UNIVERSITY^{OF} BIRMINGHAM University of Birmingham Research at Birmingham

Lipid biomarker distributions in Oligocene and Miocene sediments from the Ross Sea region, Antarctica

Duncan, Bella; McKay, Robert; Bendle, James; Naish, Timothy; Inglis, Gordon N.; Moossen, Heiko; Levy, Richard; Ventura, G. Todd; Lewis, Adam; Chamberlain, Beth; Walker, Carrie

DOI: 10.1016/j.palaeo.2018.11.028

License: Creative Commons: Attribution-NonCommercial-NoDerivs (CC BY-NC-ND)

Document Version Peer reviewed version

Citation for published version (Harvard): Duncan, B, McKay, R, Bendle, J, Naish, T, Inglis, GN, Moossen, H, Levy, R, Ventura, GT, Lewis, A, Chamberlain, B & Walker, C 2019, 'Lipid biomarker distributions in Oligocene and Miocene sediments from the Ross Sea region, Antarctica: Implications for use of biomarker proxies in glacially-influenced settings',

Palaeogeography, Palaeoclimatology, Palaeoecology, vol. 516, pp. 71-89. https://doi.org/10.1016/j.palaeo.2018.11.028

Link to publication on Research at Birmingham portal

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?) •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

1 Lipid biomarker distributions in Oligocene and Miocene sediments from

2 the Ross Sea region, Antarctica: Implications for use of biomarker proxies

3 in glacially influenced settings

- 4 Bella Duncan^a, Robert McKay^a, James Bendle^b, Timothy Naish^a, Gordon N. Inglis^c, Heiko
- 5 Moossen^{b,d}, Richard Levy^e, G. Todd Ventura^{e,f}, Adam Lewis^g, Beth Chamberlain^{b,h}, Carrie Walker^{b,i}.

6

- ^aAntarctic Research Centre, Victoria University of Wellington, P.O. Box, Wellington 6012, New
 Zealand
- ^bSchool of Geography, Earth and Environmental Sciences, University of Birmingham, Edgbaston,
 Birmingham, B15 2TT, UK
- 11 ^cOrganic Geochemistry Unit, School of Chemistry and Cabot Institute, University of Bristol,
- 12 Cantock's Close, Bristol BS8 1TS, UK

¹³ ^dPresent address: Max Planck Institute for Biogeochemistry, P.O. Box 10 01 64, 07701 Jena,

- 14 Germany
- ^eGeological and Nuclear Sciences, P.O. Box 30-368, Lower Hutt 5040, New Zealand
- ^fPresent address: Department of Geology, Saint Mary's University, 923 Robie Street, Halifax, Nova
 Scotia, B3H 3C3, Canada
- 18 ^gNorth Dakota State University, Fargo, North Dakota, 58105, USA

^hPresent address: Department of Life Sciences, Imperial College London, Silwood Park Campus,
 Buckhurst Road, BioAscot, Berkshire, SL5 7PY, UK

- 21 ⁱPresent address: School of Environment, Earth and Ecosystem Sciences, The Open University,
- 22 Milton Keynes, MK7 6AA, UK
- 23
- 24 Corresponding author: Bella Duncan, bella.duncan@vuw.ac.nz
- 25
- 26
- 27
- 28
- 29
- -
- 30
- 31

32 Abstract

33 Biomarker-based climate proxies enable climate and environmental reconstructions for regions where 34 other paleoclimatic approaches are unsuitable. The Antarctic Cenozoic record consists of widely 35 varying lithologies, deposited in rapidly changing depositional settings, with large lateral variations. Previous sedimentological and microfossil studies indicate that the incorporation of reworked older 36 37 material frequently occurs in these sediments, highlighting the need for an assessment of biomarker distribution across a range of depositional settings and ages to assess the role reworking may have on 38 39 biomarker-based reconstructions. Here, we compare sedimentary facies with the distribution of *n*-40 alkanes and hopanoids within a terrestrial outcrop, two glaciomarine cores and a deep sea core, 41 spanning the Late Oligocene to Miocene in the Ross Sea. Comparisons are also made with *n*-alkane 42 distributions in Eocene glacial erratics and Mesozoic Beacon Supergroup sediments, which are both potential sources of reworked material. The dominant *n*-alkane chain length shifts from $n-C_{29}$ to $n-C_{27}$ 43 between the Late Eocene and the Oligocene. This shift is likely due to changing plant community 44 45 composition and the plastic response of *n*-alkanes to climate cooling. Samples from glaciofluvial environments onshore, and subglacial and ice-proximal environments offshore are more likely to display 46 47 reworked *n*-alkane distributions, whereas, samples from lower-energy, lacustrine and ice-distal marine 48 environments predominantly vield immature/contemporaneous *n*-alkanes. These findings emphasise 49 that careful comparisons with sedimentological and paleontological indicators are essential when 50 applying and interpreting *n*-alkane-based and other biomarker-based proxies in glacially-influenced 51 settings.

52 Keywords: Paleoclimate, Antarctica, *n*-alkanes, biomarkers, hopanoids, reworking

53 1 Introduction

54 In Antarctic sediments, traditional microfossil-based methods of reconstructing climate can be challenging due to sparse distribution, low diversity of species, or poor preservation in sediments (i.e. 55 56 Askin and Raine, 2000; Strong and Webb, 2000; Scherer et al., 2007). In contrast, biomarkers 57 (molecular fossils preserved in the geological record) are relatively recalcitrant and have the potential to provide environmental proxy information when other methods are challenging or unsuitable. To 58 59 date, only a few studies have employed biomarkers to investigate paleoclimate changes in Antarctica, 60 with most of these conducted in offshore settings (i.e. Feakins et al., 2012; McKay et al., 2012; Pross 61 et al., 2012; Bijl et al., 2013; Feakins et al., 2014; Levy et al., 2016; Rees-Owens et al., 2018). Such 62 work is challenging because Cenozoic outcrops exposed in Antarctica are sparse, and are represented 63 by relatively superficial and poorly-dated deposits of glacially derived tills, lacustrine and fluvial 64 deposits, with occasional marine and glaciomarine sediments (Hambrey and Barrett, 1993; Marchant and Denton, 1996; Lewis et al., 2007; Lewis et al., 2008; Lewis and Ashworth, 2016). Sediments from 65 66 drillcores on the continental margin are usually glaciomarine in origin and provide better-dated

- 67 records of cyclical fluctuations of the Antarctic Ice Sheets (Barrett, 1989; Naish et al., 2001, 2009).
- 68 However, the variable lithologies in these sediments and the nature of their deposition mean that
- 69 reworking of older sediments and associated fossil material is potentially a significant issue (e.g.,
- 70 Kemp and Barrett, 1975; Askin and Raine, 2000; Prebble et al., 2006a).



71

Fig. 1: Location of sample sites in the Ross Sea region of Antarctica. WAIS: West Antarctic
Ice Sheet, EAIS: East Antarctic Ice sheet, DSDP: Deep Sea Drilling Project, CRP: Cape Roberts
Project, McMurdo S.: McMurdo Sound, V.L. Basin: Victoria Land Basin. Base map from
Quantarctica GIS package, Norwegian Polar Institute.

76 Here, lipid biomarkers (n-alkanes and hopanoids) are used to investigate how organic matter 77 varies between different lithologies and depositional environments in the Ross Sea region of 78 Antarctica. Specifically, we aim to assess whether lipid biomarkers represent organic material sourced 79 from organisms living contemporaneously with sediment deposition, or older organic material which 80 has been reworked into the sediment. Knowledge of potential reworking is critical for using and 81 interpreting biomarker-based paleoenvironmental proxies in glacially-influenced settings. Localities 82 and sediment drill cores were chosen to survey a range of depositional environments that together 83 form a transect from high elevation terrestrial deposits to the deep sea (Fig. 1). These include; (i) A

Middle Miocene (~14 Ma) lacustrine/fluvial sequence from a small mountain glacier catchment at Mt
Boreas in the Transantarctic Mountains. (ii) A Late Oligocene/Early Miocene glaciomarine sequence
in the shallow marine Cape Roberts Project 2/2A drill core, sampling a coastal sediment catchment
from an East Antarctic Ice Sheet (EAIS) outlet glacier. (iii) A deeper water Late Oligocene/Early
Miocene glaciomarine sequence in DSDP Site 270 sampling sediment sourced from now submerged

89 islands and ice caps in the central continental shelf of the Ross Sea, West Antarctica. (iv) An Early

90 Miocene to Late Miocene marine sequence from DSDP Site 274 from the Western Ross Sea abyssal

91 plain, sampling a wide sediment source catchment from both East and West Antarctica.

92 1.1 Geological setting

93 The western Ross Sea region of Antarctica is bounded by the Transantarctic Mountains 94 (TAM), which were uplifted in the early Cenozoic, with the bulk of their exhumation occurring before 95 the early Oligocene (Fitzgerald, 1994; Smellie, 2001). The basement rocks of the TAM are dominated 96 by Archean to mid-Paleozoic metasediments and intrusives (Allibone et al., 1993a; Allibone et al., 97 1993b; Goodge et al., 2002). The Devonian to Triassic Beacon Supergroup overlies this basement 98 (Barrett, 1981). Lithologies vary through the sequence, with interbedded sandstones, shales, 99 conglomerates and coals deposited in a paleoenvironmental setting moving from shallow marine to a 100 terrestrial system of lakes, braided rivers and alluvial plains (Barrett, 1981). Plant macrofossils and 101 palynomorphs are common throughout the Beacon Supergroup (Barrett, 1981). In the early Jurassic, 102 as the Gondwana super-continent began to separate, the Beacon Supergroup was intruded by the 103 Ferrar Dolerite, resulting in extensive low grade thermal metamorphism (Barrett, et al., 1986). The 104 Jurassic Ferrar Group contains extrusive volcanic rocks with fossiliferous sedimentary interbeds 105 containing terrestrial microfossil assemblages (e.g. Ribecai, 2007).

106 Scattered sedimentary outcrops and the basaltic McMurdo Volcanic Group form the Cenozoic 107 geology of the TAM (Marchant and Denton, 1996; Fielding et al., 2006; Martin et al, 2010). A significant Cenozoic sedimentary unit distributed throughout the TAM is the Sirius Group, which 108 109 comprises glacial and non-glacial sediments with well-preserved fossil woody vegetation, leaf 110 material and peat beds, deposited in terrestrial and proximal marine environments (Hambrey and Barrett, 1993; Francis and Hill, 1996; Barrett, 2013). Scattered Early Miocene to Holocene veneers of 111 112 glacial tills, colluvium and lacustrine deposits are dispersed through the TAM (Marchant and Denton, 113 1996; Lewis et al., 2007; Lewis et al., 2008; Lewis and Ashworth, 2016). Eocene to Pliocene glacial 114 erratics are found in the McMurdo region (Harwood and Levy, 2000). Much of what is currently 115 known about Cenozoic Antarctic climate is based on seismic stratigraphy and continental margin 116 drilling. Infill of sedimentary basins in the Ross Sea potentially began as early as the Late Cretaceous, with sediment accumulation continuing through the Cenozoic (Cooper et al., 1987; De Santis et al., 117 1995; Luyendyk et al., 2001; Decesari et al., 2007; Wilson and Luyendyk, 2009). Continental margin 118

- drill cores from the Ross Sea contain successions of subglacial, glaciomarine and marine sediments
- reflecting the cyclical advance and retreat of the Antarctic Ice Sheets (i.e. Barrett, 1989; Naish et al.,
- 121 2001; Naish et al., 2009; McKay et al., 2009; Levy et al., 2016).

122 **2 Methods**

123 2.1 Site description

124 This work utilises samples from: (i) The McMurdo glacial erratics from the Mt Discovery and

- 125 Minna Bluff region (Fig. 1), which yield marine and terrestrial micro- and macrofossils of mid-late
- 126 Eocene age and are interpreted as being deposited in coastal-terrestrial and nearshore marine
- 127 environments, under ice-free conditions (Harwood and Levy, 2000). (ii) Mid-Miocene terrestrial,
- 128 fossil-bearing strata from Mt Boreas in the Olympus Range, which record the last known vestige of
- 129 vegetation in the TAM before the Dry Valleys transitioned from wet- to cold-based glaciation at high
- 130 altitudes (1,425 m) (Fig. 1) (Lewis et al., 2008). (iii) Oligocene/Early Miocene glaciomarine
- 131 sediments obtained from the Cape Roberts Project core CRP-2/2A from the Victoria Land continental
- 132 slope of Antarctica (Fig. 1) (Cape Roberts Science Team, 1999). (iv) A Late Oligocene to Early
- 133 Miocene glaciomarine sequence of sediments from DSDP Site 270, drilled on the continental shelf in
- the central Ross Sea in 1973 and re-described in 2015 (Fig. 1) (The Shipboard Scientific Party, 1975a;
- 135 Kraus, 2016). (v) An Early to Late Miocene succession of ice-distal diatom-rich silty clay sediments
- from DSDP 274 on the lower continental rise in the northwestern Ross Sea (68°59.81'S,
- 137 173°25.64'W) (Fig. 1) (The Shipboard Scientific Party, 1975b). Figure 2 schematically describes the
- sampling sites and facies used in this study.



140 Fig. 2. Schematic representation of Oligocene and Miocene environments of deposition in the Ross Sea Region, and their associated sedimentary

facies. Sample sites are placed in their representative depositional setting.

142

143 2.2 Bulk analysis

144 Unless already desiccated, samples were freeze dried for 48 h prior to sample work up. All145 samples were homogenised to a powder using a Retsch 200 mixer mill.

146 Pyrolysis measurements for total organic carbon (TOC) were made using a Weatherford laboratories Source Rock Analyzer at GNS Science on ~100 mg of powdered sediment. The 147 pyrolysis program was set with the sample crucible entering the pyrolysis oven where it was held 148 149 isothermal at 300 °C for 3 mins under a continuous stream of He carrier gas using a 100 ml/min flow rate. This was followed by a 25 °C/min ramp to 650 °C. The S1 and S2 signal intensities were 150 recorded with a FID operated under a 65 ml/min stream of H₂ gas and 300 ml/min air. The pyrolysis 151 cycle was then followed by an oxidation cycle performed at 630 °C for 20 mins during which time the 152 153 oven and crucible were flushed with dry air at 250 ml/min. The generated carbon monoxide and 154 carbon dioxide gases were measured by the instrument's IR cells. All sample sequences were run 155 with three IFP 160000 analytical standard replicates (from Vinci Technologies, Institut Français du 156 Pétrole) placed at the beginning, middle and end of each sample sequence.

157 2.3 Lipid biomarker analyses

158 Organic geochemical work-up, gas chromatograph (GC)-flame ionization detector (FID) and 159 GC-Mass Spectrometer (MS) analyses was performed in the Birmingham Molecular Climatology 160 Laboratory (BMC), University of Birmingham. Lipids were extracted from ~10-15 g of homogenised sediment by ultrasonic extraction using dichloromethane (DCM):methanol (3:1). The total lipid 161 162 extract was fractionated by silica gel chromatography using *n*-hexane, *n*-hexane:DCM (2:1), DCM, and methanol to produce four separate fractions, the first of which contained the aliphatic saturated 163 164 and unsaturated hydrocarbons (e.g. n-alkanes, steranes and hopanes). Procedural blanks were also 165 analysed to ensure the absence of laboratory contaminants.

166 The aliphatic hydrocarbon fractions were analysed on an Agilent 7890B series GC, equipped 167 with a 7639ALS autosampler, a BP5-MS column (SGE Analytical Science, 60 m \times 0.32 mm \times 0.25 168 μ m) and an FID, using hydrogen (H₂) as a carrier gas. Compound separation was achieved by using 169 the following temperature program: the oven was held at 70 °C for 1 min, then heated to 120 °C at 30 170 °C/min, and then to 320 °C with 3 °C/min, where it was held for 20 mins. GC-Mass spectrometry (GC-MS) was performed using an Agilent 7890B GC, coupled to an Agilent 5977A Mass Selective 171 172 Detector (MSD). The same capillary column and temperature program was used throughout the 173 analyses for consistent compound separation. Helium (He) was used as a carrier gas. Samples were 174 bracketed with an external standard containing known abundances of certain *n*-alkanes to allow 175 identification and quantification of *n*-alkanes (average standard deviation of \pm 7.6%). *n*-Alkane peaks

- were integrated in Agilent OpenLAB Data Analysis Version A.01.01 Build 1.93.0. Relationships
- between *n*-alkane indices were investigated using Pearson's correlation coefficients and assessed as
- 178 statistically significant when p < 0.05. Hopanes and hopenes were identified based upon published
- spectra, characteristic mass fragments and retention times (e.g. Rohmer et al., 1984; Sessions et al.,
- 180 2013; Inglis et al., 2018) and integrated using GC-MS.

181 2.4 Biomarker indices

182 *n*-Alkanes of specific carbon chain lengths are known to be derived from discrete biological 183 sources. Algae and some photosynthetic bacteria typically produce dominantly $n-C_{17}$, with lesser 184 amounts of *n*-C₁₅ and *n*-C₁₉ (Clark and Blumer, 1967; Han and Calvin, 1969; Cranwell et al., 1987). 185 Other species of bacteria, including non-photosynthetic bacteria often demonstrate an even carbon 186 number preference between $n-C_{12}$ and $n-C_{22}$, commonly with high $n-C_{16}$ and $n-C_{18}$ (Han and Calvin, 1969; Grimalt and Albaigés, 1987). Non-emergent aquatic plants and Sphagnum mosses show 187 188 enhanced production of n-C₂₃ and n-C₂₅ (Baas et al., 2000; Ficken et al., 2000; Pancost et al., 2002; 189 Bingham et al., 2010). Long chain *n*-alkanes (*n*-C₂₅ and higher), usually with a high odd-over-even 190 predominance, are most abundant in the epicuticular waxes on leaves and stems of terrestrial higher 191 plants (Eglinton and Hamilton, 1963). n-Alkanes are also derived from the early diagenetic alteration of saturated and unsaturated aliphatic alcohols, ketones, esters, and di- or triterpenic acids (i.e. Tissot 192 193 and Welte, 1984; Meyers and Ishiwatari 1993). Once deposited, n-alkanes may undergo microbial or 194 geochemical alteration, modifying their distributions (Grimalt et al., 1985).

The source and maturity of higher molecular weight *n*-alkanes can be characterised by theircarbon preference index (CPI):

197
$$CPI = \frac{1}{2} \left(\left(\frac{\Sigma_{odd}(n - C_{25-33})}{\Sigma_{even}(n - C_{24-32})} \right) + \left(\frac{\Sigma_{odd}(n - C_{25-33})}{\Sigma_{even}(n - C_{26-34})} \right) \right)$$
(1)

198 Most modern sediments with terrestrially sourced organic matter have an odd-over-even 199 predominance of long chained *n*-alkanes (n-C₂₅ to n-C₃₄) and CPI values >1 (Bray and Evans, 1961; 200 Eglinton and Hamilton, 1963). A survey of modern leaf wax material demonstrates that a CPI of >1-2 201 is a reasonable threshold value indicative of relatively unmodified terrestrial plant material (Bush and McInerney, 2013). Sediments containing CPI values of <1 usually indicate either exposure to elevated 202 203 burial temperatures great enough to cause hydrocarbon cracking, or an input of organic matter that has been altered by diagenetic or catagenetic processes (Bray and Evans, 1961). Some sediments will also 204 display an unresolved complex mixture (UCM), represented by a hump in the baseline of a gas 205 206 chromatogram due to the co-elution of unresolved compounds (Gough and Rowland, 1990; Gough et 207 al., 1992). UCMs can be especially prominent in biodegraded petroleums, in which microbial degradation of the more abundant aliphatic components leads to increased concentrations of the more 208 209 recalcitrant, branched and cyclic compounds (Gough and Rowland, 1990; Gough et al., 1992). In

- 210 recent sediments, *n*-alkanes have a lower susceptibility to microbial degradation than most other types
- of organic matter as they lack functional groups, but studies on peat and lake sediments suggest that
- 212 microbial degradation does occur (Meyers and Ishiwatari, 1993; Lehtonen and Ketola, 1993). Shorter
- chain lengths appear more degradable than longer chain lengths, and microbial degradation can result
- in a decrease in CPI (Meyers and Ishiwatari, 1993; Lehtonen and Ketola, 1993).
- Average chain length (ACL) indicates the dominant *n*-alkane in a given carbon number range
 (Poynter et al., 1989; Schefuß et al., 2003):

217
$$ACL = \frac{\Sigma(C_{odd\ 25-33}, x_{odd\ 25-33})}{(x_{odd\ 25-33})}$$
(2)

218 Where $C_{odd 25-33}$ represents the carbon number of the odd chain length *n*-alkanes, and $x_{odd 25-33}$ 219 represents the concentrations of the odd *n*-alkanes in the sample. ACL is influenced by a number of 220 factors. Higher ACLs are typical of warmer, tropical regions, whilst lower ACLs are more commonly 221 observed from cooler, temperate regions, indicating that ACL could be related to air temperature 222 (Gagosian and Peltzer, 1986; Poynter et al., 1989; Dodd and Afzal-Rafii, 2000; Kawamura et al., 2003; Bendle et al., 2007; Vogts et al., 2009; Bush and McInerney, 2015). Other studies have 223 224 suggested that aridity has a strong control on ACL, with the synthesis of longer *n*-alkanes in more arid 225 environments providing plants with a more efficient wax coating to restrict water loss (Dodd et al., 1998; Dodd and Afzal-Rafii, 2000; Schefuß et al., 2003; Calvo et al., 2004; Zhou et al., 2005; 226 227 Moossen et al., 2015). ACL is also strongly controlled by the contributing vegetation, with large interand intra-species variation in *n*-alkane distributions (i.e. Vogts et al., 2009; Bush and McInerney, 228 229 2013; Feakins et al., 2016). Variation in average chain length through time therefore reflects the 230 interplay of two key factors: climate-driven plastic response of *n*-alkanes to temperature and/or aridity 231 within a plant community; or changes to the composition of the plant community, often in response to 232 climate (Bush and McInerney, 2013).

Hopanes and hopenes are C₂₇ to C₃₅ pentacyclic triterpenoids derived from a wide range of 233 234 bacteria (Rohmer et al., 1984; Talbot and Farrimond, 2007). Hopanes are ubiquitous across a variety 235 of depositional settings, and in both modern and ancient sediments (Ourisson and Albrecht, 1992). In 236 modern sediments, hopenes are mostly present in the biological 17β , 21β (H) configuration (although there are exceptions; see Inglis et al., 2018). In sediments, with increasing diagenesis, hopanes 237 238 undergo stereochemical transformations and the biologically-derived 17β , 21β (H)-hopanoid is 239 transformed into the more thermally stable $17\beta_{21\alpha}(H)$ and $17\alpha_{21\beta}(H)$ -stereoisomers (Mackenzie et 240 al., 1980; Peters and Moldowan, 1991). With increasing maturation, extended hopanoids ($>C_{30}$) also 241 undergo isomerisation at the C-22 position. As such, hopanoids are frequently used to reconstruct 242 thermal maturity (Mackenzie et al., 1980; Seifert and Moldowan, 1980; Peters and Moldowan, 1991; Farrimond et al., 1998), where decreasing $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ indices and increasing 22S/(22R+22S)243 values indicate increasing thermal maturity. The ratio of C_{27} 18 α (H)-trisnorhopane II (Ts) to C_{27} 244

245 $17\alpha(H)$ -trisnorhopane (Tm) is also commonly used as a maturity parameter, as Tm is less stable 246 during catagenesis than Ts (Seifert and Moldowan, 1978).

247 3 Results

248 3.1 Facies compilation

249 In order to compare results between sites, an internally consistent facies scheme was 250 developed based on published facies descriptions for each sample site (Fig. 2). The McMurdo erratics 251 were not included, as the context and relationship between these samples is uncertain. Instead these 252 samples were labelled by lithofacies as described by Levy and Harwood (2000) (Supplementary table 1). Samples from Mt Boreas were assigned facies based on descriptions from Lewis et al. (2008). 253 254 Facies for CRP 2/2A were developed based on descriptions of Fielding et al. (2000). Facies for DSDP 270 were assigned using descriptions of Kraus (2016), and based on previous models of glaciomarine 255 facies successions (Fielding et al. 2000; Powell and Cooper, 2002; McKay et al., 2009). For DSDP 256 274, facies were determined by using interpretations from The Shipboard Scientific Party (1975b), 257 258 Frakes (1975) and Whittaker and Müller (2006).

259 3.2 McMurdo erratics

Six Mid- Late Eocene sediment samples from the McMurdo erratics suite were analysed for 260 *n*-alkanes, sourced from a range of lithofacies (Levy and Harwood, 2000) (Supplementary table 1). 261 Three of the samples (E214, MB245 and E215) are dominated by the $n-C_{17}$ to $n-C_{20}$ short chained n-262 alkanes underlain by a UCM (Fig. 3). Samples D1, E219 and MTD95 have bimodal profiles, 263 dominated by $\sim n-C_{20}$ to $n-C_{23}$ underlain by small a UCM, and a series of longer *n*-alkanes with a 264 mode at $\sim n-C_{29}$ (Fig. 3). In all samples, $n-C_{29}$ is the dominant long-chained *n*-alkane, in contrast to 265 younger strata investigated in this study usually dominated by $n-C_{27}$. The ratio between these two 266 chain lengths has been described at all sites to investigate its variance at other localities. The CPI from 267 268 the McMurdo Erratics ranges from 1.8 to 5.5 (avg. 2.8). ACL varies from 27.9 to 28.7 (avg. 28.2), whilst the ratio of the n-C₂₉ n-alkane to n-C₂₇ ranges from 1.01 to 1.64 (avg. 1.26). The erratics 269 270 contain total abundances of *n*-alkanes ranging from 83 µg/gTOC to 1579 µg/gTOC, at an average of

271 406 μg/gTOC.



Fig. 3. Representative GC-FID-chromatograms of two samples of the McMurdo erratics.
Filled circles above peaks indicate n-alkanes, with the n-C₂₇ and n-C₂₉ labelled. UCM: unresolved
complex mixture.

276 *3.3 Mt Boreas*

272

277 12 sediment samples from Mt Boreas were analysed for *n*-alkanes (Supplementary table 1) Samples from Mt Boreas are typically dominated by long chained *n*-alkanes, particularly *n*-C₂₃, *n*-C₂₅ 278 and n-C₂₇ (Fig. 4). Some samples display small UCMs, usually underlying ~n-C₁₉ to n-C₂₀. Samples 279 were collected from three different sites within a topographic depression which held a small alpine 280 281 line, from units that are correlatable to the stratigraphic column shown in Figure 4 (Lewis et al., 282 2008). The total abundance of *n*-alkanes at these sites ranges between 4.5 μ g/gTOC to 762 μ g/gTOC, at an average of 206.5 µg/gTOC. The CPI of the long-chained *n*-alkanes ranges from 1.7 to 5.9 (avg. 283 3.4), whilst ACL varies from 26.2 to 27.4 (avg. 26.9). The ratio of the n-C₂₉ n-alkane to n-C₂₇ varies 284 from 0.15-0.97 (avg. 0.55), indicating that the *n*-C₂₇ dominates the *n*-C₂₉ in all samples from these 285 sites (Supplementary table 1). 286



Fig. 4. Stratigraphic column from a site at Mt Boreas (after Lewis et al., 2008) with the
equivalent stratigraphic positions of representative GC-FID-chromatograms of samples. Filled

- 290 *circles above peaks indicate n-alkanes, with the n-C* $_{27}$ and *n-C* $_{29}$ *labelled. Facies numbers are*
- 291 *described in Fig. 2. UCM: unresolved complex mixture.*

Pearson's correlation coefficients were estimated for each *n*-alkane variable compared to other *n*-alkane variables from these sites. Only two variables demonstrate a statistically significant correlation to each other; the ratio of $n-C_{29}/n-C_{27}$ typically decreases with increasing CPI (r = 0.711, p= 0.0096) (Fig. 5). Samples from fluvial Facies 2 typically display lower CPI, and higher ACL and *n*- $C_{29}/n-C_{27}$ values than those from lacustrine Facies 1, although it is noted that the Facies 2 is only represented by two samples (Fig. 6). Both fluvial and lacustrine samples show similar average total abundances of *n*-alkanes.



299

300 Fig. 5. Scatter plots of samples from Mt Boreas; a) CPI and ACL; b) CPI and $n-C_{29}/n-C_{27}$; c) 301 $n-C_{29}/n-C_{27}$ and ACL; d) CPI and the total abundance of n-alkanes (μ g n-alkanes/g TOC); e) ACL and 302 the total abundance of n-alkanes (μ g n-alkanes/g TOC) and f) $n-C_{29}/n-C_{27}$ and the total abundance of 303 n-alkanes (μ g n-alkanes/g TOC).

Three samples representing typical *n*-alkane distributions from the site were also analysed for additional biomarkers (Supplementary table 2). In two samples (ALS-05-21N and ALS-05-04C), hopanoids were abundant and the distribution was dominated by $17\beta(H)$ -trisnorhopane (C₂₇) and $17\beta,21\beta(H)$ -norhopane (C₂₉). Both samples are characterised by high $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ ratios (0.82 to 0.86) and indicate low thermal maturity. Within sample ALS-05 21O, hopanoids were weak and the distribution was dominated by thermally-mature C₂₇ to C₃₅ hopanes. This sample was characterised by

- 310 a low $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ ratio (0.07) and high C22S/C22R+C22S ratio (0.58) and therefore indicate high
- 311 thermal maturity.



Fig. 6: Distributions of n-alkane variables across different facies from Mt Boreas, CRP-2/2A,
DSDP 270 and DSDP 274; a) CPI, b) ACL, c) n-C₂₉/n-C₂₇ and d) the total abundance of n-alkanes
(µg n-alkanes/g TOC). Grey bars represent average values for each facies. Description of facies in
Fig. 2.

317 3.4 Cape Roberts Project 2/2A

Long chained *n*-alkanes typically dominate samples from CRP 2/2A (Fig. 7). The *n*-C₂₃, *n*-C₂₅ and *n*-C₂₇ are usually the most abundant homologs, with *n*-C₂₇ often the most prominent of these. In some samples *n*-alkanes elute with a UCM, which is usually centred between *n*-C₁₉ and *n*-C₂₃. The total abundance of *n*-alkanes is highly variable from sample to sample, ranging from 16.6 μ g/gTOC to 1893.0 μ g/gTOC, averaging 197.6 μ g/gTOC. CPI also varies over a wide range, from 1.2-5.5 (avg. 2.6), whilst ACL ranges from 26.7-28.7 (avg. 27.7). The ratio of the *n*-C₂₉*n*-alkane to *n*-C₂₇ varies

324 from 0.25-1.02 (avg. 0.67) (Supplementary table 1).

325 Pearson's correlation coefficient estimations show statistically significant correlations 326 between several *n*-alkane variables. The strongest correlations exist between CPI and $n-C_{29}/n-C_{27}$ (*r* = 327 0.837, p < 0.0001) and ACL and $n-C_{29}/n-C_{27}$ (r = 0.777, p < 0.0001), with a weaker correlation 328 between CPI and ACL (r = 0.580, p = 0.0091). Figure 8 shows that at both high and low values of 329 CPI and $n-C_{29}/n-C_{27}$, the total abundance of *n*-alkanes increases, with weak correlations between these variables (r = 0.516, p = 0.0238 and r = 0.520, p = 0.0225, respectively). Most facies contain a range 330 331 of both high and low values of CPI, with the highest average CPI in the low-energy marine mudstones 332 of facies 8 (Fig. 6). Facies 6, 7 and 8 also have broad ranges of ACL and $n-C_{29}/n-C_{27}$, with the lowest 333 average ACL and $n-C_{29}/n-C_{27}$ in the poorly sorted sandstones of facies 7, while facies 3, consisting of massive diamictites, has the highest average ACL and $n-C_{29}/n-C_{27}$. Two samples containing a much 334 higher concentration of *n*-alkanes than other samples in the facies skew the averages for facies 6 and 335 7. Without these outliers, facies 8 contains the highest average total abundance of *n*-alkanes. 336

Additional biomarkers were investigated in two samples, with a range of C_{27} - C_{32} hopanes and

338 C_{27} - C_{30} hopenes were present (Supplementary table 2). The dominant compound was 17β , 21β (H)-

bishomohopane (C_{32}) or 17α , 21β (H)-hopane (C_{30}) and samples were characterised by low-to-

340 moderate $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ ratios (0.45 to 0.65). As such, these samples are characterised by relatively

341 low thermal maturity.



Fig. 7. Stratigraphic column from CRP 2/2A with the stratigraphic positions of representative GC-FID-chromatograms of samples. Filled circles above peaks indicate n-alkanes, with the $n-C_{27}$ and $n-C_{29}$ labelled. Simplified facies groupings are labelled, and are described in table 1. UCM:

346 *unresolved complex mixture.*



347

348 Fig. 8. Scatter plots of samples from CRP 2/2A; a) CPI and ACL; b) CPI and $n-C_{29}/n-C_{27}$; c) 349 $n-C_{29}/n-C_{27}$ and ACL; d) CPI and the total abundance of n-alkanes (μ g n-alkanes/g TOC); e) ACL and 350 the total abundance of n-alkanes (μ g n-alkanes/g TOC) and f) $n-C_{29}/n-C_{27}$ and the total abundance of 351 n-alkanes (μ g n-alkanes/g TOC).

352 *3.4 DSDP 270*

Samples from DSDP 270 typically display bimodal *n*-alkane distributions with a peak at $n-C_{17}$ or $n-C_{19}$ and another peak $n-C_{27}$ (Fig. 9). Shorter chained *n*-alkanes (> $n-C_{20}$) are usually more abundant than the long chained homologs, and in four samples from the lowest sampled section of the core, long chained *n*-alkanes were not detected. Some samples display a small UCM underlying ~*n*-C₂₀ and *n*-C₂₁. CPI of the long chained *n*-alkanes ranges from 2.4-4.8 (avg. 3.4). ACL varies from 27.6-28.1 (avg. 27.9), whilst the ratio of $n-C_{29}$ to $n-C_{27}$ ranges from 0.65-0.86 (avg. 0.76). The samples have an average total abundance of *n*-alkanes of 680.92 µg/gTOC (Supplementary table 1).



- 361 Fig. 9. Stratigraphic column from DSDP 270 with the stratigraphic positions of
- 362 representative GC-FID-chromatograms of samples. Filled circles above peaks indicate n-alkanes,
- 363 with the $n-C_{27}$ and $n-C_{29}$ labelled. Simplified facies groupings are labelled, and are described in table
- 364 1. UCM: unresolved complex mixture.
- 365 Pearson's correlation coefficients show no particularly strong correlations between *n*-alkane variables. Increasing $n-C_{29}/n-C_{27}$ with decreasing CPI is very weakly correlated (r = 0.391, p =366 0.0221), with a slightly stronger correlation existing between decreasing $n-C_{29}/n-C_{27}$ with decreasing 367 ACL (r = 0.533, p = 0.0012) (Fig. 10). When grouped by facies, facies 7 demonstrates the highest 368 369 average CPI, and facies 4 contains the lowest average CPI (Fig. 6). Facies 7 shows the highest 370 average ACL and lowest average $n-C_{29}/n-C_{27}$, but most facies display similar ACL and $n-C_{29}/n-C_{27}$ 371 values. Most facies also contain a similar total abundance of *n*-alkanes, with the highest average 372 abundance in facies 3.
- Two samples with representative *n*-alkane distributions were analysed for additional biomarkers (Supplementary table 2). Samples contained a range of C_{27} – C_{32} hopanes and C_{27} – C₃₀ hopenes. The dominant compounds were 17 β (H)-trisnorhopane (C_{27}), 17 α ,21 β (H)-hopane (C30),17 β ,21 β (H)-homohopane (C_{31}). Both samples were characterised by high $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ ratios (0.69 to 1.00) and indicate low thermal maturity.



Fig. 10. Scatter plots of samples from DSDP 270; a) CPI and ACL; b) CPI and n-C₂₉/n-C₂₇;
c) n-C₂₉/n-C₂₇ and ACL; d) CPI and the total abundance of n-alkanes (μg n-alkanes/g TOC); e) ACL

- and the total abundance of n-alkanes (μg n-alkanes/g TOC) and f) n-C₂₉/n-C₂₇ and the total
- 382 abundance of n-alkanes (μg n-alkanes/g TOC).

383 *3.5 DSDP 274*

384 Samples taken from above 115m had variable, or even absent, quantities of *n*-alkanes (Fig. 11). Samples usually contained significant UCMs, which typically dominated the signal, and n-385 alkanes did not display a common dominant *n*-alkane. Samples below 115m were usually bimodal in 386 387 distribution, with a dominant *n*-alkane peak around n-C₁₉, n-C₂₀ or n-C₂₁, underlain by a UCM, and 388 another peak centred at n-C₂₇. CPI ranges from 0.6 to 3.0 (avg. 1.6), whilst ACL ranges from 25 to 389 31.0 (avg. 27.8). The ratio of $n-C_{29}$ to $n-C_{27}$ varies from 0.50 to 1.37 (avg. 0.80). The total abundance of *n*-alkanes averages 72.7 μ g/gTOC, with a range of 2.7 μ g/gTOC to 316.6 μ g/gTOC 390 391 (Supplementary table 1).

392 Pearson's correlation coefficients show the strongest correlations exist between CPI and n-393 $C_{29}/n-C_{27}$ (r = 0.809, p < 0.0000), and ACL and $n-C_{29}/n-C_{27}$ (r = 0.825, p < 0.0000) (Fig. 12). Weak 394 correlations exist between the total abundance of *n*-alkanes and the other three variables considered; 395 CPI, ACL and $n-C_{29}/n-C_{27}$ (r = 0.350, p = 0.0461; r = 0.396, p = 0.0224; r = 0.389, p = 0.0338396 respectively). Only one combination, CPI and ACL, does not indicate a statistically significant 397 correlation, as ACL becomes much more variable at low CPIs. Facies 11 has significantly lower CPI and higher $n-C_{29}/n-C_{27}$ than facies 10 (Fig. 6). ACL is much more variable in facies 11, and on 398 399 average higher, while the total abundance of *n*-alkanes is on average lower than facies 10.

400 One sample from facies 11 was analysed for additional biomarkers (Supplementary table 2). 401 The sample contained a range of thermally mature C_{27} - C_{35} hopanes. The dominant compounds were 402 $17\alpha,21\beta(H)$ -norhopane (C_{29}) and $17\alpha,21\beta(H)$ -hopane (C30). This sample was characterised by low 403 $\beta\beta/(\alpha\beta+\beta\alpha+\beta\beta)$ ratios (0), high C22S/C22R+C22S ratios (0.57) and moderate Ts/Ts+Tm ratio (0.31). 404 Collectively, this indicates high thermal maturity.

406 Fig. 11. Stratigraphic column from DSDP 274 with the stratigraphic positions of

- 407 *representative GC-FID-chromatograms of samples. Filled circles above peaks indicate n-alkanes,*
- 408 with the $n-C_{27}$ and $n-C_{29}$ labelled. Facies are described in table 1. UCM: unresolved complex mixture.

410 Fig. 12. Scatter plots of samples from DSDP 274; a) CPI and ACL; b) CPI and n-C₂₉/n-C₂₇;
411 c) n-C₂₉/n-C₂₇ and ACL; d) CPI and the total abundance of n-alkanes (μg n-alkanes/g TOC); e) ACL

412 and the total abundance of n-alkanes (μg n-alkanes/g TOC) and f) n-C₂₉/n-C₂₇ and the total

414 4 Discussion

415 4.1 Potential sources of lipid biomarkers in Cenozoic Antarctic sediments

416 *4.1.1 Contemporaneous organic matter*

417 Plants and microorganisms living contemporaneously with the accumulation of sediment 418 would have been a major contributing source for *n*-alkanes and hopanes with $\beta\beta$ stereochemistry at 419 the sites studied. Macro- and microfossils from sediment cores, onland outcrops and glacial erratics 420 indicate that the Ross Sea region of Antarctica was vegetated until at least the Mid-Miocene Climate 421 Transition (MMCT) ~14 Ma (e.g. Kemp, 1975; Mildenhall, 1989; Askin, 2000; Askin and Raine, 422 2000; Lewis et al., 2008; Warny et al., 2009; Lewis and Ashworth, 2016). Paleocene and Early Eocene sediment from cores offshore Wilkes Land indicate a highly diverse, near tropical flora 423 occupied coastal regions, with temperate rain forest inland and at higher elevations (Pross et al., 424 425 2012). Following a prolonged period of global cooling, sediments from the Ross Sea region indicate that by the Mid-Late Eocene, vegetation was largely represented by a less diverse, cool, temperate 426 427 flora dominated by Nothofagus-podocarpaceous conifer-proteaceae (Askin, 2000; Francis, 2000; Pole 428 et al., 2000).

⁴¹³ *abundance of n-alkanes (\mu g n-alkanes/g TOC).*

429 The Oligocene and early Miocene was marked by declining vegetation diversity and the 430 development of a sparse, shrubby tundra, dominated by stunted Nothofagus (Kemp, 1975; Kemp and 431 Barrett, 1975; Askin and Raine, 2000; Prebble et al., 2006a). The Mid-Miocene Climate Optimum 432 (MMCO) (~17-15 Ma), saw an increase in the abundance of pollen transported offshore indicating a 433 proliferation of woody vegetation and a possible return to more tree-like forms (Warny et al., 2009; 434 Feakins et al., 2012). This short-lived warming was followed by the MMCT, a major cooling step in 435 the Cenozoic (Shackleton and Kennett, 1975; Flower and Kennett, 1994; Lewis et al. 2007). The vegetation history of Antarctica following the MMCT has been debated, with different schools of 436 thought suggesting that demise of higher plants occurred either at the MMCT (Sugden et al., 1993; 437 Marchant et al., 1996; Lewis et al., 2008) or the Pliocene (Harwood et al. 1983; Webb et al. 1984; 438 Fielding et al., 2012). The Pliocene ages are controversial and rely on sparse diatoms present in tills of 439 440 the Meyer Desert Formation, preserved in the upper Beardmore Glacier (Barrett, 2013).

441 4.1.2 Reworked lipid biomarkers from older sediments

442 Lipid biomarkers could also be sourced from the erosion and redeposition of older sediments. 443 Two main sources are considered here; reworked Cenozoic *n*-alkanes, and reworked Permian-Jurassic 444 biomarkers sourced from the Beacon Super Group and Ferrar Group. The presence of Cretaceous 445 dinoflagellate cysts in samples of the McMurdo erratics suggests the possibility for a contribution 446 from rocks of this age now either eroded or buried, but as these occurrences are extremely rare this 447 potential contribution is considered very minor (Askin, 2000). Surface sediment from the Eastern 448 Ross Sea does contain a significant component of Late Cretaceous palynomorphs, but the location of 449 DSDP Site 270 in the central Ross Sea does not (Truswell and Drewry, 1984). Oligocene/Late 450 Miocene sediments in this core are also barren of pollen from this time period (Kemp, 1975; Duncan, 451 2017).

452 Microfossil work on Oligocene and Miocene sediments in continental margin drillcores frequently indicate the presence of older Cenozoic microfossils, likely eroded from older sedimentary 453 454 basin infill. In particular, Eocene aged dinoflagellate cysts of the 'Transantarctic flora' are used to infer reworking of Eocene material into younger sediments (e.g. Kemp, 1975; Askin and Raine, 2000; 455 Prebble et al., 2006a). Limited burial of Cenozoic sediments means that Paleogene forms reworked 456 457 into younger sediments are still light in colour and display similar autofluorescence (e.g. Askin and 458 Raine, 2000; Prebble et al., 2006a). Here, *n*-alkanes extracted from the fossiliferous McMurdo erratics 459 serve as an indication of typical Mid-Late Eocene distributions of these compounds (Section 4.2.1).

The Beacon Supergroup extends throughout the TAM and is a key source of sediment to the
sedimentary basins of the Ross Sea (e.g. Talarico et al., 2000; Smellie, 2001; Sandroni and Talarico,
2004; Sandroni and Talarico, 2011). Many of the fossil assemblages from the Beacon Supergroup
come from widespread Permian and Triassic sediments, and indicate a cool, humid Mid-Late Permian

464 climate with vegetation dominated by Glossopteris and Gangamopteris (Cúneo et al., 1993; Francis et 465 al., 1994; Collinson, 1997). By the Mid-Triassic, a more diverse flora dominated by Dicroidium 466 indicates a shift to warmer 'greenhouse' conditions (Collinson, 1997; Cúneo et al., 2003). The Beacon 467 Supergroup outcropping in the TAM has undergone widespread intrusion and thermal alteration, with altered palynomorphs in continental margin cores likely reflecting a TAM source, whilst less altered 468 specimens must have been transported from less extensively intruded sediments cratonwards of the 469 470 TAM (Askin, 1998; Askin and Raine, 2000). Fossiliferous sedimentary interbeds of the Jurassic 471 Ferrar group are also known to contribute reworked palynomorphs to offshore sediments, albeit with 472 much rarer occurrences than those sourced from the Beacon Supergroup (Askin and Raine, 2000). n-Alkanes have previously been analysed from Beacon sediments, silicified wood and coal at the Allan 473 474 Hills and Ferrar Group sediments from Carapace Nunatak in Southern Victoria Land (Matsumoto et 475 al., 1986). *n*-Alkanes ranging from $n-C_{12}$ to $n-C_{30}$ displayed a CPI varying from 0.91-1.4. Short chain 476 *n*-alkanes (< n-C₂₀) were typically more abundant than long chain lengths (Matsumoto et al., 1986). A 477 chromatogram from Matsumoto et al. (1986) indicates that UCMs are also present in these samples, 478 centred at $n-C_{18}$ and $n-C_{19}$. Hopanes in these Beacon sediments were typically dominated by $\alpha\beta$ and βα configurations indicating maturation of the sediments and alteration of hopanes from their 479 480 biologically synthesized precursors (Matsumoto et al., 1987). Variable thermal maturation of the 481 Beacon sediments in this region is suggested by two samples containing small quantities of $\beta\beta$ 482 hopanes (Matsumoto et al., 1987)

Distributions of *n*-alkanes, kerogen and palynomorphs in surface and Quaternary sediments 483 484 from the Ross Sea, and soils from the Dry Valleys, suggest the potential for recycling of *n*-alkanes from both Cenozoic and pre-Cenozoic sources is occurring via modern depositional processes. In the 485 486 Ross Sea, n-Alkanes appear in low abundances with short chained n-alkanes attributed to a mixture of 487 primary and recycled material derived from marine organisms, while long chained *n*-alkanes are 488 suggested to be higher plant material either from long-range aeolian transport or reworked from pre-489 Quaternary sediments (Kvenvolden et al., 1987; Venkatesan, 1988). A recycled source for *n*-alkanes 490 is supported by the presence of hopanes of variable maturities ($\beta\beta$, $\beta\alpha$ and $\alpha\beta$), and kerogen and pollen extensively reworked from Paleogene or pre Cenozoic sediment (Sackett et al., 1974; Truswell 491 492 and Drewry, 1984, Kvenvolden et al., 1987). The most abundant *n*-alkanes in soils from the Dry 493 Valleys are usually n-C₂₃, n-C₂₅, or n-C₂₇ (Matsumoto et al., 1990a; Matsumoto et al., 2010; Hart et al., 2011). n-Alkanes are attributed to a mixed source input, predominantly derived from endolithic 494 495 microorganisms and glacially eroded ancient plant and microorganism debris, sourced from earlier 496 Cenozoic sediments and the Beacon Sandstone (Matsumoto et al., 1990a; Matsumoto et al., 2010). 497 This is supported by the presence of mature ($\beta \alpha$ and $\alpha \beta$) isomers of hopanes, likely sourced from Beacon sediments (Matsumoto et al., 1990b) Aeolian transport as a main source for the n-alkanes in 498 499 these samples is considered unlikely, as aerosol samples near Antarctica record *n*-alkane distributions

with high ACLs and a dominant n-C₃₁, potentially as a result of large scale meridional air mass circulation transporting *n*-alkanes from the tropics to high latitudes (Bendle et al., 2007).

502 *4.1.3 In situ degradation of n-alkanes*

503 In Antarctica and the Sub-Antarctic, hydrocarbon contamination experiments indicate that 504 hydrocarbon degrading microbes are present in soils (Aislabie et al., 1998; Bej et al., 2000; Coulon et al., 2005). Longer chain length *n*-alkanes were found to be more resistant to microbial degradation, 505 506 and rates of degradation increase with increasing temperature (Coulon et al., 2005). n-Alkane 507 distributions in the studied samples are not considered the result of *in situ* thermal maturation, or 508 migration of hydrocarbons into the sediments. The sediment sampled in this study comes from near 509 surface sediment in the case of Mt Boreas, or drill cores where the deepest samples come from 385 510 mbsf in DSDP 270. None of the studied cores contained hydrocarbon residues, and heat flow measurements from CRP-2/2A and other cores in the region (CRP 3, ANDRILL 1B, ANDRILL 2A) 511 range from 24-76.7 °C/km (Bücker et al., 2000; Bücker et al., 2001; Morin et al., 2010; Schröder et 512 513 al., 2011). Basin modelling from the central and western Ross Sea shows that while the generation of 514 hydrocarbons is possible in the deeply buried sediments of the basins, expulsion and migration of 515 hydrocarbons from potential source rocks is very unlikely (Strogen and Bland, 2011).

516 *4.2 n-Alkane distributions across sample sites*

517 4.2.1 McMurdo erratics

The McMurdo erratics provide examples of mid-late Eocene *n*-alkane distributions, when 518 Antarctic vegetation was more diverse than in the Oligocene and Miocene, and the climate was 519 warmer and wetter (Askin, 2000; Francis, 2000; Pole et al., 2000). All of the samples except for 520 521 MTD95 contain a terrestrial palynomorph assemblage, with E215 and E219 also including leaves and 522 in E219, wood macrofossils (Harwood and Levy, 2000). However, these erratics do also contain a 523 minor component of reworked material sourced from the Beacon Supergroup (Askin, 2000). Despite 524 this, the occurrence of macro-fossils and dominantly Eocene-aged assemblages of palynomorphs 525 suggest that the *n*-alkanes in these samples are principally from contemporaneously-sourced organic 526 matter. The key difference in the *n*-alkane distributions of the McMurdo erratics compared to Oligocene and Miocene samples from the other studied sites is the prominence of the $n-C_{29}$ as 527 528 opposed to the n-C₂₇ (Supplementary table 1, Fig.3). We suggest the shift from a dominant n-C₂₉ to n-529 C_{27} in the Ross Sea region is due to a combination of climate cooling as the Antarctic ice sheets developed, and a shift in plant community to a flora dominated by a low diversity tundra of 530 531 Nothofagus, podocarpidites and bryophytes (Askin, 2000; Askin and Raine, 2000; Prebble et al., 2006a; Lewis et al., 2008). 532

533 *4.2.2 Mt Boreas*

534 The presence of macro and microfossils of bryophytes at Mt Boreas is represented by the prominence of n-C₂₃ and n-C₂₅ in the n-alkane distributions from this site (Fig. 4) (Lewis et al., 2008). 535 The *n*-C₂₇ homolog is also particularly abundant and is likely sourced from shrubs and trees such as 536 Nothofagidites lachlaniae in the lake catchment (Lewis et al., 2008). A correlation between increasing 537 n-C₂₉ and decreasing CPI could be the result of either: 1) microbial degradation lowering CPI and 538 preferentially degrading the shorter chain n-C₂₇; or 2) incorporation of recycled material, likely from 539 540 weathered, thermally-degraded Beacon Supergroup, and older Cenozoic sediments in the catchment. This is supported by the presence of thermally matured hopanes in sample ALS-05 21O, which is 541 542 from the base of the lacustrine section, just above a glacial till containing clasts of Beacon Supergroup 543 (Lewis et al., 2008). Other samples in which hopanes were investigated were dominated by $\beta\beta$ 544 hopanes, supporting an interpretation that much of the biomarkers in the rest of the lacustrine 545 sediments are contemporaneously sourced. Lacustrine depositional environments (Facies 1) have 546 higher average CPIs and lower average $n-C_{29}/n-C_{27}$ values than fluvial samples (Facies 2) (Fig. 6). Fluvial environments can be erosive settings as coarser sediments require greater water velocity for 547 548 suspension and movement (Miller et al., 1977), suggesting a fluvially influenced environment is more likely to rework *n*-alkanes. In the lacustrine setting, high CPI, low ACL and $n-C_{29}/n-C_{27}$ in particular 549 550 occur directly below, and almost directly above the moss peat (Fig. 4). n-alkane distributions are 551 likely sampling the aquatic plants and mosses deposited during a shallow water phase of the lake. 552 Samples from beds representing a deeper water phase of the lake (Lewis et al., 2008) are marked by a 553 similar average CPI and *n*-C₂₉/*n*-C₂₇ as the laminated silts, but a higher ACL, reflecting an increased input of emergent and terrestrial plant matter from the surrounding catchment. 554

555 *4.2.3 CRP 2/2A*

556 In CRP 2/2A, *n*-alkane distributions typically show the n-C₂₇ as the dominant homolog, although n-C23, n-C25 and n-C29 were also commonly abundant. The prominence of these n-alkane 557 homologs is in line with palynomorph evidence which suggests input from trees, shrubs and 558 bryophytes (Prebble et al., 2006a). Fluctuating abundances of reworked palynomorphs (thermally 559 altered, poorly preserved or of a known older range) often coincide with larger abundances of Eocene 560 561 Transantarctic flora dinoflagellates (Prebble et al., 2006a). This indicates reworked samples were 562 sourced from both Permian/Triassic Beacon sediments and earlier Cenozoic sediments. The correlations between low CPI, and high ACL and $n-C_{29}/n-C_{27}$ (Fig. 8) can be explained by a mixed 563 source input of n-alkanes, from contemporaneous material, early Cenozoic sediments, and both 564 altered and unaltered areas of the Beacon Supergroup. This is supported by the presence of both 565 biologically synthesized and thermally matured hopane configurations. While a contribution from 566 567 more recent recycled material (i.e Early Oligocene n-alkanes) cannot be ruled out, Prebble et al.

- 568 (2006b) found little evidence for reworking between Early and Late Oligocene sequences. UCMs in
- these samples typically underlie the lower chain lengths, and may be the result of post-depositional
- 570 microbial alteration, or could be inherited from the Beacon Supergroup (Matsumoto et al., 1986).
- 571 Facies groupings reflect depositional environments which are predominantly influenced by the
- proximity of glaciers near the site. More ice-distal, marine facies (7 and 8) have on average high CPIs,
- 573 low ACL and low $n-C_{29}/n-C_{27}$ (Fig. 6), while samples from ice-proximal or subglacial settings tend to
- show the opposite trends. This suggests that low-energy, more ice-distal marine environments are
- 575 more likely to contain well-preserved *n*-alkane distributions reflecting contemporaneously sourced *n*-
- alkanes, whilst more ice-proximal and subglacial environments have a higher likelihood of containingreworked *n*-alkanes.

578 *4.2.4 DSDP 270*

n-Alkane distributions from DSDP 270 are typically bi-modal suggesting two primary sources 579 for *n*-alkanes in this drill core (Fig. 9). Algae and bacteria the likely source for the shorter chain 580 581 lengths, with terrestrial higher plants contributing to longer chain lengths (section 2.5). The presence 582 of a contemporaneous pollen assemblage with almost no reworked contribution indicates the long 583 chained *n*-alkanes predominantly reflect contemporary onshore vegetation. This is shown in the CPI values that vary less than the other sites sampled and all sit above 2.4 (Fig. 6), and the predominance 584 585 of hopanes in $\beta\beta$ configurations. Facies representing more ice-proximal settings (facies 3 and 4) show the lowest average CPIs suggesting that these settings are likely to contain more degraded *n*-alkane 586 587 distributions, whether as the result of post-depositional processes or some sediment recycling due to 588 glacial erosion and redeposition (Fig. 6).

589 *4.2.5 DSDP 274*

590 *n*-Alkane distributions in DSDP 274 are separated into two distinct groups, above and below an unconformity/condensed section at 113.6 mbsf (Figs. 6 and 11). Samples taken from below 113.6 591 592 mbsf show bi-modal distributions in chromatograms suggesting a mixed contribution from both algae 593 and bacteria, and terrestrial plants. Other than the uppermost 2 samples from this section of the core, all samples are considered to be part of facies 10, which was deposited with a high terrigenous and 594 595 biogenic sedimentation rate, under a regime of weak bottom currents (Fig. 2) (Frakes, 1975; 596 Whittaker and Müller, 2006). Reworked palynomorphs are present in this section (Kemp, 1975), 597 which, coupled with variable CPI and UCMs suggest that some contribution of reworked *n*-alkanes is 598 likely. However, the generally high CPI, dominance of the n-C₂₇ and lack of variation in ACL and n- C_{29}/n - C_{27} suggests that much of the *n*-alkanes present reflect comparatively more contemporaneous 599 600 input than the overlying interval, or at least material recycled from the Oligocene or younger.

601 Above the unconformity or condensed section at 113.6 mbsf, the sedimentation rate slows and 602 manganese nodules provide evidence for winnowing by a strong bottom current regime. This interval 603 is also associated with an increase in coarse sediment which could result from ice rafting (Frakes, 604 1975; Whittaker and Müller, 2006) or winnowing of the fine fraction due to intensification of bottom 605 currents. These sediments date to the Late Miocene, and Antarctic glacial expansion at this time could 606 explain the increase in ice rafting or bottom water current intensity (McKay et al., 2009; Herbert et al., 607 2016). Sediments from this part of the core are also include and post-date the MMCT when it has been debated that higher plants became extinct on Antarctica (Sugden et al., 1993; Marchant et al., 608 609 1996; Lewis et al., 2008). This indicates that *n*-alkanes from this section of the core may 610 predominantly be derived from older sediments, an interpretation supported by a hopane distribution 611 dominated by thermally matured configurations. Low CPIs, high $n-C_{29}/n-C_{27}$, often large and dominant UCMs and low sedimentation rates suggest that *n*-alkanes in these samples have also been 612 613 extensively degraded, likely by microbial activity as sediments are winnowed and reworked in the 614 surface layers of the seabed.

615 **5.** Synthesis

n-Alkane distributions in Eocene to Miocene sediments from the Ross Sea region vary with 616 age and sample site. Between the Eocene and Oligocene, the dominant chain length recorded in 617 618 sediments changes from $n-C_{29}$ to $n-C_{27}$, concomitant with a significant climate cooling and a shift in plant community (section 4.2.1). The dominance of the n-C₂₇ in sediments sourced from wide 619 620 catchments incorporating a cool, low diversity vegetation dominated by Nothofagus is in contrast to 621 lower latitudes where $n-C_{29}$ and $n-C_{31}$ are often more abundant (e.g. Poynter et al., 1989; Kawamura 622 et al., 2003; Sachse et al., 2006; Bendle et al., 2007). At least one modern species of Nothofagus (N. 623 menziesii) from New Zealand has been shown to produce n-C₂₇ as its dominant n-alkane (Burrington, 624 2015), while other species in New Zealand and South America are typically dominated by *n*-C₂₉ and 625 n-C₃₁ (Schellekens et al., 2009; Schellekens et al., 2011; Burrington, 2015). The high abundance of n-C27 in samples from Mt Boreas also containing abundant pollen from N. lachlaniae suggest that this 626 species was likely producing large proportions of this *n*-alkane. The prominence of the $n-C_{27}$ across 627 the Oligocene and Miocene sites of this study likely reflects both a climate adaption by plants 628 growing in the Antarctic tundra to cold temperatures, and the abundance of Nothofagus in the 629 630 catchments.

631 While the Oligocene and Miocene vegetation of Antarctica was a main source of *n*-alkanes to 632 the sample sites, reworked *n*-alkanes and hopanes from early and pre-Cenozoic sediments were also 633 evident. In particular, variables that often characterised samples with more reworked material were 634 low CPI values, but higher ACLs and $n-C_{29}/n-C_{27}$ ratios. In a sample containing material solely from 635 reworked thermally altered sections of the Beacon Supergroup, a low CPI, ACL and $n-C_{29}/n-C_{27}$

- 636 would be expected (Matsumoto et al., 1986). The association of low CPIs with high ACL and $n-C_{29}/n-$
- C_{27} therefore suggests that reworked samples likely contain a mixture of *n*-alkanes derived from
- 638 thermally matured Beacon sediments, coupled with material from early Cenozoic and less altered pre-
- 639 Cenozoic sediments containing a higher abundance of longer chained n-alkanes such as n-C₂₉ and n-
- C_{31} . In some instances, this distribution could also result from microbial degradation, which could
- 641 lower CPI, whilst also preferentially scavenging shorter chain lengths.

642 Sediments deposited by glacio-fluvial, ice-proximal glaciomarine and subglacial processes 643 are more likely to contain reworked *n*-alkane distributions than those from lacustrine or ice-distal 644 marine environments, although careful site specific consideration of sediment provenance must be 645 undertaken, regardless of relative proximity to glaciers or rivers. Prior to the MMCT, glaciers 646 throughout the TAM, and likely the exposed areas of the Ross Sea, were warm-based (Marchant and 647 Denton, 1996; Lewis et al., 2007). This regime would have favoured high rates of glacial erosion of 648 underlying strata, resulting in rapid remobilisation, deposition and burial of sediment in glacial 649 proximal regions (Sugden and Denton, 2004; Powell et al. 2000). These processes likely led to the 650 deposition of greater proportions of reworked *n*-alkanes in ice-proximal environments. Although ice-651 distal settings also record glaciomarine processes and may be subject to reworking, they are likely a 652 more integrated record of aeolian and glacio-fluvial sediment transport offshore. However, as 653 Antarctica became progressively more arid during the Late Miocene and Pliocene, it is feasible that 654 offshore transport of contemporaneous *n*-alkanes via glacio-fluvial action reduced, and thus the 655 relative input of reworked *n*-alkanes became more prominent (e.g. at DSDP 274).

The varying contribution of contemporaneous and reworked biomarkers across sediments 656 657 sourced from different depositional environments, catchments and ages emphasizes how caution must be exercised when applying biomarker-based paleoclimate proxies in glacially-influenced settings 658 (e.g. *n*-alkane δ^{13} C and δ^{2} H). In particular, several aspects should be considered when determining if 659 an *n*-alkane distribution is a contemporaneously-sourced organic matter signature. These include the 660 values and variation of factors such as CPI and ACL, maturation indices of other biomarkers such as 661 662 hopanes, whether the catchment and depositional setting of a site is more likely to accumulate and preserve a contemporary distribution, and assemblages of other fossil material such as palynomorphs. 663 When constructing timeseries of biomarker assemblages, it is also important to consider other aspects 664 665 of the depositional environment in the Ross Sea. The coastal setting of the Ross Sea could be influenced by pulses of reworked material, given the potential for point source glacial meltwater 666 667 discharge, and large-scale meltwater discharge events (i.e. Powell and Domack, 2002; Lewis et al., 2006), which may focus erosion to a certain lithological source. Input of reworked material via 668 669 episodic, erosive hydrological events in Paleocene-Eocene Thermal Maximum sediments from Tanzania has been invoked to explain the highly variable *n*-alkane δ^{13} C values in these sediments 670 (Carmichael et al., 2017). Glacially-influenced environments have a high potential to erode and 671

almost instantaneously redeposit older biomarkers and pollen offshore in concentrated numbers, and

673 indeed biomarker distributions could be used as a potential tool to identify such reworking events.

674 6 Conclusions

675 n-Alkane and hopanoid distributions have been characterised in Eocene to Miocene sediments from a range of depositional environments in the Ross Sea region of Antarctica. Between the Late 676 Eocene and the Oligocene, a shift in *n*-alkane dominant chain length is observed from $n-C_{29}$ to $n-C_{27}$. 677 678 This is inferred to be a result of both a shift in plant community, as well as a response to significant 679 climate cooling. Biomarker distributions in Oligocene and Miocene samples varyingly display a 680 contribution from both contemporaneous and reworked sources. n-Alkane distributions typical of a 681 reworked sample were a low CPI, and high ACL and $n-C_{29}/n-C_{27}$ values. Reworked samples likely reflect a mixed contribution from thermally altered and less thermally altered regions of the Mesozoic 682 683 Beacon Super Group, coupled with material sourced from earlier Cenozoic sediments. Microbial degradation during transport and post-deposition may also contribute to these distributions. Samples 684 685 dominated by contemporaneously-sourced organic matter display a higher CPI, and lower ACL and n- C_{29}/n - C_{27} values. These *n*-alkanes were sourced from the sparse, cold tundra which existed during this 686 687 time. Fluvial environments onshore, and subglacial and ice-proximal environments offshore were 688 more likely to contain reworked *n*-alkanes. Lacustrine environments onshore, and ice-distal 689 environments offshore, were more likely to contain contemporary *n*-alkanes. These findings indicate 690 the possibility of reworking should be taken into account when biomarkers are used for paleoclimate 691 studies in ice-marginal environments.

692 7 Acknowledgments

693 The authors are grateful for access to samples from the IODP core repository at Texas A&M 694 University for DSDP Sites 270 and 274. This study was funded via an Antarctica New Zealand Sir Robin Irvine PhD Scholarship and Scientific Committee of Antarctic Research Fellowship awarded to 695 696 Bella Duncan, with additional funding from a Royal Society of New Zealand Rutherford Discovery 697 Fellowship awarded to Rob McKay (RDF-13-VUW-003) and New Zealand Ministry of Business Innovation and Employment Contract C05X1001. Field activities were supported by Antarctica New 698 699 Zealand. The authors are grateful for support from IODP and support in kind from the University of 700 Birmingham. The authors thank two anonymous reviewers for their constructive comments.

701 Data Availability

702

Data associated with this study can be found in the supplementary tables.

703 **References**

704 Aislabie, J., McLeod, M., Fraser, R., 1998. Potential for biodegradation of hydrocarbons in soil from 705 the Ross Dependency, Antarctica. Applied Microbiology and Biotechnology 49, 210-214. 706 Allibone, A.H., Cox, S.C., Graham, I.J., Smillie, R.W., Johnstone, R.D., Ellery, S.G., Palmer, K., 707 1993. Granitoids of the Dry Valleys area, southern Victoria Land, Antarctica: plutons, field 708 relationships, and isotopic dating. New Zealand journal of geology and geophysics 36, 281-297. 709 Allibone, A.H., Cox, S.C., Smillie, R.W., 1993. Granitoids of the Dry Valleys area, southern Victoria 710 Land: geochemistry and evolution along the early Paleozoic Antarctic Craton margin. New 711 Zealand journal of geology and geophysics 36, 299-316. 712 Askin, R.A., 1998. Palynological investigations of Mount Feather Sirius Group samples: recycled 713 Triassic assemblages, In: Wilson, G.S., Barron, J. (Eds.), Mount Feather Sirius Group Core Workshop and Collaborative Sample Analysis. Byrd Polar Research Center Report No. 14. 714 Byrd Polar Research Center, The Ohio State University, Columbus, Ohio, pp. 59-65. 715 716 Askin, R.A., 2000. Spores and pollen from the McMurdo Sound erratics, Antarctica. In: Stilwel, J.D., 717 Feldman, R.M. (Eds.), Paleobiology and Paleoenvironments of Eocene Rocks: McMurdo Sound, East Antarctica. American Geophysical Union, Washington, D.C., pp. 161-181. 718 719 Askin, R.A., Raine, J.I., 2000. Oligocene and Early Miocene terrestrial palynology of the Cape 720 Roberts Drillhole CRP-2/2A, Victoria Land Basin, Antarctica. Terra Antarctica 7, 493-501. 721 Barrett, P.J., 1981. History of the Ross Sea region during the deposition of the Beacon Supergroup 722 400-180 million years ago. Journal of the Royal Society of New Zealand 11, 447-458. 723 Barrett, P.J., Elliot, D.H., Lindsay, J.F., 1986. The Beacon Supergroup (Devonian- Triassic) and 724 Ferrar Group (Jurrasic) in the Beardmore Glacier Area, Antarctica. In: Turner, M.D., Splettstoesser J.E. (Eds.), Geology of the central Transantarctic Mountains. American 725 Geophysical Union, Washington, D.C., pp. 339-428. 726 727 Barrett, P.J. (Ed.). 1989. Antarctic Cenozoic history from the CIROS-1 drillhole, McMurdo Sound. 728 DSIR Publishing, Wellington, New Zealand. 729 Barrett, P.J., 2013. Resolving views on Antarctic Neogene glacial history-the Sirius debate. Earth and 730 Environmental Science Transactions of the Royal Society of Edinburgh 104, 31-53. 731 Bej, A.K., Saul, D., Aislabie, J., 2000. Cold-tolerant alkane-degrading Rhodococcus species from 732 Antarctica. Polar Biology 23, 100-105. 733 Bendle, J., Kawamura, K., Yamazaki, K., Niwai, T., 2007. Latitudinal distribution of terrestrial lipid 734 biomarkers and n-alkane compound-specific stable carbon isotope ratios in the atmosphere over 735 the western Pacific and Southern Ocean. Geochimica et Cosmochimica Acta 71(24), 5934-736 5955. 737 Bijl, P.K., Bendle, J.A., Bohaty, S.M., Pross, J., Schouten, S., Tauxe, L. Stickley, C.E., McKay, R.M., 738 Röhl, U., Olney, M., Sluijs, A., Escutia, C., Brinkhuis, H., Expedition 318 Scientists, 2013.

- 739Eocene cooling linked to early flow across the Tasmanian Gateway. Proceedings of the
- 740 National Academy of Sciences 110, 9645-9650.
- Bray, E.E., Evans, E.D., 1961. Distribution of n-paraffins as a clue to recognition of source
 beds. Geochimica et Cosmochimica Acta 22, 2-15.
- Bücker, C.J., Wonik, T., Jarrard, R., 2000. The temperature and salinity profile in CRP-2/2A, Victoria
 Land Basin, Antarctica. Terra Antarctica 7, 255-259.
- Bücker, C., Jarrard, R.D., Wonik, T., 2001. Downhole temperature, radiogenic heat production, and
 heat flow from the CRP-3 drillhole, Victoria Land Basin, Antarctica. Terra Antarctica 8, 151160.
- Burrington, P., 2015. How to be a Prehistoric Weatherman: Using n-alkanes as a Proxy for Holocene
 Climate and Hydrology, Southwest South Island, New Zealand (Unpublished Masters Thesis).
 University of Otago, New Zealand.
- Bush, R.T., McInerney, F.A., 2013. Leaf wax n-alkane distributions in and across modern plants:
 implications for paleoecology and chemotaxonomy. Geochimica et Cosmochimica Acta 117, 161-179.
- Bush, R.T., McInerney, F.A., 2015. Influence of temperature and C₄ abundance on n-alkane chain
 length distributions across the central USA. Organic Geochemistry 79, 65-73.
- Calvo, E., Pelejero, C., Logan, G.A., De Deckker, P., 2004. Dust-induced changes in phytoplankton
 composition in the Tasman Sea during the last four glacial cycles. Paleoceanography 19.
- Cape Roberts Science Team, 1999. Studies from the Cape Roberts Project, Ross Sea Antarctica,
 Initial report on CRP-2/2A. Terra Antarctica 6(1), 1-173.
- 760 Carmichael, M. J., Inglis, G. N., Badger, M. P., Naafs, B. D. A., Behrooz, L., Remmelzwaal, S.,
- 761 Monteiro, F. M., Rohrssen, M., Farnsworth, A., Buss, H. L., Dickson, A. J., Valdes, P. J., Lunt,
- D. J., Pancost, R. D., 2017. Hydrological and associated biogeochemical consequences of rapid
 global warming during the Paleocene-Eocene Thermal Maximum. Global and Planetary
 Change 157, 114-138.
- Clark Jr, R.C., Blumer, M., 1967. Distribution of n-paraffins in marine organisms and
 sediment. Limnology and Oceanography 12, 79-87.
- Collinson, J.W., 1997. Paleoclimate of Permo-Triassic Antarctica. International Symposium on
 Antarctic Earth Sciences 7, 1029-1034.
- Cooper, A.K., Davey, F.J., Behrendt, J.C., 1987. Seismic stratigraphy and structure of the Victoria
 Land basin, western Ross Sea, Antarctica. In: Cooper, A.K., Davey, F.J. (Eds.), The Antarctic
 Continental Margin: Geology and Geophysics of the Western Ross Sea. Circumpacific Council
 for Energy and Mineral Resources, Houston, T.X., pp. 27-65.
- Coulon, F., Pelletier, E., Gourhant, L., Delille, D., 2005. Effects of nutrient and temperature on
 degradation of petroleum hydrocarbons in contaminated sub-Antarctic soil. Chemosphere 58,
 1439-1448.

- Cranwell, P.A., Eglinton, G., Robinson, N., 1987. Lipids of aquatic organisms as potential
 contributors to lacustrine sediments—II. Organic Geochemistry 11, 513-527.
- Cúneo, N.R., Isbell, J., Taylor, E.L., Taylor, T.N., 1993. The Glossopteris flora from Antarctica:
 taphonomy and paleoecology. Comptes Rendus XII ICC-P 2, 13-40.
- 780 Cúneo, N.R., Taylor, E.L., Taylor, T.N., Krings, M., 2003. In situ fossil forest from the upper
- Fremouw Formation (Triassic) of Antarctica: paleoenvironmental setting and paleoclimate
 analysis. Palaeogeography, Palaeoclimatology, Palaeoecology 197, 239-261.
- 783 Decesari, R.C., Sorlien, C.C., Luyendyk, B.P., Wilson, D.S., Bartek, L.R., Diebold, J., Hopkins, S.E.,
- 784 2007. Regional seismic stratigraphic correlations of the Ross Sea: implications for the tectonic
- 785 history of the West Antarctic Rift System. In: Cooper, A.K., Raymond, C.R., 10th ISAES
- 786Editorial Team (Eds.), Antarctica: A Keystone in a Changing World- Online Proceedings of the
- 787 10th ISAES, 2007-1047, USGS Open-File Report, Short Research Paper 052, 4p.
- 788 De Santis, L., Anderson, J.B., Brancolini, G., Zayatz, I., 1995. Seismic record of late Oligocene
- through Miocene glaciation on the central and eastern continental shelf of the Ross Sea. In:
- Cooper, A.K., Barker, P.F., Brancolini, G. (Eds.), Geology and Seismic Stratigraphy of the
 Antarctic Margin. American Geophysical Union, Washington, D.C., pp. 235-260.
- Dodd, R.S., Rafii, Z.A., Power, A.B., 1998. Ecotypic adaptation in Austrocedrus chilensis in cuticular
 hydrocarbon composition. New Phytologist 138, 699-708.
- Dodd, R.S., Afzal-Rafii, Z., 2000. Habitat-related adaptive properties of plant cuticular
 lipids. Evolution 54, 1438-1444.
- Duncan, B.J., 2017. Cenozoic Antarctic climate evolution based on molecular and isotopic biomarker
 reconstructions from geological archives in the Ross Sea region (Unpublished PhD Thesis).
 Victoria University of Wellington, New Zealand.
- Eglinton, G., Hamilton, R.J., 1963. The distribution of alkanes. Chemical plant taxonomy 187, 217.
- Farrimond, P., Taylor, A., TelnÆs, N., 1998. Biomarker maturity parameters: the role of generation
 and thermal degradation. Organic Geochemistry 29, 1181-1197.
- Feakins, S.J., Warny, S., Lee, J.E., 2012. Hydrologic cycling over Antarctica during the middle
 Miocene warming. Nature Geoscience 5, 557-560.
- Feakins, S.J., Warny, S., DeConto, R.M., 2014. Snapshot of cooling and drying before onset of
 Antarctic Glaciation. Earth and Planetary Science Letters 404, 154-166.
- Feakins, S.J., Peters, T., Wu, M.S., Shenkin, A., Salinas, N., Girardin, C.A., Bentley, L.P., Blonder,
 B., Enquist, B.J., Martin, R.E., Asner, G.P., Asner, G.P., 2016. Production of leaf wax n-
- alkanes across a tropical forest elevation transect. Organic Geochemistry 100, 89-100.
- Ficken, K.J., Li, B., Swain, D.L., Eglinton, G., 2000. An n-alkane proxy for the sedimentary input of
 submerged/floating freshwater aquatic macrophytes. Organic Geochemistry 31, 745-749.
- Fielding, C.R., Naish, T.R., Woolfe, K., Lavelle, M., 2000. Facies analysis and sequence stratigraphy
 of CRP-2/2A, Victoria Land Basin, Antarctica. Terra Antarctica 7, 323-338.

- Fielding, C.R., Henrys, S.A., Wilson, T.J., 2006. Rift history of the western Victoria Land Basin: a
- new perspective based on integration of cores with seismic reflection data. In D.K. Fütterer, D.
 Damaske, G. Kleinschmidt, H. Miller & F. Tessensohn (Eds.), Antarctica (pp. 309-318). Berlin
 Heidelberg: Springer.
- Fielding, C.R., Harwood, D.M., Winter, D.M., Francis, J.E., 2012. Neogene stratigraphy of Taylor
- 818 Valley, Transantarctic Mountains, Antarctica: evidence for climate dynamism and a vegetated
 819 early Pliocene coastline of McMurdo Sound. Global and Planetary Change 96-97, 97-104.
- Fitzgerald, P.G., 1994. Thermochronologic constraints on post-Paleozoic tectonic evolution of the
 central Transantarctic Mountains, Antarctica. Tectonics 13, 818-836.
- Flower, B.P., Kennett, J.P., 1994. The middle Miocene climatic transition: East Antarctic ice sheet
 development, deep ocean circulation and global carbon cycling. Palaeogeography,
 palaeoclimatology, palaeoecology 108, 537-555.
- Frakes, L.A., 1975. Paleoclimatic significance of some sedimentary components at Site 274. Initial
 Reports of the Deep Sea Drilling Project 28, 785-787.
- Francis, J.E., Woolfe, K.J., Arnott, M.J., Barrett, P.J., 1994. Permian climates of the southern margins
 of Pangea: evidence from fossil wood in Antarctica. Pangea: Global Environments and
 Resources- Memoir 17, 275-282.
- Francis, J.E., Hill, R.S., 1996. Fossil plants from the Pliocene Sirius Group, Transantarctic Mountains:
 Evidence for climate from growth rings and fossil leaves. Palaios 11, 389-396.
- Francis, J.E., 2000. Fossil Wood from Eocene High Latitude Forests: Mcmurdo Sound, Antarctica. In:
 Stilwel, J.D., Feldman, R.M. (Eds.), Paleobiology and Paleoenvironments of Eocene Rocks:
- McMurdo Sound, East Antarctica. American Geophysical Union, Washington, D.C., pp. 253260.
- Gagosian, R.B., Peltzer, E.T., 1986. The importance of atmospheric input of terrestrial organic
 material to deep sea sediments. Organic Geochemistry 10, 661-669.
- Goodge, J.W., Myrow, P., Williams, I.S., Bowring, S.A., 2002. Age and provenance of the
 Beardmore Group, Antarctica: constraints on Rodinia supercontinent breakup. The Journal of
 geology 110, 393-406.
- Gough, M.A., Rowland, S.J., 1990. Characterization of unresolved complex mixtures of hydrocarbons
 in petroleum. Nature 344, 648-650.
- Gough, M.A., Rhead, M.M., Rowland, S.J., 1992. Biodegradation studies of unresolved complex
 mixtures of hydrocarbons: model UCM hydrocarbons and the aliphatic UCM. Organic
 Geochemistry 18, 17-22.
- Grimalt, J., Albaigés, J., Al-Saad, H.T., Douabul, A.A.Z., 1985. n-Alkane distributions in surface
 sediments from the Arabian Gulf. Naturwissenschaften 72, 35-37.

- 67 Grimalt, J., Albaigés, J., 1987. Sources and occurrence of C_{12} - C_{22} n-alkane distributions with even carbon-number preference in sedimentary environments. Geochimica et Cosmochimica Acta 51, 1379-1384.
- Hambrey, M.J., Barrett, P.J. 1993. Cenozoic sedimentary and climatic record, Ross Sea region,

Antarctica. In: Kennett, J.P., Warnke, D.A. (Eds.) The Antarctic Paleoenvironment: A

- Perspective on Global Change: Part Two, American Geophysical Union, Washington, D.C., pp.91-124.
- Han, J., Calvin, M., 1969. Hydrocarbon distribution of algae and bacteria, and microbiological
 activity in sediments. Proceedings of the National Academy of Sciences 64, 436-443.
- Hart, K.M., Szpak, M.T., Mahaney, W.C., Dohm, J.M., Jordan, S.F., Frazer, A.R., Allen, C.C.R.,
 Kelleher, B.P., 2011. A bacterial enrichment study and overview of the extractable lipids from
 paleosols in the Dry Valleys, Antarctica: implications for future Mars
- reconnaissance. Astrobiology 11, 303-321.
- Harwood, D.M., 1983. Diatoms from the Sirius Formation, Transantarctic Mountains. Antarctic
 Journal of the United States 18, 98-100.
- Harwood, D.M., Levy, R.H., 2000. The McMurdo Erratics: introduction and overview. In: Stilwel,
 J.D., Feldman, R.M. (Eds.), Paleobiology and Paleoenvironments of Eocene Rocks: McMurdo
 Sound, East Antarctica, American Geophysical Union, Washington, D.C., pp. 1-18.
- 866 Herbert, T.D., Lawrence, K.T., Tzanova, A., Peterson, L.C., Caballero-Gill, R., Kelly, C.S., 2016.
- Late Miocene global cooling and the rise of modern ecosystems. Nature Geoscience 9, 843-868 847.
- Inglis, G. N., Naafs, B. D. A., Zheng, Y., McClymont, E. L., Evershed, R. P., Pancost, R. D., 2018.
 Distributions of geohopanoids in peat: Implications for the use of hopanoid-based proxies in
 natural archives. Geochimica et Cosmochimica Acta 224, 249-261.
- Kawamura, K., Ishimura, Y., Yamazaki, K., 2003. Four years' observations of terrestrial lipid class
 compounds in marine aerosols from the western North Pacific. Global Biogeochemical
 Cycles 17, 3-1-3-19.
- Kemp, E.M., 1975. Palynology of Leg 28 drill sites, Deep Sea Drilling Project. Initial Reports of the
 Deep Sea Drilling Project 28, 599-623.
- Kemp, E.M., Barrett, P. J., 1975. Antarctic glaciation and early Tertiary vegetation. Nature 258, 507508.
- Kraus, C., 2016. Oligocene to early Miocene glacimarine sedimentation of the central Ross Sea, and
 implications for the evolution of the West Antarctic Ice Sheet (Unpublished Masters Thesis).
 Victoria University of Wellington, New Zealand.
- Kvenvolden, K.A., Rapp, J.B., Golan-Bac, M., Hostettler, F.D., 1987. Multiple sources of alkanes in
 Quaternary oceanic sediment of Antarctica. Organic geochemistry 11, 291-302.

- Lehtonen, K., Ketola, M., 1993. Solvent-extractable lipids of Sphagnum, Carex, Bryales and CarexBryales peats: content and compositional features vs peat humification. Organic
 Geochemistry 20, 363-380.
- Levy, R.H., Harwood, D.M., 2000. Tertiary marine palynomorphs from the McMurdo Sound erratics,
 Antarctica. In Stilwel, J.D., Feldman, R.M. (Eds.), Paleobiology and Paleoenvironments of
 Eocene Rocks: McMurdo Sound, East Antarctica, American Geophysical Union, Washington,
 D.C., pp. 183-242.
- Levy, R., Harwood, D., Florindo, F., Sangiorgi, F., Tripati, R., von Eynatten, H., Gasson, E., Kuhn.
 G., Tripati, A., DeConto, R., Fielding, C., Field, B., Golledge, N., McKay, R., Naish, T., Olney,
 M., Pollard, D., Schouten, S., Talarico, F., Warny, S., Willmott, V., Acton, G., Panter, K.,
 Paulsen, T., Taviani, M., SMS Science Team, 2016. Antarctic ice sheet sensitivity to
 atmospheric CO₂ variations in the early to mid-Miocene. Proceedings of the National Academy
 of Sciences 113, 3453-3458.
- Lewis, A.R., Marchant, D.R., Kowalewski, D.E., Baldwin, S.L., Webb, L.E., 2006. The age and
 origin of the Labyrinth, western Dry Valleys, Antarctica: Evidence for extensive middle
 Miocene subglacial floods and freshwater discharge to the Southern Ocean. Geology 34, 513516.
- Lewis, A.R., Marchant, D.R., Ashworth, A.C., Hemming, S.R., Machlus, M.L., 2007. Major middle
 Miocene global climate change: Evidence from East Antarctica and the Transantarctic
 Mountains. Geological Society of America Bulletin 119, 1449-1461.
- Lewis, A.R., Marchant, D.R., Ashworth, A.C., Hedenäs, L., Hemming, S.R, Johnson, J.V., Leng,
 M.J., Machlus, M.L., Newton, A.E., Raine, J.I., Willenbring, J.K., Williams, M., Wolfe, A.P.,
 2008. Mid-Miocene cooling and the extinction of tundra in continental Antarctica. Proceedings
- 907 of the National Academy of Sciences 105, 10676-10680.
- Lewis, A.R., Ashworth, A.C., 2016. An early to middle Miocene record of ice-sheet and landscape
 evolution from the Friis Hills, Antarctica. Geological Society of America Bulletin 128, 719738.
- Luyendyk, B.P., Sorlien, C.C., Wilson, D.S., Bartek, L.R., Siddoway, C.S., 2001. Structural and
 tectonic evolution of the Ross Sea rift in the Cape Colbeck region, Eastern Ross Sea,
- 913 Antarctica. Tectonics 20, 933-958.
- Madureira, L.A., Piccinini, A., 1999. Lipids as indicators of paleoclimatic changes, II: terrestrial
 biomarkers. Revista Brasileira de Oceanografia 47, 115-125.
- 916 Marchant, D.R., Denton, G.H., 1996. Miocene and Pliocene paleoclimate of the Dry Valleys region,
 917 southern Victoria Land: a geomorphological approach. Marine Micropaleontology 27, 253-271.
- 918 Marchant, D.R., Denton, G.H., Swisher, C.C., Potter, N., 1996. Late Cenozoic Antarctic paleoclimate
- 919 reconstructed from volcanic ashes in the Dry Valleys region of southern Victoria
- 20 Land. Geological Society of America Bulletin 108, 181-194.

- 921 Martin, A.P., Cooper, A.F., Dunlap, W.J., 2010. Geochronology of Mount Morning, Antarctica: two-922 phase evolution of a long-lived trachyte-basanite-phonolite eruptive center. Bulletin of 923 Volcanology 72, 357-371.
- 924 Matsumoto, G.I., Funaki, M., Machihara, T., Watanuki, K., 1986. Alkanes and alkanoic acids in the 925 Beacon Supergroup samples from the Allan Hills and the Carapace Nunatak in
- 926 Antarctica. Memoirs of National Institute of Polar Research. Special issue 43, 149-158.
- 927 Matsumoto, G. I., Machihara, T., Suzuki, N., Funaki, M., Watanuki, K., 1987. Steranes and
- triterpanes in the Beacon Supergroup samples from southern Victoria Land in Antarctica. 928 929 Geochimica et Cosmochimica Acta 51, 2663-2671.
- Matsumoto, G.I., Akiyama, M., Watanuki, K., Torii, T., 1990a. Unusual distributions of long-chain n-930 931 alkanes and n-alkenes in Antarctic soil. Organic Geochemistry 15, 403-412.
- 932 Matsumoto, G. I., Hirai, A., Hirota, K., Watanuki, K., 1990b. Organic geochemistry of the McMurdo 933 dry valleys soil, Antarctica. Organic Geochemistry 16, 781-791.
- 934 Matsumoto, G.I., Honda, E., Sonoda, K., Yamamoto, S., Takemura, T., 2010. Geochemical features 935 and sources of hydrocarbons and fatty acids in soils from the McMurdo Dry Valleys in the 936 Antarctic. Polar Science 4, 187-196.
- 937 McKay, R., Browne, G., Carter, L., Cowan, E., Dunbar, G., Krissek, L., Naish, T., Powell, R., Reed, 938 J., Talarico, F., Wilch, T., 2009. The stratigraphic signature of the late Cenozoic Antarctic Ice 939
- Sheets in the Ross Embayment. Geological Society of America Bulletin 121, 1537-1561.
- 940 McKay, R., Naish, T., Carter, L., Riesselman, C., Dunbar, R., Sjunneskog, C., Winter, D., Sangiorgi,
- 941 F., Warren, C., Pagini, M., Schouten, S., Willmott, V., Levy, R., DeConto R., Powell, R. D.,
- 942 2012. Antarctic and Southern Ocean influences on Late Pliocene global cooling. Proceedings of the National Academy of Sciences 109, 6423-6428. 943
- 944 Mackenzie, A. S., Patience, R. L., Maxwell, J. R., Vandenbroucke, M., Durand, B., 1980. Molecular 945 parameters of maturation in the Toarcian shales, Paris Basin, France-I. Changes in the configurations of acyclic isoprenoid alkanes, steranes and triterpanes. Geochimica et 946 947 Cosmochimica Acta 44, 1709-1721.
- 948 Meyers, P.A., Ishiwatari, R., 1993. Lacustrine organic geochemistry—an overview of indicators of organic matter sources and diagenesis in lake sediments. Organic geochemistry 20, 867-900. 949
- 950 Mildenhall, D.C., 1989. Terrestrial palynology. In: Barrett, P.J. (Ed.), Antarctic Cenozoic history from
- 951 the CIROS-1 drillhole, McMurdo Sound. DSIR Publishing, Wellington, New Zealand, pp. 119-952 127.
- Miller, M.C., McCave, I.N., Komar, P., 1977. Threshold of sediment motion under unidirectional 953 954 currents. Sedimentology 24, 507-527.
- Moossen, H., Bendle, J., Seki, O., Quillmann, U., Kawamura, K., 2015. North Atlantic Holocene 955 climate evolution recorded by high-resolution terrestrial and marine biomarker 956
- 957 records. Quaternary Science Reviews 129, 111-127.

- Morin, R.H., Williams, T., Henrys, S.A., Magens, D., Niessen, F., Hansaraj, D., 2010. Heat flow and
 hydrologic characteristics at the AND-1B borehole, ANDRILL McMurdo Ice Shelf Project,
 Antarctica. Geosphere 6, 370-378.
- 961 Naish, T.R., Barrett, P.J., Dunbar, G.B., Woolfe, K.J., Dunn, A.G., Henrys, S.A., Claps, M., Powell,
 962 R.D., Fielding, C.R., 2001. Sedimentary cyclicity in CRP drillcore, Victoria Land Basin,

963 Antarctica. Terra Antarctica 8, 225-244.

- Naish, T., Powell, R., Levy, R., Wilson, G., Scherer, R., Talarico, F., Krissek, L., Niessen, F.,
- 965 Pompilio, M., Wilson, T., Carter, L., DeConto, R., Huybers, P., McKay, R., Pollard, D., Ross,
- 966 J., Winter, D., Barrett, P., Browne, G., Cody, R., Cowan, E., Crampton, J., Dunbar, G., Dunbar,
- 967 N., Florindo, F., Gebhardt, C., Graham, I., Hannah, M., Hansaraj, D., Harwood, D., Helling, D.,
- 968 Henrys, S., Hinnov, L., Kuhn, G., Kyle, P., Läufer, A., Maffioli, P., Magens, D., Mandernack,
- 969 K., McIntosh, W., Millan, C., Morin R., Ohneiser, C., Paulsen, T., Persico, D., Raine, I., Reed,
- 970 J., Riesselman, C., Sagnotti, L., Schmitt, D., Sjunneskog, C., Strong, P., Taviani, M., Vogel, S.,
- 971 Wilch T., Williams, T., 2009. Obliquity-paced Pliocene West Antarctic ice sheet oscillation.
- 972 Nature 458, 322-328.
- Ourisson, G., Albrecht, P., 1992. Hopanoids. 1. Geohopanoids: the most abundant natural products on
 Earth? Accounts of Chemical Research 25, 398-402.
- Peters, K. E., Moldowan, J. M., 1991. Effects of source, thermal maturity, and biodegradation on the
 distribution and isomerization of homohopanes in petroleum. Organic geochemistry 17, 47-61.
- 977 Pole, M., Hill, B., Harwood, D., 2000. Eocene plant macrofossils from erratics, McMurdo Sound,
- Antarctica. In: Stilwel, J.D., Feldman, R.M. (Eds.), Paleobiology and Paleoenvironments of
 Eocene Rocks: McMurdo Sound, East Antarctica, American Geophysical Union, Washington,
 D.C., pp. 243-251.
- Powell, R., Krissek, L.A., Van der Meer, J., 2000. Preliminary depositional environmental analysis of
 CRP-2/2A, Victoria Land Basin, Antarctica: palaeoglaciological and palaeoclimatic
 inferences. Terra Antarctica 7, 313-322.
- Powell, R.D., Cooper, J.M., 2002. A glacial sequence stratigraphic model for temperate, glaciated
 continental shelves. Geological Society, London, Special Publications 203, 215-244.
- Powell, R.D., Domack, E.W., 2002. Modern glacimarine environments. In: Menzies, J. (Ed.), Modern
 and Past Glacial Environments, Butterworth-Heinemann, Boston, pp. 361-390.
- 988 Poynter, J.G., Farrimond, P., Robinson, N., Eglinton, G., 1989. Aeolian-derived higher plant lipids in
- 989 the marine sedimentary record: Links with palaeoclimate. In: Leinen, M., Sarnthein, M.,
- 990 (Eds.), Paleoclimatology and paleometeorology: modern and past patterns of global991 atmospheric transport, Springer, Netherlands, pp. 435-462
- 992 Prebble, J.G., Raine, J.I., Barrett, P.J., Hannah, M.J., 2006a. Vegetation and climate from two
- 993 Oligocene glacioeustatic sedimentary cycles (31 and 24 Ma) cored by the Cape Roberts Project,

- 994 Victoria Land Basin, Antarctica. Palaeogeography, Palaeoclimatology, Palaeoecology 23, 41-995 57.
- Prebble, J.G., Hannah, M.J., Barrett, P.J., 2006b. Changing Oligocene climate recorded by
 palynomorphs from two glacio-eustatic sedimentary cycles, Cape Roberts Project, Victoria
 Land Basin, Antarctica. Palaeogeography, Palaeoclimatology, Palaeoecology 231, 58-70.
- 999 Pross, J., Contreras, L., Bijl, P.K., Greenwood, D. R., Bohaty, S.M., Schouten, S., Bendle, J.A., Röhl,
- 1000 U., Tauxe, L., Raine, J.I., Huck, C.E., van de Flierdt, T., Jamieson, S.S.R., Stickley, C.E., van
- 1001 de Scootbrugge, B., Escutia, C., Brinkhuis, H., Integrated Ocean Drilling Program Expedition
- 318 Scientists, 2012. Persistent near-tropical warmth on the Antarctic continent during the earlyEocene epoch. Nature 488, 73-77.
- 1004 Rees-Owen, R.L., Gill, F.L., Newton, R.J., Ivanović, R.F., Francis, J.E., Riding, J.B., Vane, C.H., dos
 1005 Santos, R.A.L., 2018. The last forests on Antarctica: Reconstructing flora and temperature from
- 1006 the Neogene Sirius Group, Transantarctic Mountains. Organic Geochemistry
- doi.org/10.1016/j.orggeochem.2018.01.001.
- 1008 Ribecai, C., 2007. Early Jurassic miospores from Ferrar Group of Carapace Nunatak, South Victoria
 1009 Land, Antarctica. Review of Palaeobotany and Palynology 144, 3-12.
- Rohmer, M., Bouvier-Nave, P., Ourisson, G., 1984. Distribution of hopanoid triterpenes in
 prokaryotes. Microbiology 130, 1137-1150.
- Sachse, D., Radke, J., Gleixner, G., 2006. δD values of individual n-alkanes from terrestrial plants
 along a climatic gradient–Implications for the sedimentary biomarker record. Organic
 Geochemistry 37, 469-483.
- Sackett, W.M., Poag, C.W., Eadie, B.J., 1974. Kerogen recycling in the Ross Sea,Antarctica. Science 185, 1045-1047.
- Sandroni, S., Talarico, F., 2004. Petrography and provenance of basement clasts in CIROS-1 core,
 McMurdo Sound, Antarctica. Terra Antarctica 11, 93-114.
- Sandroni, S., Talarico, F.M., 2011. The record of Miocene climatic events in AND-2A drill core
 (Antarctica): Insights from provenance analyses of basement clasts. Global and Planetary
 Change 75, 31-46.
- Schefuß, E., Ratmeyer, V., Stuut, J.B.W., Jansen, J.H.F., Sinninghe Damsté, J.S., 2003. Carbon
 isotope analyses of n-alkanes in dust from the lower atmosphere over the central eastern
- 1024Atlantic. Geochimica et Cosmochimica Acta 67, 1757-1767.
- Schellekens, J., Buurman, P., Pontevedra-Pombal, X., 2009. Selecting parameters for the
 environmental interpretation of peat molecular chemistry–a pyrolysis-GC/MS study. Organic
 Geochemistry 40, 678-691.
- Schellekens, J., Buurman, P., 2011. n-Alkane distributions as palaeoclimatic proxies in ombrotrophic
 peat: the role of decomposition and dominant vegetation. Geoderma 164, 112-121.

- Scherer, R., Hannah, M., Maffioli, P., Persico, D., Sjunneskog, C., Strong, C. P., Taviani, M., Winter,
 D., 2007. Palaeontologic characterisation and analysis of the AND-1B core, ANDRILL
- 1032 McMurdo Ice Shelf Project, Antarctica. Terra Antarctica 14, 223-254.
- Schröder, H., Paulsen, T., Wonik, T., 2011. Thermal properties of the AND-2A borehole in the
 southern Victoria Land Basin, McMurdo Sound, Antarctica. Geosphere 7, 1324-1330.
- 1035 Seifert, W. K., Moldowan, J. M., 1978. Applications of steranes, terpanes and monoaromatics to the
- 1036 maturation, migration and source of crude oils. Geochimica et Cosmochimica Acta 42, 77-95.
- Seifert, W. K., Moldowan, J. M., 1980. The effect of thermal stress on source-rock quality as
 measured by hopane stereochemistry. Physics and Chemistry of the Earth 12, 229-237.
- Shackleton, N.J., Kennett, J.P., 1975. Paleotemperature history of the Cenozoic and the initiation of
 Antarctic glaciation: oxygen and carbon isotope analyses in DSDP Sites 277, 279, and

1041 281. Initial reports of the deep sea drilling project 29, 743-755.

- Smellie, J.L., 2001. History of Oligocene erosion, uplift and unroofing of the Transantarctic
 Mountains deduced from sandstone detrital modes in CRP-3 drillcore, Victoria Land Basin,
 Antarctica. Terra Antarctica 8, 481-490.
- Strogen, D.P., Bland, K.J., 2011. Hydrocarbon Risk for Drilling on the Coulman High, GNS Science
 Consultancy Report Volume 2011/185, GNS Science, Lower Hutt, New Zealand.
- Strong, C. P., Webb, P. N., 2000. Oligocene and Miocene Foraminifera from CRP-2/2A, Victoria
 Land Basin, Antarctica. Terra Antarctica 7, 461-472.
- Sugden, D.E., Marchant, D.R., Denton, G.H., 1993. The case for a stable East Antarctic ice sheet: the
 background. Geografiska Annaler. Series A. Physical Geography 75, 151-154.
- Sugden, D., Denton, G., 2004. Cenozoic landscape evolution of the Convoy Range to Mackay Glacier
 area, Transantarctic Mountains: onshore to offshore synthesis. Geological Society of America
 Bulletin 116, 840-857.
- Talarico, F., Sandroni, S., Fielding, C. R., Atkins, C., 2000. Variability, petrography and provenance
 of basement clasts in core from CRP-2/2A, Victoria Land Basin, Antarctica. Terra Antarctica 7,
 529-544.
- Talbot, H. M., Farrimond, P., 2007. Bacterial populations recorded in diverse sedimentary
 biohopanoid distributions. Organic Geochemistry 38, 1212-1225.
- The Shipboard Scientific Party, 1975a. Shipboard Site Reports: Sites 270, 271, 272. Initial Reports of
 the Deep Sea Drilling Project 28, 211-334.
- 1061 The Shipboard Scientific Party, 1975b. Shipboard Site Reports: Site 274. Initial Reports of the Deep1062 Sea Drilling Project 28, 369-433.
- Tissot, B.P., Welte, D.H., 1984. Petroleum Formation and Occurrence (2nd ed.). Springer-Verlag,
 Berlin Heidelberg.
- Truswell, E.M., Drewry, D.J., 1984. Distribution and provenance of recycled palynomorphs in
 surficial sediments of the Ross Sea, Antarctica. Marine Geology 59, 187-214.

- 1067 Venkatesan, M.I., 1988. Organic geochemistry of marine sediments in Antarctic region: marine lipids
 1068 in McMurdo Sound. Organic Geochemistry 12, 13-27.
- Vogts, A., Moossen, H., Rommerskirchen, F., Rullkötter, J., 2009. Distribution patterns and stable
 carbon isotopic composition of alkanes and alkan-1-ols from plant waxes of African rain forest
 and savanna C₃ species. Organic Geochemistry 40, 1037-1054.
- Warny, S., Askin, R.A., Hannah, M.J., Mohr, B.A., Raine, J.I., Harwood, D.M., Florindo, F., 2009.
 Palynomorphs from a sediment core reveal a sudden remarkably warm Antarctica during the
 middle Miocene. Geology 37, 955-958.
- 1075 Webb, P.N., Harwood, D.M., McKelvey, B.C., Mercer, J.H., Stott, L.D., 1984. Cenozoic marine
 1076 sedimentation and ice-volume variation on the East Antarctic craton. Geology 12, 287-291.
- 1077 Whittaker, J.M., Müller, R.D., 2006. Seismic stratigraphy of the Adare Trough area,

1078 Antarctica. Marine geology 230, 179-197.

- 1079 Wilson, D.S., Luyendyk, B.P., 2009. West Antarctic paleotopography estimated at the Eocene-
- 1080 Oligocene climate transition. Geophysical Research Letters 36, L16302,
- doi:10.1029/2009GL039297.
- Zhou, W., Xie, S., Meyers, P.A., Zheng, Y., 2005. Reconstruction of late glacial and Holocene
 climate evolution in southern China from geolipids and pollen in the Dingnan peat
- 1083 climate evolution in southern China from geolipids and pollen in the Dingnan peat
- sequence. Organic Geochemistry 36, 1272-1284.