (~40 Ma) and PrOM event (~37.3 Ma) can be identified in the benthic  $\delta^{18}$ O and  $\delta^{13}$ C isotope profiles and compared to the middle 21 Eccene-to-early Oligocene benthic isotope stratigraphy from ODP Site 689 (Maud Rise; 22 23 Diester-Haass and Zahn, 1996) (Figure 2). The EOT interval is characterized by a large 24 (~1‰) positive shift in benthic oxygen and carbon isotopes between Cores 277-20R and 19R 25 (183.64–171.28 mbsf) (Shackleton and Kennett, 1975; Keigwin, 1980), which is slightly lower than the full magnitude of the benthic  $\delta^{18}$ O shift seen at other Southern Ocean sites on 26 27 the Kerguelen Plateau and Maud Rise (Diester-Haass and Zahn, 1996; Zachos et al., 1996; 28 Bohaty et al., 2012).

#### **4.2 Site 277 oxygen and carbon isotopes**

30 Site 277  $\delta^{18}$ O results show a typical surface-to-deep gradient with more negative values in 31 bulk and planktic foraminifers compared to benthic foraminifers (Figure 3, Supplementary 1 Tables 7–10). Foraminiferal  $\delta^{13}$ C values also display typical gradients, with more positive 2 values in bulk and planktic foraminifers compared to benthic foraminifers (Figure 3). 3 However, all planktic foraminifera analysed from Site 277 are characterized by a 'frosty' 4 preservation state, indicating some diagenetic alteration (Sexton et al., 2006). We have 5 therefore focused our interpretation on benthic foraminifera because their isotopic signatures 6 are likely less affected by diagenesis.

7 Several short-lived climatic events are identified in the benthic stable isotope records at Site 8 277 (Figures 2 and 3, Supplementary Table 7). The body of the MECO was not recovered 9 due to a 16-m sampling gap between the top of Core 277-33R and the base of Core 277-32R, 10 but MECO onset and recovery is well constrained by a 0.5% negative shift in benthic  $\delta^{18}$ O values at ~313 mbsf (between Samples 277-33R-2, 106–108 cm and -33R-1, 129–130.5 cm) 11 12 and a ~0.4‰ positive shift in  $\delta^{18}$ O values at ~296 mbsf (between samples 32R-3, 107–109) cm and 32R-3, 77–79 cm), indicating that the MECO spans ~17 m (Figure 2). The MECO is 13 more strongly expressed in the benthic  $\delta^{18}$ O but this may relate to the poor recovery of the 14 body of the event at this site or diagenetic impacts on planktic  $\delta^{18}$ O values (Pearson et al., 15 2001; Sexton et al., 2006). In agreement with other records (Bohaty and Zachos, 2003; 16 Bohaty et al., 2009), a positive  $\delta^{13}C$  shift is observed in conjunction with the onset of the 17 18 MECO in the benthic and bulk carbonate records (Figure 2).

The PrOM event (Scher et al., 2014) is well-defined in the  $\delta^{18}$ O record from DSDP Site 277 19 20 but also spans three significant recovery gaps at the base of Cores 277-26R, 25R and 24R (~244.5 to 225.5 mbsf) (Figure 3). The ~0.4‰ positive shift in  $\delta^{18}$ O that marks the onset of 21 22 the PrOM, spans upper Core 277-26R and lower Core 277-25R (~240-230 mbsf), and is followed by an interval of relatively low  $\delta^{18}$ O values in upper Core 277-25R, prior to 23 24 reaching maximum values in uppermost Core 277-25R (~226 mbsf) (Figure 2). A gradual decrease in  $\delta^{18}$ O occurs through Core 277-24R. We define the PrOM at DSDP Site 277 as the 25 interval within these three cores in which benthic  $\delta^{18}$ O exceeds ~0.6‰, with the exception of 26 a narrow interval in upper Core 277-25R. These benthic  $\delta^{18}$ O values are lower than those 27 reported by Scher et al. (2014), but it is likely that peak  $\delta^{18}$ O values are not captured at Site 28 277. Consequently the PrOM is placed between 240.62 and 219.57 mbsf (spanning a ~21-m 29 section). The planktic  $\delta^{18}$ O records show similar trends to the benthic record in the PrOM 30 interval, but lacks the maximum excursion in uppermost Core 277-25R. At the onset of the 31 32 PrOM event, short-lived negative  $\delta^{13}C$  excursions are evident in the benthic, bulk and

1 planktic records. However, a longer-term positive trend for planktic and benthic  $\delta^{13}$ C values 2 is associated with the benthic  $\delta^{18}$ O maximum.

Directly above the PrOM event, there is a short-lived ~0.4‰ decrease in  $\delta^{18}$ O values in Core 277-23R (210.74 to 207.41 mbsf), evident in benthic and planktic foraminifera as well as bulk carbonate, prior to the increase in  $\delta^{18}$ O that spans the EOT (Figure 3). Benthic and planktic  $\delta^{13}$ C also exhibit a small negative excursion at this level. This interval may be correlated to the late Eocene warming interval reported from ODP sites 689 (Maud Rise), 738, 744, and 748 (Kerguelen Plateau) (Diester-Haass and Zahn, 1996; Bohaty and Zachos, 2003; Villa et al., 2008; Villa et al., 2014).

10 A large positive shift in  $\delta^{18}$ O occurs at Site 277 between the base of Core 277-20R and Core 11 277-19R, with maximum values in benthic and planktic  $\delta^{18}$ O and  $\delta^{13}$ C occurring in Core 277-

12 19R (171.28 to 169.65 mbsf). This can be correlated to the large  $\delta^{18}$ O shift across the EOT

13 documented at many deep-sea sites, which is characterised by two distinct steps (EOT-1 and

14 Oi-1) in more complete sections (e.g., Coxall et al., 2005; Katz et al., 2008).

We note that the stable isotope record at Site 277 exhibits high amplitude cyclical variation in the range of 0.5‰ for benthic  $\delta^{18}$ O and slightly more for  $\delta^{13}$ C (Figure 3). The presence of at least 10 cycles within the 6 million years between the MECO and the EOT is consistent with orbital-scale forcing. Although the record is too incomplete to establish the frequency of these cycles, their presence in this expanded Paleogene section bodes well for future drilling at this location.

#### 21 **4.3 Radiolarian assemblages at DSDP Site 277**

22 In total, 16 families, 56 genera and 98 radiolarian species were identified at DSDP Site 277 23 (Supplementary Table 1). Radiolarian abundance is generally low (10-100 specimens/g) and 24 preservation is moderate throughout the middle Eocene-to-lower upper Eocene interval 25 (349.2 to 227.2 mbsf) (Figure 4). In the uppermost Eocene and lower Oligocene (226.1-143.9 26 mbsf) radiolarians are abundant to very abundant (>1500 specimens/g) and well preserved. 27 Diversity increases during the MECO (313.5-296 mbsf) and in the upper Eocene (226.10-28 186.5 mbsf) and drops in the lower Oligocene (162.2-134.5 mbsf) (Figure 4). A short-lived 29 drop in radiolarian abundance (<500 specimens/g) and diversity is observed at 210.5–207.5 30 mbsf during the late Eocene warming event. Diversity closely parallels trends in abundance 31 and preservation. Simpson Evenness is strongly correlated with diversity but exhibits greater troughs where samples are sparse (Figure 4). Spumellarians are dominant in most samples
ranging between ~44 and 96% (~71% average). The main families are the Actinommidae,
Litheliidae, Spongodiscidae, Artostrobiidae, Lychnocaniidae and Lophocyrtiidae
(Supplementary Table 1).

5 Three samples from the middle Eocene section of Site 277 (313.5 mbsf, 312.7 mbsf, 296 6 mbsf; Cores 277-32R and 33R) that lie within the onset and recovery of the MECO, show 7 improved preservation, a peak in diversity, and mark the first significant occurrence of 8 diatoms (Figure 4). Amphicraspedum murrayanum and A. prolixum gr. have isolated 9 occurrences in this interval, while A. prolixum gr. also has trace occurrences in five samples 10 in the uppermost Eocene to lowermost Oligocene (Cores 277-24R to -20R at 217.70 mbsf, 11 209 mbsf, 207.5 mbsf, 197.82 mbsf and 186.50 mbsf). Several species are restricted to the 12 MECO, including Artobotrys titanothericeraos, Sethocyrtis chrysallis, Eusyringium fistuligerum and Stichopilium cf. bicorne. Lophocyrtis jacchia hapsis, which is a high-latitude 13 14 variant of L. jacchia jacchia (Sanfilippo and Caulet, 1998) and endemic to the Southern 15 Ocean, is also common during the MECO and uppermost Eocene (217.7–206.83 mbsf), but is 16 absent from the remaining middle and lower upper Eocene. Furthermore, the LOs of several 17 (albeit rare) species are recorded at this site during the MECO interval (Axoprunum pierinae, Zealithapium mitra, Periphaena spp., Larcopyle hayesi, L. polyacantha, Zygocircus 18 19 buetschli, Siphocampe? amygdala, Eucyrtidium montiparum, Lychnocanium amphitrite, 20 Clinorhabdus anantomus, Lophocyrtis keraspera, Lophocyrtis dumitricai, Cryptocarpium 21 ornatum and Lamprocyclas particollis) (Supplementary Table 1).

22 A major change in siliceous assemblages occurs within the PrOM interval (~226 mbsf; Core 277-25R), coincident with maximum values in benthic  $\delta^{18}$ O (Figure 4). A pronounced 23 24 increase in radiolarian abundance (from <50 to ~4000 specimens/gram), preservation and 25 diversity occurs at 226.10 mbsf (Sample 277-25R-1, 60 cm). Diatoms also become abundant 26 at the same level as the increase in radiolarian abundance. The most abundant nassellarian 27 families are the Artostrobiidae (~22%), Lophocyrtiidae (~6%) and Lychnocaniidae (~2.5%). 28 Plagiacanthidae account for ~2% of the total assemblage. The following taxa have their LO 29 within the PrOM at Site 277: Lithelius (?) foremanae, Ceratocyrtis spp., Lithomelissa ehrenbergi, L. gelasinus, L. sphaerocephalis, Siphocampe nodosaria, Artostrobus annulatus, 30 31 Artostrobus cf. pretabulatus, Clathrocyclas universa, Dictyophimus? aff. archipilium,

Lychnocanium waiareka, Aphetocyrtis rossi and Theocyrtis tuberosa (Supplementary Table
 1).

Five samples were investigated at Site 277 that lie within the late Eocene warming event (210.5–207.5 mbsf). During this event, radiolarian abundance and diversity decrease significantly, as well as diatom abundance (Figure 4). The radiolarian assemblages of these five samples differ from the other upper Eocene samples. Lychnocaniidae are more abundant (~12%), whereas Artostrobiidae are absent. Furthermore, Lophocyrtiidae decrease (~4%) and Plagiacanthidae and *Larcopyle* spp. are very rare (0.5% and 0.9%, respectively; Supplementary Table 1).

10 Immediately after the warming event, a second pronounced increase in radiolarian abundance 11 (from <200 to 9600 specimens/gram) and diversity is observed at 206.83 mbsf, together with 12 an increase in diatom abundance (Figure 4). In the uppermost Eocene-to-lowermost 13 Oligocene interval (206.83–186.5 mbsf), Plagiacanthidae (~5%), Artostrobiidae (~7%) and 14 Lophocyrtiidae (~10%) increase again, whereas Lychnocaniidae decrease (~2%; 15 Supplementary Table 1). Theocyrtis tuberosa has a very rare occurrence from the upper 16 Eocene to lower Oligocene (~226–143.9 mbsf; Core 277-25R to -16R). This species is also 17 known to have had isolated occurrences in the southern Atlantic and southern Indian oceans 18 in the late Eocene (Takemura, 1992; Takemura and Ling, 1997) and is common in latest 19 Eccene to early late Oligocene assemblages from low to middle latitudes of all ocean basins 20 (Sanfilippo et al., 1985).

A significant decline in radiolarian abundance and diversity is observed through the lower Oligocene (186.5 to 134.5 mbsf; Cores 277-20R to -15R) (Figure 4). Radiolarian abundance declines from 6400 to 750 radiolarians/gram. Many nassellarian taxa decline or disappear, especially within the Lophocyrtiidae and Plagiacanthidae. Spumellarians increase from ~73% to ~97% of the total fauna, with Litheliidae and Actinommidae being the most abundant families (Supplementary Table 1).

Rarefaction analysis of Site 277 radiolarian data (Figure 4) indicate that counts of at least 300 specimens are required to achieve a reliable measure of diversity and taxic richness. However, poor preservation in the middle Eocene and lower upper Eocene intervals (~350 to ~227 mbsf) has resulted in poor recovery of radiolarians with 9 samples containing <300 specimens and 9 samples of <100 specimens. Because these samples span an interval in which significant changes in diversity and assemblage composition occur, we include metrics</p>

1 for all samples in Figure 4 (samples of <100 specimens, <300 specimens and >300 specimens are highlighted) and metrics for samples with >100 specimens in Figures 6 and 7. To 2 3 investigate whether the diversity drop between ~292 to ~227 mbsf is a preservational artefact 4 or a real feature of the assemblage, we also determined range-through taxic richness (Figure 5 4). We have chosen sample sizes of 100 and 300 (both with a subsampling of 1000), respectively, which show a similar pattern to the original observation. The decrease in range-6 7 through taxic richness at the top and bottom of the record is due to edge effects. According to 8 this analysis, range-through taxic richness is higher than observed in Core 277-32 to -26 9 (292.2–235.5 mbsf). Chert nodules are present down-core from ~246 mbsf, so the scarcity of 10 taxa in the interval between ~350 and 246 mbsf is likely to be an artefact of diagenesis. 11 However, the increase in taxic richness in the MECO appears to be supported by this 12 analysis, at least for the uppermost sample. The analysis also indicates that there is a distinct 13 increase in diversity related to the PrOM event around ~226 mbsf, although it is more muted 14 than the raw data suggest. It is notable that the decrease in diversity evident in the raw data during the late Eocene warming event is not shown in the range-through data. In fact, there 15 may be a further increase in taxic richness within this interval. We conclude that range-16 17 through taxic richness is a helpful tool for determining if diversity changes are due to 18 diagenesis or environmental variation, especially when coupled with consideration of the 19 lithologic changes (e.g. chertification).

#### 20 **4.4 Radiolarian assemblages at other Southwest Pacific sites**

To establish the significance and nature of radiolarian faunal turnover associated with the PrOM event regionally, we investigated the upper Eocene-to-lower Oligocene intervals of DSDP sites 280, 281 and 283 and ODP Site 1172.

#### 24 4.4.1 DSDP Site 280

Four samples were investigated at DSDP Site 280 from Cores 280-7R, 6R and 5R (123.4 to 92.54 mbsf). In previous work, the E-O boundary in Hole 280 was placed at the base of Core 280-6R (110.5 mbsf) (Crouch and Hollis, 1996). However, due to the presence of *Eucyrtidium antiquum* (Caulet, 1991) and *Larcopyle frakesi* (Chen, 1975), both of which have LOs in the lower Oligocene, we place the studied interval (123.4–92.54 mbsf) in lower Oligocene Zone RP15(SH) (Figure 5, Supplementary Table 3). This is in agreement with O'Connor (2000), who found upper Eocene assemblages were restricted to Cores 280-10R to
-8R (205.5 to 139 mbsf). The absence of the zonal marker *Axoprunum? irregularis* indicates
correlation with lower RP15(SH). *Eucyrtidium spinosum*, which according to Funakawa and
Nishi (2005) has its HO in the lower Oligocene, is absent in the Site 280 study interval.
However, the HO of this species is recorded within the upper Eocene interval at Site 277,
suggesting a diachronous HO between the Southwest Pacific and the South Atlantic.

7 In total, 15 families, 35 genera and 50 radiolarian species were identified at Site 280. 8 Radiolarians are abundant (1000-2500 specimens/g) and well preserved in all samples. 9 Diatoms are also very abundant (D/R ratio ~10) (Figure 5). Diversity and Evenness is stable 10 and high in all samples. Spumellarians are slightly more abundant than nassellarians (52-11 66% of the assemblage). The most abundant families are Litheliidae (20-37%), Plagiacanthidae (14–22 %), Actinommidae (4–12%), Spongodiscidae (5–9%), Eucyrtidiidae 12 13 (4-8%) and Lophocyrtiidae (3-8%) (Supplementary Table 3). Compared to DSDP Site 277, 14 this site has higher diatom abundance and better overall preservation, which may explain the 15 higher diversity. More species of the genera Lithomelissa (7) and Larcopyle (5) are present, 16 as well as a higher abundance of Lophocyrtiidae. Lychnocaniids are very rare at this site 17 (<1%) and the genus *Lychnocanium* is absent (Supplementary Table 3).

#### 18 **4.4.2 DSDP Site 281**

19 Seven samples were investigated from DSDP Site 281 in the interval between 149 and 122.5 20 mbsf (Cores 281-16R to -14R) (Figure 5). Results from three of these samples were 21 previously reported in Crouch and Hollis (1996) but have been re-examined for this study. 22 Due to the presence of Eucyrtidium spinosum and Eucyrtidium nishimurae, the latter with a 23 HO in the late Eocene at ~36.9–36.7 Ma (Funakawa and Nishi, 2005), we correlate the Site 24 281 study interval with lower Zone RP14(SH) (~Kaiatan local stage). A hiatus spanning the 25 uppermost Eocene and Oligocene is inferred from the presence of abundant glauconite in the 26 upper part of Core 281-14R as well as from common Cyrtocapsella tetrapera in Core 281-27 13R, which indicates a Miocene age (Crouch and Hollis, 1996).

In total, 14 families, 34 genera and 46 species were identified at Site 281. Radiolarians are abundant (2000–4000 specimens/g) and well preserved. Diversity is lower than at Site 280, but Evenness is still high and similar to the other sites (Figure 5). The D/R ratio is also high and comparable to Site 280, except in the upper two samples in Core 281-14R (125.5–122.5

1 mbsf). The radiolarian assemblages are dominated by spumellarians (55-93%), with 2 Litheliidae (17–42%), Spongodiscidae (12–30%) and Actinommidae (10–20%) the most 3 abundant families. The most common nassellarians belong to the Plagiacanthidae (1-15%), 4 Lophocyrtiidae (3–7%) and Eucyrtidiidae (1–7%) (Supplementary Table 4). Although sites 5 280 and 281 were relatively close to each other (Figure 1), the radiolarian assemblages are distinctly different, indicating different oceanographic conditions. Crouch and Hollis (1996) 6 7 concluded that Site 281 was shallower and closer to terrigenous influx than Site 280. The 8 depositional environment of Site 280 is interpreted as more oceanic. The greater abundance 9 of Spongodiscidae at Site 281 supports a shallower oceanic setting for this locality (Casey, 10 1993). Compared to the early upper Eocene assemblage of Site 277, where radiolarian 11 abundance and diversity is very low, with several samples containing less than ~100 12 specimens, Site 281 contains more Spongidiscidae (~20%), Plagiacanthiidae (~7%) and 13 Litheliidae (~20%), whereas the genus *Lychnocanium* is absent at Site 281.

#### 14 **4.4.3 DSDP Site 283**

15 Six samples were examined from Site 283 between 192.25 and 87.75 mbsf (Cores 283-8R to -5R) (Figure 5). The lowermost sample at 192.25 mbsf is correlated to RP13(SH) due to the 16 17 absence of *Eucyrtidium spinosum*. The uppermost five samples are of early late Eocene age 18 based on the presence of E. spinosum and nannofossil age control (Edwards and Perch-19 Nielsen, 1975). The age of the Site 281 and 283 successions are poorly defined and the PrOM 20 event cannot be located at these sites. Both sites contain *Eucyrtidium nishimurae*: at Site 283 21 in all samples, at Site 281 its HO is in 125.5–122.5 mbsf. According to Funakawa and Nishi 22 (2005) its HO is in C17n1n (~36.7 Ma; Gradstein et al., 2012). E. nishimurae is absent at Site 23 277. The deposition of siliceous ooze in the upper middle to upper Eocene and the absence 24 (or very rare) occurrence of foraminifera suggests a deep oceanic setting close or below the 25 Calcite Compensation Depth (CCD) for Site 283.

A total of 16 families, 50 genera and 81 radiolarian species were recorded at Site 283. Radiolarians are very abundant (4700–21150 radiolarians/gram), with the highest abundance in Cores 283-6R and 5R, well preserved, and diverse (59–77 taxa per sample, Fisher  $\alpha$  Index of 10–13, Evenness of 0.75–0.89). Diatoms are present in low abundance with D/R ratios <1 (Figure 5). Spumellarians account for 59–87 % of the assemblage, with the Litheliidae (23– 38%), Actinommidae (5–19%) and the Spongodiscidae (2–8%) the most abundant families.

The Trissocyclidae (2-11%), Eucyrtidiidae (2-11%), Lophocyrtiidae (3-8%) and 1 2 Plagiacanthidae (2–8%) are the most common nassellarian families (Supplementary Table 5). 3 *Theocyrtis tuberosa* is very abundant in the uppermost sample. The acme of this taxon might 4 be correlated to its rare occurrence at Site 277 in the upper Eocene. Several taxa appear 5 earlier at Site 283 than at Site 277. These include the following taxa that occur in the upper 6 middle Eocene (e.g. Axoprunum bispiculum, Amphicentria sp. 1 sensu Suzuki, Ceratocyrtis 7 spp., Lithomelissa ehrenbergi, L. cf. haeckeli, L. sphaerocephalis, L. tricornis, 8 Pseudodictyophimus gracilipes gr., Tripodiscinus clavipes, Siphocampe nodosaria, 9 Spirocyrtis joides, Aspis sp. A sensu Hollis, Clathrocyclas universa, Eurystomoskevos 10 petrushevskaae, Lychnocanium waiareka, Aphetocyrtis gnomabax) or lower upper Eocene (Spirocyrtis greeni, Eurystomoskevos cauleti, Lophocyrtis jacchia hapsis, Lamprocyclas 11 12 particollis) at Site 283.

#### 13 **4.4.4 ODP Site 1172**

14 Forty-one samples were analysed from ODP Site 1172 spanning the middle Eocene-to-lower 15 Oligocene interval, including four samples from Core 1172D-2R (356.875–355.675 mbsf) and thirty-seven from Cores 1172A-48X to -39X (445.01-354.625 mbsf). The faunal 16 17 assemblages of ODP Site 1172 were described by Suzuki et al. (2009), who did not correlate 18 them to RP Zones. Many taxa used to define Southern Hemisphere RP zones at Site 277 are 19 absent at Site 1172 or have diachronous ranges. Eucyrtidium spinosum, the marker for Zone 20 RP14(SH), has its LO at 373.75-371.21 mbsf, but Lithomelissa tricornis and 21 Pseudodictyophimus gracilipes are absent. Eucyrtidium antiquum has a single LO at 22 365.21 mbsf, but is absent in the early Oligocene. E. nishimurae is present within the middle 23 and upper Eocene. Axoprunum irregularis is very abundant in the lower Oligocene interval at 24 this site (356.875–354.625 mbsf), which we correlate to the upper RP15(SH) zone.

Spumellarians dominate the Site 1172 assemblages throughout the middle Eocene to lower Oligocene (~82%). The Litheliidae are the most abundant family comprising about 20% on average in the middle Eocene, 35% in the upper Eocene, and 25% in the lower Oligocene. Plagiacanthidae (0.5–2.5%), Eucyrtidiidae (0.5–3%), Lophocyrtiidae (1.5–8%) and Lychnocaniidae (0.5–2.7%) account for most of the nassellarians. Fisher  $\alpha$  Diversity and Simpson Evenness are very high throughout the succession, ranging between ~10–20 and 0.82–0.96, respectively. Similar to Site 277, diversity and evenness decrease in the lower
 Oligocene (Supplementary Table 6).

3 Eocene sediments at Site 1172 consist of silty claystone with abundant diatoms. This 4 sequence is overlain by a transitional unit in the uppermost Eocene consisting of glauconitic 5 siltstones, which indicate increased bottom-water current activity in the uppermost Eocene 6 (Kennett and Exon, 2004; Stickley et al., 2004). There is a sharp transition in the lowermost 7 Oligocene to a pelagic carbonate sequence consisting of nannofossil chalk (Exon et al., 8 2004). Diatoms are more abundant and of inner neritic nature in the middle Eocene until 9 ~408 mbsf (~39 Ma), where they become more oceanic and may indicate a change to a more 10 outer neritic regime. Above ~376 mbsf (~38 Ma) the diatom assemblage indicates an inner to 11 outer neritic regime (Röhl et al., 2004).

#### 12 **4.5 Trends in biogeographic affinities**

13 The radiolarian assemblages at our five sites include 92 species or species groups that can be 14 assigned to one of three biogeographic categories: high-latitude (50 taxa), cosmopolitan (38 15 taxa), and low-latitude (4 taxa) (Table 1, Supplementary Table 12). Biogeographic affinities 16 remain poorly known for the remaining 39 taxa encountered at DSDP sites 277, 280, 281 and 17 283, and for ~100 taxa at Site 1172 reported by Suzuki et al. (2009). Within the high-latitude 18 group, six taxa are bipolar (Artostrobus annulatus, Axoprunum bispiculum, Ceratocyrtis spp., 19 Cycladophora cosma cosma, Pseudodictyophimus gracilipes gr. and Spongopyle osculosa), 20 whereas 45 taxa are inferred to be endemic to the Southern Ocean. Almost all species in the 21 Litheliidae, Lophocyrtiidae and Plagiacanthidae are high latitude. The biogeographic affinity 22 of Lithelius minor gr. is considered to be cosmopolitan, but because this group is very 23 abundant in some assemblages, we separate it out in Figures 6 and 7. For Site 277, we also 24 differentiate key high-latitude taxa within the three families noted above, namely *Larcopyle* 25 spp., Lophocyrtis longiventer and Lithomelissa spp., and the actinommid Axoprunum 26 *irregularis* (Figure 6).

At Site 277, taxa with high-latitude affinities are present from the base of the study section in the middle Eocene (Figure 6). The MECO is characterized by the presence of high-latitude taxa of ~23% (*Larcopyle* spp., *Lophocyrtis jacchia hapsis*, *L. longiventer*), but also the appearance of low-latitude species *Amphicraspedum murrayanum* and *A. prolixum* gr. (up to ~10%). *Lophocyrtis jacchia hapsis* is considered to be a high-latitude variant of *L. jacchia*  *jacchia* and has a short stratigraphic range in the middle to late Eocene in the Southern Ocean
(Sanfilippo and Caulet, 1998). In our study this taxon has a common appearance only during
the MECO and in the upper Eocene (Figure 6). In the middle of the PrOM event (~225 mbsf),
diversity and high-latitude taxa increase (average of 28%) in conjunction with the appearance
of *Lithomelissa* spp. and other high-latitude Lophocyrtiidae.

6 During the late Eocene warming event, high-latitude taxa decrease to  $\sim 13\%$  at Site 277 and 7 only rare occurrences of *Lithomelissa* spp. and high-latitude *Lophocyrtis* spp. are noted 8 (Figure 6, Table 2). Late Eocene warming however coincides with the abundant occurrence 9 of the low-latitude taxon *Thyrsocyrtis pinguisicoides* (up to 20%) and the trace occurrence of 10 A. prolixum. Cosmopolitan taxa are dominated by Lychnocanium spp. but general diversity also decreases within the warming event (Supplementary Table 1). After this event, high-11 12 latitude taxa increase to up to ~50% in the uppermost Eocene and lowermost Oligocene with 13 the reappearance of all high-latitude taxa and an overall diversification (Figure 6, Table 2). 14 During the lower Oligocene, diversity declines and especially the Plagiacanthidae and 15 Lophocyrtiidae decrease in abundance. Lithelius minor gr. is dominant until ~144 mbsf. 16 Above 144 mbsf, *Lithelius minor* gr. decreases in abundance and high-latitude actinommids 17 Axoprunum bispiculum and A. irregularis make up ~97% of the high-latitude assemblage 18 (Figure 6, Supplementary Table 1).

19 At Site 1172, high-latitude taxa are present in the middle and upper Eocene, although varying 20 between ~3 and 40% of the assemblage for which biogeographic affinities have been 21 established (Figure 7). The MECO interval at Site 1172 (Core 1172D-45X; Bijl et al., 2010) 22 corresponds to a minimum in high-latitude taxa, which is part of a longer minimum in high-23 latitude taxa from 430 to 410 mbsf. The most profound increase in high-latitude taxa at Site 24 1172 occurs in the lower Oligocene (~50-80%) with an increase in abundance of A. 25 irregularis to dominant levels, similar to Site 277. None of the low-latitude taxa found at the 26 other sites are present at Site 1172.

At Site 283, high-latitude taxa are present from the middle Eocene and range between ~12 and 35%. *Lithelius minor* gr. is very abundant and varies between ~20–40% in all samples (Figure 7). We tentatively correlate the relatively high abundance in the low-latitude species *Theocyrtis tuberosa* (~9%) in the upper part of the studied section (87.75 mbsf) to the late Eocene warming event at Site 277. Sites 280 and 281 both have a higher proportion of highlatitude taxa in the lower upper Eocene to lower Oligocene than all other sites. High-latitude taxa range between ~40 and 73% in the lower upper Eocene at Site 281 and between ~50– 73% in the lower Oligocene at Site 280, respectively (Figure 7). Several taxa that are present in the lower Oligocene at Site 280 are absent at Site 277, including *Lithomelissa challengerae*, *Larcopyle frakesi*, *Lithomelissa sakaii*, and *Antarctissa* spp. The abundance of *Lithelius minor* gr. is also high at sites 280 and 281, ranging between ~20–40%.

# 6 **5 Discussion**

#### 7 5.1 Comparison with geochemical temperature proxies

8 The radiolarian assemblages documented at sites 277 and 1172 within the MECO interval 9 lack typical Tropical taxa such as Thyrsocyrtis spp. or Podocyrtis spp. (e.g. Kamikuri et al., 2012). Taxa with low-latitude affinities, such as Amphicraspedum murrayanum and A. 10 11 prolixum gr., account for only 5% of the total assemblage at Site 277 and are absent at Site 12 1172. The persistence of high-latitude taxa and the variety of cosmopolitan species at both 13 sites suggests a warm-temperate climate (15–20°C, Nelson and Cooke, 2001), in contrast to 14 geochemical proxies suggesting a tropical climate (>25°C) for the MECO at Site 1172 (Bijl 15 et al., 2010) and ~27°C for the latest Eocene at Site 277 (Liu et al., 2009). The sea surface temperature estimates were derived from organic proxies (TEX<sub>86</sub> and  $U_{37}^{K'}$ ) and may be 16 17 biased towards summer temperatures (Liu et al., 2009; Hollis et al., 2012). Although the 18 interval of peak warmth may not be preserved in the MECO at Site 277, the relatively low 19 abundance of Tropical radiolarian taxa within the PETM and early Eocene climatic optimum 20 in the Southwest Pacific has also been previously noted by Hollis (2006; Hollis et al., 2014).

#### 21 **5.2 Nature of the Antarctic assemblage**

22 High-latitude taxa existed from at least the middle Eocene at sites 277, 283 and 1172. Many 23 taxa that are present from the earliest late Eocene (~38 Ma) at sites 281 and 283 appear later 24 at Site 277, during the PrOM event (~37 Ma), coinciding with an increase in radiolarian 25 abundance, diversity and preservation. A comparison of all high-latitude groups is shown in 26 Table 2. We assigned all *Lithomelissa* spp. and *Larcopyle* spp. to the high-latitude group as 27 they are more abundant at higher-latitude sites. Although we assigned a cosmopolitan affinity 28 to *Lithelius minor* gr., the paleoecology of this group is not yet fully understood, as it tends to 29 be most abundant at high-latitude sites. The sudden appearance of Lithomelissa spp. and

other high-latitude taxa and diatoms at Site 277 indicates the expansion of high-latitude water masses across the southern Campbell Plateau in two phases: first during the PrOM event and again after the late Eocene warming event when a second diversification and influx of highlatitude taxa is observed.

#### 5 5.3 High-latitude cooling and eutrophication during the PrOM event

#### 6 5.3.1 Diagenesis

7 One possible explanation for the pronounced increase in radiolarian abundance and diversity 8 in the upper Eocene at Site 277 is that these trends are an artefact of biogenic opal diagenesis. 9 Chert nodules are recorded throughout the upper Paleocene-to-middle Eocene section of the 10 cored sequence at Site 277, with a transition between chert-bearing nannofossil chalk and 11 overlying nannofossil ooze at 246 mbsf (lower upper Eocene) (Kennett et al., 1975). The 12 presence of chert combined with the generally poorer preservation of radiolarians in the 13 lower Paleogene interval indicates some degree of diagenesis, which is also reflected in the 14 range-through taxic richness analysis. However, the first major radiolarian turnover event 15 occurs ~20 m above the lithological transition from chert-bearing nannofossil chalk to 16 nannofossil oozes, which implies that the event represents a real increase in radiolarian and 17 diatom abundance and not an artefact of diagenesis. No lithological changes are present at 18 that levels which could explain the observed diversity decrease during the late Eocene 19 warming event and the increase in diversity thereafter.

#### 20 **5.3.2 Climate cooling**

21 The long-term cooling trend through the middle and late Eocene, which was interrupted by 22 the short-lived MECO warming event, does not explain the sudden radiolarian diversification 23 in the late Eocene at Site 277. If gradual, long-term cooling was the driver of the expansion 24 of high-latitude taxa, a progressive increase in such taxa would be expected over a longer 25 time period. A gradual increase of high-latitude taxa is observed at Site 1172 from the middle 26 Eocene but not at Site 277. Instead, the short-lived PrOM event appears to have been the 27 trigger for the northward expansion of high-latitude taxa onto the Campbell Plateau. Whether 28 this event was caused by an abrupt decline in atmospheric CO<sub>2</sub> concentrations or was related 29 to the opening of the Tasmanian Gateway, which may have been open to surface circulation 30 in early middle Eocene (Bijl et al., 2013), cannot be determined. Furthermore, astronomically 1 induced changes also may have had a role. Laskar et al. (2004) calculated nodes in the 2 amplitude modulation of eccentricity and obliquity at ~37 Ma and Röhl et al. (2004) found 3 evidence at Site 1172 for the increasing dominance of the 100 kyr eccentricity cycle at  $\sim$ 37 4 Ma. Although there are nodes in amplitude modulation throughout the Eocene (Laskar et al, 5 2004), it is possible that a combination of these phenomena (e.g., a decrease in atmospheric CO<sub>2</sub>, gateway opening and nodes in amplitude modulation) caused a cooling event. The 6 7 amplitude modulation of obliquity, in particular, has been linked to climatic cooling in the 8 Oligocene (Pälike et al., 2006).

9 The PrOM event at ~37 Ma may have been associated with the formation of small Antarctic 10 ice sheets (Scher et al., 2014), which in turn may have caused an intensification of currents in the Southern Ocean. Funakawa and Nishi (2008) reported a marked increase in radiolarian 11 taxa with Antarctic faunal affinities at ODP Site 689 (Maud Rise, southern Atlantic) in the 12 earliest late Eocene (~38.6-36.9 Ma; Chron C17), which they interpreted to signify the 13 14 northward expansion of the polar front that may be related to the PrOM. A subsequent 15 decrease in Antarctic taxa, spanning Chrons 16 and 15 (~37-35 Ma), was related to the late 16 Eccene warming event of Bohaty and Zachos (2003). These cooling and warming events 17 appear to have caused longer-lived changes in radiolarian faunal assemblages than those observed at Site 277, lasting 1.7 and 2.0 Ma, respectively. This may reflect specific 18 19 differences in the oceanographic settings of the two sites. Although the late Eocene warming 20 event appears to be short-lived at Site 277, it may have spanned a longer interval of time 21 elsewhere in the Southwest Pacific. Incursions of warm-water foraminifera, including the 22 low-latitude genus Hantkenina, are known to have occurred in the middle late Eocene (late 23 Kaiatan-early Runangan) in sedimentary basins of southern New Zealand (Hornibrook et al., 24 1989; Hornibrook, 1992).

#### 25 **5.3.3 Radiolarian biogeographic reconstruction**

During the middle Eocene, high-latitude radiolarian taxa were present at sites 277, 283, and 1172 (Figure 7). The short-lived increase in abundance, diversity and the influx of lowlatitude radiolarian *Amphicraspedum murrayanum* and *A. prolixum* gr. during the MECO at Site 277 and a high percentage of cosmopolitan taxa at Site 1172 suggest moderately warm temperatures at both sites, which may have been the result of a slightly stronger influence of an East Australian Current (EAC) (Figure 8A). However, radiolarians and diatoms were abundant only at Site 1172 during the middle Eocene, which suggests a higher productivity
region, perhaps a consequence of local of upwelling. The Tasmanian Gateway was open to a
shallow westward flowing Antarctic Slope Current (ASC), driven by the polar easterlies (Bijl
et al., 2013; Scher et al. 2015).

5 During the onset of the PrOM event in the early late Eocene (~38–37 Ma, Figure 8B), the 6 abundance of high-latitude taxa increased at sites 1172 and 277. Additionally, sites 281 and 7 283 were characterized by high radiolarian abundance, with an average of ~61 and ~27% 8 high-latitude taxa, respectively. The region of high-productivity expanded in this time, with 9 the southernmost Site 281 having the highest D/R ratio in the interval (Figure 8B). The region 10 might have experienced an intensification of the Ross gyre, extending the region of high productivity onto the Campbell Plateau and creating a Subtropical Front (STF) (Nelson and 11 12 Cooke, 2001) (Figure 8B).

13 During the late Eocene warming event (~36 Ma, Figure 8C), radiolarian diversity decreased 14 abruptly at Site 277, together with a decrease in high-latitude taxa (Lithomelissa spp., 15 Larcopyle spp., Lophocyrtiidae, Table 2) and diatoms, and the appearance of low-latitude 16 taxa at sites 283 and 277. Site 281 contains a late Eocene hiatus, implying an increase in the 17 strength of bottom water currents across the Tasmanian Gateway. We suggest that these 18 changes were associated with a southward shift of the EAC that pushed the high-productivity 19 zone of the STF towards the south, explaining the low radiolarian abundance and drop in 20 diversity at Site 277. There is little evidence that the large Tasman Current as proposed by 21 Huber et al. (2004) and Bijl et al. (2010) existed in the middle and late Eocene. Instead, our 22 data suggest Site 277 was positioned at the northernmost limit of the influence of the Ross 23 Gyre and the southernmost site of the influence of warm water delivered by the EAC.

24 During the latest Eocene-earliest Oligocene interval (~35-32 Ma, Figure 8D), Site 277 experienced a second siliceous plankton bloom, associated with high radiolarian and diatom 25 26 abundance and the reappearance of high-latitude taxa (Table 2). This suggests that latest 27 Eccene cooling led to the expansion of the Ross Gyre to encroach on Campbell Plateau 28 (Figure 8D). At the same time and perhaps reflecting strengthening of northward and westward flowing bottom currents, the area of non-deposition widened across the Tasmanian 29 30 Gateway over sites 281, 283 and 1172. During the earliest Oligocene, the abundance of 31 radiolarians and diatoms at Site 280 suggests high primary productivity region. This may

have been a consequence of intensified upwelling associated with the ASC in conjunction
 with deepening in this sub-basin.

3 In contrast, diversity declines at Site 277 in the early Oligocene (Figure 8E) and diatoms become scarce. The radiolarian fauna becomes dominated by Lithelius minor gr. and 4 5 Actinommidae and many high-latitude taxa disappear (e.g. Lithomelissa spp., Table 2). This 6 may be related to the development of the ACC. The ACC is inferred to have developed at 7 ~30 Ma as the Tasmanian Gateway became fully open (Carter et al., 2004) and its northward 8 expansion brought it in line with the westerly wind belt (Scher et al., 2015). This resulted in 9 the zone of non-deposition extending over Site 280 as it moved into the path of the ACC. At 10 Site 277, the radiolarian fauna is dominated by the high-latitude species Axoprunum irregularis, which is also dominant at Site 1172. Thus, the general low diversity of 11 12 radiolarians and the scarcity of diatoms at Site 277 suggests the establishment of a cold-water 13 nutrient-depleted environment, similar to the modern setting (Hollis and Neil, 2005). The 14 development of the ACC restricted the northward extent of Ross Gyre and served to establish 15 the Subantarctic Front on the southern margin of the Campbell Plateau (Carter et al., 2004).

# 16 6 Conclusions

17 Middle Eocene-to-early Oligocene radiolarian assemblages from DSDP sites 277, 280, 281, 18 283 and ODP Site 1172 were examined to investigate the relative influence of low- and high-19 latitude water masses in the southern Southwest Pacific Ocean as global climate cooled and 20 ice sheets expanded in Antarctica. In contrast to temperature reconstructions based on 21 geochemical proxies that indicate subtropical-tropical temperatures at high-latitudes during 22 the middle and late Eocene (Liu et al., 2009; Bijl et al., 2010), Eocene radiolarian 23 assemblages in this region lack significant numbers of warm-water taxa. Furthermore, we 24 show that many high-latitude and taxa endemic to the Antarctic are already present in the middle Eocene. The MECO has been identified at Site 277 from for aminiferal  $\delta^{18}O$  records 25 26 and is associated with a short-lived incursion of two taxa with low-latitude affinities, 27 Amphicraspedum prolixum gr. and A. murrayanum. The absence of definitive Tropical taxa 28 suggests warm temperate rather than tropical conditions during this short-lived event.

Radiolarians are very abundant and well preserved at high-latitude sites 281, 283 and 1172 during the early late Eocene and at Site 280 during the early Oligocene. For taxa with identified biogeographic affinities, those with high-latitude affinities comprise ~60% at sites

1 280 and 281 and ~30% at sites 283 and 1172. During the early late Eocene (~37 Ma), a positive shift in foraminiferal  $\delta^{18}$ O values at Site 277 marks the onset of the PrOM event. A 2 3 pronounced increase in diversity, abundance and preservation of radiolarians occurs in 4 conjunction with this event at Site 277 in addition to a marked increase in diatom abundance. 5 Many high-latitude taxa that are very abundant at sites 281 and 283 in the late middle Eocene 6 and early late Eocene become abundant or have their LOs at Site 277 at ~37 Ma, including: 7 Lithelius minor gr., Larcopyle hayesi, L. polyacantha, Spongopyle osculosa, Lithomelissa 8 sphaerocephalis, L. gelasinus, L. ehrenbergi, Ceratocyrtis spp., Dictyophimus aff. 9 archipilium, Lamprocyclas particollis, and Antarctic morphotypes of Aphetocyrtis 10 gnomabax, A. rossi, Lophocyrtis aspera, L. keraspera and L. longiventer. This northward 11 extension of high-latitude taxa onto the Campbell Plateau appears to have been triggered by 12 cooling during the PrOM event, which may have been associated with a short-lived 13 development of an Antarctic ice sheet.

14 A late Eocene warming event at ~36 Ma is accompanied by a decrease in radiolarian 15 diversity, high-latitude taxa and low diatom abundance at Site 277. Two low-latitude taxa, 16 Theocyrtis tuberosa and Thyrsocyrtis pinguisicoides, make short-lived incursions into the 17 Southwest Pacific at this time. After this event, radiolarian diversity increases again with the 18 reappearance of high-latitude taxa and abundant diatoms at Site 277. Through the EOT, 19 radiolarians decrease in abundance and diversity at Site 277. Most nassellarian taxa within 20 the Plagiacanthidae and Lophocyrtiidae decline, whereas Lithelius minor gr. and 21 Actinommidae become dominant. Together with the scarcity of diatoms, we infer that 22 conditions over the Campbell Plateau became nutrient-depleted as a consequence of the 23 development of the ACC. The establishment of the ACC at around 30 Ma is inferred to have 24 caused widespread non-deposition in the Southwest Pacific and restricted the northward flow 25 of Ross Gyre.

26

#### 27 Acknowledgements

This study used bulk material and reference slides stored in the DSDP/ODP Micropaleontology Reference Centre, which is located at GNS Science, Lower Hutt, New Zealand. We greatly appreciate the reviews of David Lazarus and an anonymous referee that provided helpful comments in improving our manuscript. We acknowledge the technical support of Sonja Bermudez (GNS Science), James Crampton (GNS Science) and Johan Renaudie (Museum f
ür Naturkunde, Berlin) and editorial handling by Gerald Dickens. This
 project is funded by the New Zealand Marsden Fund (Contract GNS1201).

# 3 **References**

- Barron, J. A., Stickley, C. E., and Bukry, D.: Paleoceanographic, and paleoclimatic
  constraints on the global Eocene diatom and silicoflagellate record, Palaeogeogr.
  Palaeoclimatol. Palaeoecol., 422, 85-100, doi:10.1016/j.palaeo.2015.01.015, 2015.
- 7 Bijl, P. K., Houben, A. J., Schouten, S., Bohaty, S. M., Sluijs, A., Reichart, G.-J., Damsté, J.
- S. S., and Brinkhuis, H.: Transient Middle Eocene atmospheric CO2 and temperature
  variations, Science, 330, 819-821, 2010.
- 10 Bijl, P. K., Bendle, J. A., Bohaty, S. M., Pross, J., Schouten, S., Tauxe, L., Stickley, C. E.,
- 11 McKay, R. M., Röhl, U., and Olney, M.: Eocene cooling linked to early flow across the
- 12 Tasmanian Gateway, Proc. Natl. Acad. Sci. U.S.A., 110, 9645-9650, 2013.
- Bird, P.: An updated digital model of plate boundaries, Geochem. Geophys. Geosyst., 4(3),
  1027, doi:10.1029/2001GC000252, 2003.
- Bohaty, S. M., and Zachos, J. C.: Significant Southern Ocean warming event in the late
  middle Eocene, Geology, 31, 1017-1020, doi: 10.1130/G19800.1, 2003.
- Bohaty, S. M., Zachos, J. C., Florindo, F., and Delaney, M. L.: Coupled greenhouse warming
  and deep-sea acidification in the middle Eocene, Paleoceanography, 24, PA2207,
  doi:10.1029/2008PA001676, 2009.
- Bohaty, S. M., Zachos, J. C., and Delaney, M. L.: Foraminiferal Mg/Ca evidence for
  Southern Ocean cooling across the Eocene-Oligocene transition, Ear. Planet. Sci. Lett., 317,
  251-261, doi:10.1016/j.epsl.2011.11.037, 2012.
- 23 Boyden, J. A., Müller, R. D., Gurnis, M., Torsvik, T. H., Clark, J. A., Turner, M., Ivey-Law,
- 24 H., Watson, R. J., and Cannon, J. S.: Next-generation plate-tectonic reconstructions using
- 25 GPlates, Geoinformatics: cyberinfrastructure for the solid earth sciences, 95-114, 2011.
- 26 [http://www.gplates.org/index.html; download on 25-09-2015]
- Cande, S. C., and Stock, J. M.: Pacific—Antarctic—Australia motion and the formation of
  the Macquarie Plate, Geophys. J. Int., 157, 399-414, 2004.
  - 28

- Carter, L., Carter, R., and McCave, I.: Evolution of the sedimentary system beneath the deep
   Pacific inflow off eastern New Zealand, Marine Geology, 205, 9-27, 2004.
- Casey, R. E.: Radiolaria, in: Fossil Prokaryotes and Protists, edited by: Lipps, J. H.,
  Blackwell Scientific Publications, Oxford/London, UK, 249-284, 1993.
- 5 Caulet, J. P.: Radiolarians from the Kerguelen Plateau, Leg 119, in: Barron, J.A., Larsen, B.
- 6 et al., Proceedings ODP, Scientific Results, 119, College Station, TX (Ocean Drilling
  7 Program), 513-546, 1991.
- 8 Chen, P. H.: Antarctic Radiolaria, in: Hayes, D.E., Frakes, L.A., et al., Initial Reports of the
  9 Deep Sea Drilling Project, Vol. 28, U.S. Government Printing Office, Washington, D.C.,
  10 437-513, 1975.
- Coxall, H. K., Wilson, P. A., Pälike, H., Lear, C. H., & Backman, J.: Rapid stepwise onset of
  Antarctic glaciation and deeper calcite compensation in the Pacific Ocean. Nature,
  433(7021), 53-57, 2005.
- Croon, M. B., Cande, S. C., and Stock, J. M.: Revised Pacific-Antarctic plate motions and
  geophysics of the Menard Fracture Zone, Geochem. Geophys. Geosyst., 9, Q07001,
  doi:10.1029/2008GC002019, 2008.
- Crouch, E. M., and Hollis, C. J.: Paleogene palynomorph and radiolarian biostratigraphy of
  DSDP Leg 29, sites 280 and 281 South Tasman Rise, Institute of Geological and Nuclear
  Sciences science report 96/19, 46p., 1996.
- 20 Diester-Haass, L., and Zahn, R.: Eocene-Oligocene transition in the Southern Ocean: History
- of water mass circulation and biological productivity, Geology, 24, 163-166, 1996.
- Diester-Haass, L., Robert, C., and Chamley, H.: The Eocene-Oligocene preglacial-glacial
  transition in the Atlantic sector of the Southern Ocean (ODP Site 690), Mar. Geol., 131, 123149, 1996.
- 25 Edwards, A. R., and Perch-Nielsen, K.: Calcareous nannofossils from the southern southwest
- 26 Pacific, Deep Sea Drilling Project, Leg 29, in: Kennett, J. P., Houtz, R. E., et al., Initial
- 27 Reports of the Deep Sea Drilling Project, Vol. 29, Washington, DC, US Government Printing
- 28 Office, 469-539, 1975.
- Exon, N. F., Kennett, J. P., and Malone, M. J.: Leg 189 synthesis: Cretaceous-Holocene
  history of the Tasmanian gateway, in: Proceedings ODP, Scientific Results, 2004.

- Florindo, F., and Roberts, A. P.: Eocene-Oligocene magnetobiochronology of ODP Sites 689
   and 690, Maud Rise, Weddell Sea, Antarctica, Geol. Soc. Am. Bull., 117, 46-66, 2005.
- Funakawa, S., and Nishi, H.: Late middle Eocene to late Oligocene radiolarian
  biostratigraphy in the Southern Ocean (Maud Rise, ODP Leg 113, Site 689), Mar.
  Micropaleontol., 54, 213-247, 2005.
- Funakawa, S., and Nishi, H.: Radiolarian faunal changes during the Eocene-Oligocene
  transition in the Southern Ocean (Maud Rise, ODP Leg 113, Site 689) and its significance in
  paleoceanographic change, Micropaleontology, 54, 15-26, 2008.
- 9 Funakawa, S., Nishi, H., Moore, T. C., and Nigrini, C. A.: Radiolarian faunal turnover and
- 10 paleoceanographic change around Eocene/Oligocene boundary in the central equatorial
- 11 Pacific, ODP Leg 199, Holes 1218A, 1219A, and 1220A, Palaeogeogr. Palaeoclimatol.
- 12 Palaeoecol., 230, 183-203, 2006.
- Gradstein, F., Ogg, J., Schmitz, M., and Ogg, G.: The geologic time scale 2012, vol. 2,
  Elsevier New York, 2012.
- Granot, R., Cande, S., Stock, J., and Damaske, D.: Revised Eocene-Oligocene kinematics for
  the West Antarctic rift system, Geophys. Res. Lett., 40, 279-284, doi:
  10.1029/2012GL054181, 2013.
- Hammer, Ø., Harper, D., and Ryan, P.: Past: Paleontological Statistics Software Package for
  education and data analysis. Paleontología Electrónica 4: 1-9, http://palaeo-electronica.
  org/2001\_1/past/issue1\_01. html, 2001. [download of version 3.17 on 24-07-2015,
  http://folk.uio.no/ohammer/past/]
- Hollis, C. J.: Biostratigraphy and paleoceanographic significance of Paleocene radiolarians
  from offshore eastern New Zealand, Mar. Micropaleontol., 46, 265-316, 2002.
- Hollis, C. J.: Radiolarian faunal change across the Paleocene-Eocene boundary at Mead
  Stream, New Zealand, Eclogae Geol. Helv., 99, S79-S99, 2006.
- Hollis, C., and Neil, H.: Sedimentary record of radiolarian biogeography, offshore eastern
  New Zealand, New Zeal. J. Mar. Fresh., 39, 165-192, 2005.
- 28 Hollis, C. J., Waghorn, D. B., Strong, C. P., and Crouch, E. M.: Integrated Paleogene
- 29 Biostratigraphy of DSDP Site 277 (Leg 29): Foraminifera, Calcareous Nannofossils,
- 30 Radiolaria, and Palynomorphs, Institute of Geological & Nuclear Sciences Limited, 1997.

- 1 Hollis, C. J., Dickens, G. R., Field, B. D., Jones, C. M., and Percy Strong, C.: The Paleocene-
- 2 Eocene transition at Mead Stream, New Zealand: a southern Pacific record of early Cenozoic
- 3 global change, Palaeogeogr. Palaeoclimatol. Palaeoecol., 215, 313-343, 2005.
- 4 Hollis, C. J., Taylor, K. W. R., Handley, L., Pancost, R. D., Huber, M., Creech, J. B., Hines,
- 5 B. R., Crouch, E. M., Morgans, H. E. G., Crampton, J. S., Gibbs, S., Pearson, P. N., and
- 6 Zachos, J. C.: Early Paleogene temperature history of the Southwest Pacific Ocean:
- 7 Reconciling proxies and models, Earth Planet. Sci. Lett., 349, 53-66, DOI
- 8 10.1016/j.epsl.2012.06.024, 2012.
- 9 Hollis, C. J., Pascher, K. M., Hines, B. R., Littler, K., Kulhanek, D. K., Strong, C. P., Zachos,
- 10 J. C., Eggins, S. M. and Philips, A.: Was the Early Eocene ocean unbearably warm or are the
- 11 proxies unbelievably wrong? Rendiconti Online 31, 109-110, 2014.
- 12 Hornibrook, N. d. B.: New Zealand Cenozoic marine paleoclimates: a review based on the
- 13 distribution of some shallow water and terrestrial biota, Pacific Neogene: environment,
- 14 evolution, and events. University of Tokyo Press, Tokyo, 83-106, 1992.
- 15 Hornibrook, N. de B., Brazier, R. C., and Strong, C. P.: Manual of New Zealand Permian to
- Pleistocene foraminiferal biostratigraphy, Paleontological bulletin/New Zealand Geological
  Survey, 56, 1-175, 1989.
- 18 Houben, A. J., Bijl, P. K., Pross, J., Bohaty, S. M., Passchier, S., Stickley, C. E., Röhl, U.,
- 19 Sugisaki, S., Tauxe, L., and van de Flierdt, T.: Reorganization of Southern Ocean Plankton
- 20 Ecosystem at the Onset of Antarctic Glaciation, Science, 340, 341-344, 2013.
- Huber, M., and Sloan, L. C.: Heat transport, deep waters, and thermal gradients: Coupled
  simulation of an Eocene greenhouse climate, Geophys. Res. Lett., 28, 3481-3484, 2001.
- Huber, M., Sloan, L. C., and Shellito, C.: Early Paleogene oceans and climate: A fully
  coupled modeling approach using the NCAR CCSM, Geological Society of America Special
  Papers, 369, 25-47, 2003.
- 26 Huber, M., Brinkhuis, H., Stickley, C. E., Döös, K., Sluijs, A., Warnaar, J., Schellenberg, S.
- 27 A., and Williams, G. L.: Eocene circulation of the Southern Ocean: Was Antarctica kept
- warm by subtropical waters?, Paleoceanography, 19, PA4026, doi:10.1029/2004PA001014,
- 29 2004.

- 1 Jenkins, D. G.: Cenozoic planktic foraminiferal biostratigraphy of the southwestern Pacific
- 2 and Tasman Sea DSDP Leg 29, in: Kennett, J.P., Houtz, R.E. et al., Initial Reports of the
- 3 Deep Sea Drilling Project, Vol. 29, U.S. Government Printing Office, Washington, D.C. ,
  4 449-467, 1975.
- Kamikuri, S.-i., Moore, T. C., Lyle, M., Ogane, K., and Suzuki, N.: Early and Middle Eocene
  radiolarian assemblages in the eastern equatorial Pacific Ocean (IODP Leg 320 Site U1331):
  Faunal changes and implications for paleoceanography, Mar. Micropaleontol., 98, 1-13,
  doi:10.1016/j.marmicro.2012.09.004, 2013.
- 9 Katz, M. E., Miller, K. G., Wright, J. D., Wade, B. S., Browning, J. V., Cramer, B. S.,
- 10 Rosenthal, Y.: Stepwise transition from the Eocene greenhouse to the Oligocene icehouse.
- 11 Nature Geoscience, 1(5), 329-334, 2008.
- 12 Keigwin, L.: Palaeoceanographic change in the Pacific at the Eocene-Oligocene boundary,
- 13 Nature, 287, 722-725, 1980.
- Keller, W.R.: Cenozoic plate tectonic reconstructions and plate boundary processes in the
  Southwest Pacific. Unpub. PhD Thesis: California Institute of Technology. Pasadena, 2003.
- 16 Kennett, J. P.: Cenozoic evolution of Antarctic glaciation, the circum-Antarctic Ocean, and
- 17 their impact on global paleoceanography, J. Geophys. Res., 82, 3843-3860, 1977.
- 18 Kennett, J. P.: The development of planktonic biogeography in the Southern Ocean during
  19 the Cenozoic, Mar. Micropaleontol., 3, 301-345, 1978.
- 1) the central, mar. meropatontol., 5, 501 545, 1976.
- 20 Kennett, J. P., and Exon, N. F.: Paleoceanographic evolution of the Tasmanian Seaway and
- 21 its climatic implications, in: The Cenozoic Southern Ocean: Tectonics, Sedimentation, and
- Climate Change Between Australia and Antarctica, Geoph. Monog. Series 151, 345-367,2004.
- 24 Kennett, J. P., Houtz, R. E., Andrews, P. B., Edwards, A. R., Gostin, V. A., Hajós, M.,
- 25 Hampton, M., Jenkins, D. G., Margolis, S., Ovenshine, T., and Perch-Nielsen, K.: Initial
- 26 Reports of the Deep Sea Drilling Project, Vol. 29, U.S. Government Printing Office,
- 27 Washington, D.C., 1975.
- 28 Laskar, J., Robutel, P., Joutel, F., Gastineau, M., Correia, A., and Levrard, B.: A long-term
- numerical solution for the insolation quantities of the Earth, Astron. Astrophys., 428, 261285, 2004.

- Lazarus, D., and Caulet, J. P.: Cenozoic Southern Ocean reconstructions from
   sedimentologic, radiolarian, and other microfossil data, Antarct. Res. Ser., 60, 145-174, 1993.
- Lazarus, D.: Neptune: A marine micropaleontology database, Math. Geol., 26, 817-832,
  10.1007/BF02083119, 1994. [last access: 21-08- 2015]
- 5 Lazarus, D., Hollis, C., and Apel, M.: Patterns of opal and radiolarian change in the Antarctic
- mid-Paleogene: Clues to the origin of the Southern Ocean, Micropaleontology, 54, 41-48,
  2008.
- 8 Liu, Z., Pagani, M., Zinniker, D., DeConto, R., Huber, M., Brinkhuis, H., Shah, S. R., Leckie,
- 9 R. M., and Pearson, A.: Global cooling during the Eocene-Oligocene climate transition,
- 10 Science, 323, 1187-1190, 2009.
- 11 Liu, J., Aitchison, J. C., and Ali, J. R.: Upper Paleocene radiolarians from DSDP Sites 549
- 12 and 550, Goban Spur, NE Atlantic, Palaeoworld, 20, 218-231, 2011.
- 13 Lunt, D. J., Dunkley Jones, T., Heinemann, M., Huber, M., LeGrande, A., Winguth, A.,
- Loptson, C., Marotzke, J., Roberts, C., and Tindall, J.: A model–data comparison for a multimodel ensemble of early Eocene atmosphere–ocean simulations: EoMIP, Clim. Past, 8, 1717-
- 16 1736, 2012.
- 17 Matthews, K. J., Williams, S. E., Whittaker, J. M., Müller, R. D., Seton, M., and Clarke, G.
- 18 L.: Geologic and kinematic constraints on Late Cretaceous to mid Eocene plate boundaries in
- 19 the southwest Pacific, Earth Sci. Rev., 140, 72-107, 2015.
- Morgans, H. E. G.: Late Paleocene to Middle Eocene foraminiferal biostratigraphy of the
  Hampden Beach section, eastern South Island, New Zealand. New Zeal. J. Geol. Geop., 52,
  273-320, 2009.
- Nelson, C. S., and Cooke, P. J.: History of oceanic front development in the New Zealand
  sector of the Southern Ocean during the Cenozoic—a synthesis, New Zeal. J. Geol. Geop.,
  44, 535-553, 2001.
- O'Connor, B.: Stratigraphic and geographic distribution of Eocene Miocene Radiolaria from
  the southwest Pacific, Micropaleontology, 46, 189-228, 2000.
- 28 Pälike, H., Shackleton, N. J., and Röhl, U.: Astronomical forcing in Late Eocene marine
- 29 sediments, Ear. Planet. Sci. Lett., 193, 589-602, 2001.

- 1 Pälike, H., Frazier, J., & Zachos, J. C.: Extended orbitally forced palaeoclimatic records from
- 2 the equatorial Atlantic Ceara Rise. Quaternary Sci. Rev., 25(23), 3138-3149, 2006.
- 3 Pearson, P. N., Ditchfield, P. W., Singano, J., Harcourt-Brown, K. G., Nicholas, C. J., Olsson,
- 4 R. K., Shackleton, N. J., and Hall, M. A.: Warm tropical sea surface temperatures in the Late
- 5 Cretaceous and Eocene epochs, Nature, 413, 481-487, 2001.
- 6 Petrushevskaya, M. G.: Cenozoic radiolarians of the Antarctic, Leg 29, DSDP, in: Initial
- 7 Reports of the Deep Sea Drilling Project, edited by: Kennett, J. P., Houtz, R. E., et al., US
- 8 Government Printing Office, Washington, DC, vol. 29, 541-675, 1975.
- 9 Raine, J. I., Beu, A. G., Boyes, A. F., Campbell, H. J., Cooper, R. A., Crampton, J. S.,
- 10 Crundwell, M. P., Hollis, C. J., and Morgans, H. E. G.: Revised calibration of the New
- 11 Zealand Geological Timescale : NZGT2015/1, Lower Hutt, N.Z.: GNS Science. GNS
- 12 Science report 2012/39. 53 p, 2015.
- Röhl, U., Brinkhuis, H., Stickley, C. E., Fuller, M., Schellenberg, S. A., Wefer, G., and
  Williams, G. L.: Sea level and astronomically induced environmental changes in middle and
  late Eocene sediments from the East Tasman Plateau, in: Exon, N.F., Kennett, J.P., and
  Malone, M.J. (Eds.), The Cenozoic Southern Ocean: tectonics, sedimentation, and climate
  change between Australia and Antarctica. Am. Geophys. Union, Geophys. Monogr., 151,
  127-151, 2004.
- Sanfilippo, A., and Caulet, J. P.: Taxonomy and evolution of Paleogene Antartic and Tropical
  Lophocyrtid radiolarians, Micropaleontology, 44, 1-43, 1998.
- Sanfilippo, A., Westberg-Smith, M. J., and Riedel, W. R.: Cenozoic radiolaria, in: Plankton
  stratigraphy: Volume 2, Radiolaria, Diatoms, Silicoflagellates, Dinoflagellates and
  Ichthyoliths, edited by: Bolli, H. M., Saunders, J. B., and Perch-Nielsen, K., 631-712, 1985.
- Scher, H. D., Bohaty, S. M., Smith, B. W., and Munn, G. H.: Isotopic interrogation of a
  suspected late Eocene glaciation, Paleoceanography, 29, 2014PA002648,
  10.1002/2014PA002648, 2014.
- 27 Scher, H. D., Whittaker, J. M., Williams, S. E., Latimer, J. C., Kordesch, W. E., and Delaney,
- 28 M. L.: Onset of Antarctic Circumpolar Current 30 million years ago as Tasmanian Gateway
- aligned with westerlies, Nature, 523, 580-583, 2015.

- 1 Seton, M., Müller, R., Zahirovic, S., Gaina, C., Torsvik, T., Shephard, G., Talsma, A.,
- Gurnis, M., Turner, M., and Maus, S.: Global continental and ocean basin reconstructions
  since 200Ma, Earth Sci. Rev., 113, 212-270, 2012.
- Sexton, P. F., Wilson, P. A., and Norris, R. D.: Testing the Cenozoic multisite composite
  delta(18)O and delta(13)C curves: New monospecific Eocene records from a single locality,
  Demerara Rise (Ocean Drilling Program Leg 207), Paleoceanography, 21, PA2019,
  doi:10.1029/2005PA001253, 2006.
- Shackleton, N., and Kennett, J.: Paleotemperature history of the Cenozoic and the initiation
  of Antarctic glaciation: oxygen and carbon isotope analyses in DSDP Sites 277, 279, and
  281, in: Kennett, J.P., Houtz, R. E., et al., Initial reports of the deep sea drilling project, Vol.
  29, 743-755, 1975.
- 12 Spencer-Cervato, C.: The Cenozoic deep sea microfossil record: explorations of the 13 DSDP/ODP sample set using the Neptune database, Palaeontologia Electronica, 2, 270, 1999.
- Spiess, V.: Cenozoic magnetostratigraphy of Leg 113 drill sites, Maud Rise, Weddell Sea,
  Antarctica, Proceedings ODP, Scientific Results, 113, Ocean Drilling Program, College
  Station, TX, 261–315, doi:10.2973/odp.proc.sr.113.182.1990, 1990.
- Stickley, C. E., Brinkhuis, H., Schellenberg, S. A., Sluijs, A., Röhl, U., Fuller, M., Grauert,
  M., Huber, M., Warnaar, J., and Williams, G. L.: Timing and nature of the deepening of the
- 19 Tasmanian Gateway, Paleoceanography, 19, PA4027, doi:10.1029/2004PA001022, 2004.
- Sutherland, R.: The Australia-Pacific boundary and Cenozoic plate motions in the SW
  Pacific: Some constraints from Geosat data, Tectonics, 14, 819-831, 1995.
- Suzuki, N., Ogane, K., and Chiba, K.: Middle to Late Eocene polycystine radiolarians from
  the Site 1172, Leg 189, Southwest Pacific, News of Osaka Micropaleontologists, special
  volume, 14, 239-296, 2009.
- 25 Takemura, A.: Radiolarian Paleogene biostratigraphy in the southern Indian Ocean, Leg 120,
- 26 in: Wise, S.W. Jr., Schlich, R. et al., Proceedings ODP, Scientific Results, 120, Ocean
- 27 Drilling Program, College Station, TX, 735–756, doi:10.2973.odp.proc.sr.120.177, 1992.
- 28 Takemura, A., and Ling, H. Y.: Eocene and Oligocene radiolarian biostratigraphy from the
- 29 Southern Ocean correlation of ODP Legs 114 (Atlantic Ocean) and 120 (Indian Ocean),
- 30 Mar. Micropaleontol., 30, 97-116, 1997.

- 1 Torsvik, T. H., Van der Voo, R., Preeden, U., Mac Niocaill, C., Steinberger, B., Doubrovine,
- P. V., van Hinsbergen, D. J., Domeier, M., Gaina, C., and Tohver, E.: Phanerozoic polar
  wander, palaeogeography and dynamics, Ear. Sci. Rev., 114, 325-368, 2012.
- 4 van Hinsbergen, D. J., de Groot, L. V., van Schaik, S. J., Spakman, W., Bijl, P. K., Sluijs, A.,
- 5 Langereis, C. G., and Brinkhuis, H.: A Paleolatitude Calculator for Paleoclimate Studies,
- 6 PloS one, 10, e0126946, 2015.
- Villa, G., Fioroni, C., Pea, L., Bohaty, S., and Persico, D.: Middle Eocene–late Oligocene
  climate variability: calcareous nannofossil response at Kerguelen Plateau, Site 748, Mar.
- 9 Micropaleontol., 69, 173-192, 2008.
- Vonhof, H. B., Smit, J., Brinkhuis, H., Montanari, A., and Nederbragt, A. J.: Global cooling
  accelerated by early late Eocene impacts?, Geology, 28, 687-690, 2000.
- 12 Westerhold, T., Röhl, U., Pälike, H., Wilkens, R., Wilson, P., and Acton, G.: Orbitally tuned
- 13 time scale and astronomical forcing in the middle Eocene to early Oligocene, Clim. Past, 10,
- 14 955-973, doi:10.5194/cp-10-955-2014, 2014.
- Zachos, J. C., Quinn, T. M., and Salamy, K. A.: High-resolution (104 years) deep-sea
  foraminiferal stable isotope records of the Eocene-Oligocene climate transition,
  Paleoceanography, 11, 251-266, 1996.
- 18 Zachos, J. C., Pagani, M., Sloan, L., Thomas, E., and Billups, K.: Trends, rhythms, and
- aberrations in global climate 65 Ma to present, Science, 292, 686-693, 2001.
- 20

- 1 Table 1: Summary of species for which biogeographic affinities have been established and
- 2 their presence (x) at sites 277, 280, 281, 283, and 1172. H=high-latitude (>45°N/S), L=low-
- 3 latitude (<25°N/S) and C=cosmopolitan. Location of photographic images on plates for

# 4 selected species.

Таха	Biogeogr. affinity	Site 277	Site 280	Site 281	Site 283	ODP 1172	Plate
Amphicentria sp. 1 sensu Suzuki	Н	x		x	x	x	Pl. 2, Fig. 1
Amphicraspedum murrayanum Haeckel	L	х					Pl. 1, Fig. 14
Amphicraspedum prolixum Sanfilippo and Riedel gr.	L	х	х				Pl. 1, Fig. 15-17
Amphisphaera coronata (Ehrenberg) gr.	С	х			х	х	Pl. 1, Fig. 2
Amphisphaera spinulosa (Ehrenberg)	С	х			х		Pl. 1, Fig. 5
Amphymenium splendiarmatum Clark and Campbell	С	x	x	x	x		Pl. 1, Fig. 18,19
Antarctissa cylindrica Petrushevskaya	Н		x				
Antarctissa robusta Petrushevskaya	Н		х				
Aphetocyrtis bianulus (O'Connor)	Н	х			х	х	Pl. 5, Fig. 1
Aphetocyrtis gnomabax Sanfilippo and Caulet	Н	х	х	х	х		Pl. 5, Fig. 2-7
Aphetocyrtis rossi Sanfilippo and Caulet	Н	х	x		х		Pl. 5, Fig. 8-11
Artobotrys auriculaleporis (Clark and Campbell)	С	х				х	
Artostrobus annulatus (Bailey)	Н	х			х		
Artostrobus cf. pretabulatus Petrushevskaya	Н	х					Pl. 3, Fig. 13
Aspis sp. A sensu Hollis	Н	х	х		х		Pl. 3, Fig. 14-16
Axoprunum bispiculum (Popofsky)	Н	х			х		
Axoprunum pierinae (Clark and Campbell) gr.	С	х	х	х	х	х	Pl. 1, Fig. 10,11
Axoprunum? irregularis Takemura	Н	х				х	Pl. 1, Fig. 12
Ceratocyrtis spp.	Н	х	х		х	х	Pl. 2, Fig. 3-5
Cinclopyramis circumtexta (Haeckel)	С	х	х	х	х	х	
Clathrocyclas universa Clark and Campbell	С	х		х	х	х	
Clinorhabdus anantomus Sanfilippo and Caulet	Н	х		х	х		Pl. 5, Fig. 12,13
Clinorhabdus robusta (Abelmann)	Н					х	
Cornutella profunda Ehrenberg	С	х	х	х	х	х	
Cryptocarpium bussonii (Carnevale) gr.	С	х	х	х	х	х	Pl. 5, Fig. 25a,b, 26a,b
Cryptocarpium ornatum (Ehrenberg)	С	х			х		
Cycladophora cosma cosma Lombari and Lazarus	Н		х				Pl. 3, Fig. 17
Cycladophora humerus (Petrushevskaya)	Н		x	х	х		Pl. 3, Fig. 18
Cycladophora spp.	Н	х		х	х		
Cyrtolagena laguncula Haeckel	С	х			х		
Dictyophimus pocillum Ehrenberg	С	х					
Dictyophimus? aff. archipilium Petrushevskaya	Н	х		х	х		Pl. 4, Fig. 3a,b-8
Dictyophimus? archipilium Petrushevskaya	Н	х	х		х		Pl. 4, Fig. 1a,b, 2
Eucyrtidium antiquum Caulet	Н	х	х			х	Pl. 3, Fig. 19
Eucyrtidium mariae Caulet	Н	х					
Eucyrtidium nishimurae Takemura and Ling	Н			х	х	х	Pl. 3, Fig. 20a, b
Eucyrtidium spinosum Takemura	Н	х		х	х	х	Pl. 3, Fig. 21
Eucyrtidium montiparum Ehrenberg	С	х			х		Pl. 3, Fig. 22
Eurystomoskevos cauleti O'Connor	Н	х	х	х	х		Pl. 3, Fig. 23a, b
Eurystomoskevos petrushevskaae Caulet	Н	х	х	x	x	x	Pl. 3, Fig. 24
Eusyringium fistuligerum (Ehrenberg)	С	х				x	Pl. 3, Fig. 25
Eusyringium lagena (Ehrenberg)	С				x		
Glycobotrys nasuta (Ehrenberg) gr.	С	х	х	х	х	х	Pl. 3, Fig. 5-7
Lamprocyclas particollis O'Connor	Н	х	х	х	х		Pl. 5, Fig. 27
Larcopyle cf. pylomaticus (Riedel)	Н		x	х			Pl. 1, Fig. 25a, b
Larcopyle frakesi (Chen)	Н		x				Pl. 1, Fig. 20
Larcopyle hayesi (Chen)	Н	х	х	x	x		Pl. 1, Fig. 21
Larcopyle labyrinthusa Lazarus	Н		x				Pl. 1, Fig. 22
Larcopyle polyacantha (Campbell and Clark) gr.	Н	х	х	x	x		Pl. 1, Fig. 23, 24
Larcopyle spp.	Н	х	x	х			
Lithelius minor Jörgensen gr.	С	x	х	x	х	x	Pl. 1, Fig. 26-28
Lithomelissa challengerae Chen	Н		х				Pl. 2, Fig. 6-8
Lithomelissa ehrenbergi Bütschli	Н	x	х	х	х	х	Pl. 2, Fig. 10, 11
Lithomelissa gelasinus O'Connor	Н	x	x	x	x		Pl. 2, Fig. 12, 13
Lithomelissa robusta Chen	Н	1	х		х		Pl. 2, Fig. 16

Lithomelissa sphaerocephalis Chen	Н	х	х	х	х		Pl. 2, Fig. 17
Lithomelissa spp.	Н	х	х	х	х		
Lithomelissa tricornis	Н	х	х	х	х		Pl. 2, Fig. 18
Lithomelissa? sakaii O'Connor	Н		х				Pl. 2, Fig. 19
Lophocyrtis (Apoplanius) aspera (Ehrenberg)	Н	х		х	х		Pl. 5, Fig. 14a, b-16
Lophocyrtis (Apoplanius) keraspera Sanfilippo and Caulet	Н	х			х	х	Pl. 5, Fig. 17-19
Lophocyrtis (Lophocyrtis) jacchia hapsis Sanfilippo and Caulet	Н	х			х		Pl. 5, Fig. 20-22
Lophocyrtis (Paralampterium) dumitricai Sanfilippo	С	х				х	
Lophocyrtis (Paralampterium) longiventer (Chen)	Н	х	х	х	х	х	Pl. 5, Fig. 23, 24
Lophocyrtis spp.	Н				х		
Lophophaena capito Ehrenberg	С	х		х	х		
Lychnocanium amphitrite (Foreman)	С	х			х	х	Pl. 4, Fig. 11a, b, c, 12
Lychnocanium babylonis (Clark and Campbell)	С	х			х		Pl. 4, Fig. 13a, b, 14
Lychnocanium bellum Clark and Campbell	С	х			х	х	Pl. 4, Fig. 15, 16
Periphaena decora Ehrenberg	С	х	х	х	х	х	
Periphaena heliastericus (Clark and Campbell)	С	х	х	х	х	х	
Plectodiscus circularis (Clark and Campbell)	С	х	х	х	х	х	
Pseudodictyophimus galeatus Caulet	Н		х				Pl. 2, Fig. 20
Pseudodictyophimus gracilipes (Bailey) gr.	Н	x	x	x	x		Pl. 2, Fig. 21-23
Pseudodictyophimus spp.	Н		х				Pl. 2, Fig. 24-27
Sethocyrtis chrysallis Sanfilippo and Blome	С	х					Pl. 3, Fig. 26a, b
Siphocampe nodosaria (Haeckel)	С	х		х	х	х	
Siphocampe quadrata (Petrushevskaya and Kozlova)	С	х		х	х	х	
Siphocampe? amygdala (Shilov)	С	х			х		Pl. 3, Fig. 11, 12
Sphaeropyle tetrapila (Hays)	Н	х					Pl. 1, Fig. 29
Spirocyrtis joides (Petrushevskaya)	С	х	х	х	х		
Spongodiscus cruciferus (Clark and Campbell)	С	х		х		х	
Spongodiscus festivus (Clark and Campbell)	С	x				x	
Spongopyle osculosa Dreyer	Н	x	х	x	x	x	Pl. 1, Fig. 13
Spongurus bilobatus Clark and Campbell	С	х		х	х	х	
Stylosphaera minor Clark and Campbell gr.	С	х	х		х	х	Pl. 1, Fig. 7
Theocampe amphora (Haeckel)	С	х					
Theocampe urceolus (Haeckel)	С	х	х	х	х		
Theocyrtis tuberosa Riedel	L	х			х		Pl. 5, Fig. 30
Thyrsocyrtis pinguisicoides O'Connor	L	х			x		Pl. 3, Fig. 27
Tripodiscinus clavipes (Clark and Campbell)	С	х		x	x		
Zealithapium mitra (Ehrenberg)	С	х			х		Pl. 1, Fig. 8

1 Table 2: Average of total % of high-latitude species, groups, genera and high-latitude

2 members of families for five time slices: Middle Eocene Climatic Optimum (MECO,

3 ~40 Ma), early late Eocene/PrOM (~38–37 Ma), late Eocene warming event (~36 Ma), latest

4 Eocene-earliest Oligocene (~35–32 Ma) and early Oligocene (~30 Ma).

5

	Site 277					Site 280	Site 281	Site 283		Site 1172	
	40 Ma	38-37 Ma	36 Ma	35-32 Ma	30 Ma	e. Olig.	38-37 Ma	38-37 Ma	36 Ma	40 Ma	
% high-latitude species	23.2	28.9	13.7	39.0	100.0	62.6	61.2	28.0	25.6	7.8	
Larcopyle spp. %	6.9	2.9	2.5	6.2	-	18.4	26.5	3.0	1.8	-	
Lithomelissa spp. %	0.1	1.8	0.1	5.9	-	16.4	11.8	4.1	4.8	0.4	
High-lat. Lophocyrtiidae %	14.9	20.8	8.0	16.7	3.3	10.4	14.2	8.5	6.9	4.1	
High-lat. Eucyrtidiidae %	-	0.4	0.5	1.8		8.8	6.3	7.4	9.1	3.3	
Other high-lat. Plagiacanthidae %	-	0.2	-	1.4		6.5	1.4	1.8	1.5	-	
Other high-lat. species %	1.3	2.8	2.7	7.0	96.7	2.0	1.1	3.1	1.5	0.1	
% cosmopolitan species	72.6	71.1	80.9	59.9	-	37.3	38.8	71.8	65.2	92.2	
% low-latitude species	4.2	0.1	5.4	1.0	-	0.1	-	0.1	9.2	-	

6

# 1 Figures



Figure 1. Modern location of DSDP and ODP study sites in the Southwest Pacific; dark
grey=coastline, light grey=2000 m isobath of continental boundary, STF=Subtropical Front,
SAF=Subantarctic Front, SAW=Subantarctic Water.





3 Figure 2. DSDP Site 277 stratigraphy, NZ Stages (Raine et al. (2015), Southern Hemisphere 4 radiolarian zones (RP), nannofossil zones (NP), lithology, core recovery, selected bioevents (ages calibrated to the 2012 geological timescale; Gradstein et al., 2012; Raine et al., 2015) 5 and benthic  $\delta^{18}$ O and  $\delta^{13}$ C data of DSDP Site 277. The dashed lines correlate Site 277 based 6 7 on the ages of the bioevents to Southern Ocean Cibicidoides data of ODP Site 689 Hole B (Maud Rise) (Diester-Haass and Zahn, 1996) calibrated to the GTS2012 timescale using the 8 9 magnetostratigraphy data of Florindo and Roberts (2005) and Spiess (1990). LO=Lowest Occurrence; LCO=Lowest Common Occurrence; HO=Highest Occurrence; MECO=Middle 10 Eocene Climatic Optimum; PrOM=Priabonian Oxygen Isotope Maximum; EOT=Eocene-11 12 Oligocene transition.





Figure 3. DSDP Site 277  $\delta^{18}$ O and  $\delta^{13}$ C records and location of studied radiolarian samples within the MECO and late Eocene warming interval (red stars) and radiolarian-rich upper Eocene-lower Oligocene interval (blue stars).





Figure 4. DSDP Site 277 benthic  $\delta^{18}$ O record; radiolarian abundance and diatom/radiolarian 3 4 (D/R) ratio; Taxic richness (number of taxa) derived from individual rarefaction and range-5 through analyses for different sample sizes; Fisher a Index and Simpson Evenness Index for 6 radiolarian assemblages. Red bars indicate sample sizes <100 specimens, blue bars for 7 sample sizes <300 specimens and black bars for samples sizes >300 specimens. 8



Figure 5. Stratigraphy, Southern Hemisphere radiolarian zones (RP), lithology and core
recovery at DSDP sites 280, 281 and 283. Variation in radiolarian abundance,
diatom/radiolarian (D/R) ratio, Fisher α Index and Simpson Evenness for radiolarian
assemblages at all sites.



Figure 6. Biogeographic affinities of radiolarian assemblages at DSDP Site 277 and the
abundance of high-latitude taxa/families. MECO=Middle Eocene Climatic Optimum,
PrOM=Priabonian Oxygen Isotope Maximum, EOT=Eocene-Oligocene transition.



Figure 7. Variation in faunal affinities for radiolarians assemblages and Fisher α Diversity at all sites. Dashed black lines indicate correlation between sites. The location of the MECO at Site 1172 is taken from Bijl et al. (2010).



