- 1 How likely was a green Antarctic Peninsula during warm Pliocene interglacials? A critical
- 2 reassessment based on palynofloras from James Ross Island

4 Ulrich Salzmann<sup>a\*</sup>, James B. Riding<sup>b</sup>, Anna E. Nelson<sup>c</sup>, John L. Smellie<sup>d</sup>

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- 6 a School of the Built and Natural Environment, Northumbria University, Ellison Building,
- 7 Newcastle upon Tyne NE1 8ST, U.K., ulrich.salzmann@northumbria.ac.uk
- 8 b British Geological Survey, Kingsley Dunham Centre, Keyworth, Nottingham NG12 5GG,
- 9 U.K., j.riding@bgs.ac.uk
- 10 c British Antarctic Survey, High Cross, Madingley Road, Cambridge CB3 0ET, U.K.,
- annaelizabethlaloe@googlemail.com
- d Department of Geology, University of Leicester, University Road, Leicester LE1 7RH,
- 13 U.K., <u>ils55@le.ac.uk</u>

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- 15 <u>Corresponding author:</u> Tel. +44 (0)191 2273874. *E-mail address*:
- 16 ulrich.salzmann@northumbria.ac.uk (U. Salzmann)

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ABSTRACT

- 20 The question whether Pliocene climate was warm enough to support a substantial vegetation
- 21 cover on Antarctica is of great significance to the ongoing and controversial debate on the
- stability or dynamism of Antarctic ice sheets during warm periods with high greenhouse gas
- 23 concentrations. Here we present a systematic palynological comparison of pollen and
- 24 dinoflagellate cyst assemblages from Early Pliocene diamictites collected from the northern
- and eastern Antarctic Peninsula. The sedimentary successions are exceptionally well dated
- using a combination of <sup>40</sup>Ar/<sup>39</sup>Ar and <sup>87</sup>Sr/<sup>86</sup>Sr isotope analyses on interbedded lavas and
- 27 pristine bivalve molluscs. Well-preserved pectinid shells and cheilostome bryozoans suggest
- that the palynomorph-bearing sediments were deposited during warmer Pliocene
- 29 interglacials. The palynological analyses presented here do not identify any *in-situ* pollen and
- 30 spores which indicate the presence of substantial vegetation cover. Direct comparisons
- 31 between palynomorph assemblages of Pliocene diamictites and the underlying Cretaceous
- 32 succession, which included fluorescence microscopy, show that most of the palynomorphs
- are reworked from Upper Campanian and Lower Maastrichtian sediments. Our study

suggests a local provenance for most of the Hobbs Glacier diamictites, whilst sediments at the western coast might have been produced by ice sheets from the Antarctic Peninsula. The presence of the acritarch *Leiosphaeridia* indicates sea-ice during Pliocene interglacials with summer sea surface temperatures ranging from -2 to +5°C. Whilst this implies near modern climate conditions during the Late Neogene, the presence of the dinoflagellate cyst *Bitectatodinium tepikiense* at one location suggests that sea surface temperatures might have been substantially warmer during some interglacials. The absence of *in-situ* pollen and spores in the James Ross Island diamictites cannot be taken as proof of non-existent vegetation. However, this paper presents indirect multiple proxy evidence which makes the presence of a substantial Pliocene vegetation cover on James Ross Island unlikely and supports previous reconstructions of a permanent ice sheet on the West Antarctic Peninsula throughout the Late Neogene.

Keywords: Pliocene; Antarctica; vegetation; palynomorphs; Neogene; sea ice.

#### 1. Introduction

A recent research focus has been on polar regions which may experience environmental conditions similar to those of the warm interglacial periods of the Pliocene (ca. 5.3 to 2.6 Ma ago) towards the end of this century (IPCC, 2007; Salzmann et al., 2009; Lunt et al., 2010). During the Pliocene, global mean annual surface temperature was about 2-3°C higher than pre-industrial levels (Haywood et al., 2000) with elevated atmospheric CO<sub>2</sub> concentrations ranging between 270 and 450 ppmv (Pagani et al., 2010; Seki et al., 2010). This warming was particularly accentuated at high latitudes. For the Canadian high Arctic, recent multi-proxy studies suggest Pliocene mean annual temperatures were about 19 °C warmer than at present (Ballantyne et al., 2010). In contrast to the Arctic, Pliocene palaeenvironmental reconstructions of the southern polar regions are scarce and there is a great uncertainty in estimating the magnitude of temperature increase and ice sheet extent during the warm interglacials of the Neogene. Antarctica holds by far the largest amount of freshwater on Earth. Given current climate concerns, it is of vital importance to understand how southern polar environments and ice sheets responded and might respond in the future to increasing temperatures under high greenhouse gas concentrations.

This paper presents palynological analyses of exceptionally well dated, Early Pliocene to Late Miocene samples from the Antarctic Peninsula located at the northernmost edges of the Antarctic Ice Sheet (Fig. 1). Due to its geographical position and topography, the Peninsula environment is very sensitive to climate change and temperatures have risen rapidly throughout the last decades resulting in a recent substantial thinning of the West Antarctic Ice Sheet (Vaughan et al., 2003; Pritchard et al., 2009; Steig et al., 2009). For the Middle Miocene (ca. 14 Ma), multiple palaeoecological proxies suggest that East Antarctica was still warm enough to support a substantial tundra vegetation in coastal regions, before further cooling brought a full polar climate to this region (Lewis et al., 2008; Williams et al., 2008; Warny et al., 2009). However, despite further rapid cooling, temperatures during Pliocene interglacials might still have been warm enough to cause a substantial reduction in sea-ice cover (e.g. Hillenbrand and Ehrmann, 2005) and fluctuation of the West Antarctic ice sheet (Naish et al., 2009), which might have collapsed during the maxima of Pliocene warmth (Pollard and DeConto, 2009). Pliocene vegetation reconstructions based on climate models indicate the presence of woody Nothofagus-dominated tundra shrub vegetation in coastal regions of Antarctica during Pliocene warm periods (Fig. 1, Salzmann et al., 2008). However, palaeobotanical data which could support these model simulations are extremely scarce. Most uncertainties in reconstructing Pliocene Antarctic vegetation are caused by the dynamic origin of sedimentary deposits which have been disturbed and reworked through glacial transport. Reworking strongly hampers an unambiguous identification of in-situ material which provoked controversial debates about existing Pliocene environmental reconstructions and dating using age-diagnostic microfossils (Francis and Hill, 1996; Stroeven et al., 1996; Harwood and Webb, 1998; Wilson et al., 2002).

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The aim of this paper is to test whether the Antarctic Peninsula was covered by a substantial vegetation cover during warm Pliocene interglacials. We present a systematic analysis and comparison of palynomorph assemblages from Pliocene diamictites taken from coastal, ice-free regions of James Ross Island, northern Antarctic Peninsula, (Fig. 1, Nelson et al., 2009). Well-preserved bivalve shells and interbedded lavas allow an exceptionally good dating control via <sup>40</sup>Ar/<sup>39</sup>Ar and <sup>87</sup>Sr/<sup>86</sup>Sr isotope analyses (Smellie et al., 2006; Smellie et al., 2008). To distinguish between *in-situ* and reworked material, we compare Neogene and underlying Cretaceous sediments using a multi-method approach, including:

a) fluorescence microscopy to identify taphonomic histories (e.g. van Gijzel, 1967; Yeloff and Hunt, 2005)

- b) colour and preservation of palynomorhs to identify thermal maturation (e.g. Pirrie et al.,
   1997; Prebble et al., 2006b)
- 101 c) age indicative dinoflagellate cysts (e.g. Troedson and Riding, 2002; Williams et al., 2004)
- d) potentially age indicative acritarchs (e.g. Hannah, 2006; Warny et al., 2006; Warny et al., 2007)
  - e) potentially age indicative ratio of pollen taxa (Fleming and Barron, 1996)

The paper presents both a reconstruction of the Neogene environmental history of James Ross Island and a systematic feasibility study of different methods previously applied in Antarctic palynological studies.

# 2. Geographical setting and previous work on the Pliocene of James Ross Island

James Ross Island is a volcanic island located on the eastern side of the northern Antarctic Peninsula (Fig. 1). The island is occupied by a large central polygenetic shield volcano which is covered by a 200-400 m thick ice cap rising to more than 1600 m a.s.l. (Nelson et al., 2009). The regional stratigraphy is characterised by 5 km of marine volcaniclastic sediments of the James Ross Basin comprising a stratigraphical succession of Jurassic deep marine mudstones at its western margins, overlain by Lower Cretaceous to Upper Eocene back-arc basin sediments (e.g. Pirrie et al., 1992). Neogene glaciomarine sediments unconformably overlie Cretaceous sediments, or are interbedded within extensive basaltic lavas of the James Ross Island Volcanic Group (JRIVG). The Neogene diamictites sampled here belong to the Hobbs Glacier Formation of Pirrie et al. (1997), and are exposed at the low-lying, ice-free regions of the northern and eastern coasts of James Ross Island (Fig. 1a; Smellie et al., 2006). The diamictites were deposited as debris flows presumably close to the grounding-line of marine-terminating glaciers. A full explanation of the stratigraphy and geology for each sample site was given by Nelson et al. (2009).

Samples from Forster Cliff, Cascade Cliff, Pecten Spur and Ekelöf Point (Fig. 1a) contain numerous, relative intact fossil pectinid shells of *Austrochlamys* sp. and cheilostome bryozoans (Fig. 2). The good preservation of macrofossils and a dominance of local JRIVG clast lithologies suggest that the bivalves have not been transported far within the glacigenic debris flow deposits (Nelson et al., 2009). The bivalves were probably living on the sea floor

during periods of ice advance during warm Late Neogene interglacials. Growth increment analyses coupled with stable isotope data ( $\delta^{18}$ O/ $\delta^{13}$ C) indicate much reduced sea ice with environmental conditions allowing growth throughout much of the year, even during the coldest winters (Williams et al., 2010). Mean Annual Range of Temperature (zs-MART) analyses of fossil bryozoans sampled at Cascade Cliff (Fig. 1a) indicate increased seasonality, and an overall warmer climate for the Weddell Sea region during the Early Pliocene (Clark et al., 2010).

<sup>87</sup>Sr/<sup>86</sup>Sr isotope dates of pectinid shells (*Austrochlamys* sp.) and <sup>40</sup>Ar/<sup>39</sup>Ar dates from underlying and overlying lavas show that most palynomorph-bearing diamictites were deposited after 3.69 Ma, during the Zanclean stage (Table 1; Smellie et al., 2006; Smellie et al., 2008; unpublished data). The <sup>40</sup>Ar/<sup>39</sup>Ar ages provide the best estimate for the depositional ages of the sediments, whereas the wide range of <sup>87</sup>Sr/<sup>86</sup>Sr isotope dates indicate some reworking of the pectinid shells. Previous palynological analyses of Neogene diamictites collected near Hamilton Point on southeast James Ross (Fig. 1) recorded only few *in-situ* Late Neogene dinoflagellate cysts (Pirrie et al., 1997). The vast majority of palynomorphs were interpreted as being reworked from the Upper Cretaceous. Neogene pollen and spores could not be identified and this prevented any reconstruction of potential vegetation on James Ross Island during warm Pliocene interglacials.

**3. Methods** 

### 3.1 Palynology

Fifty-four Neogene diamictites and five underlying Cretaceous sandstones, collected in the Austral summer of 2006-2007 on the northern and western snow-free coastal regions of James Ross Island (Fig. 1a, Nelson et al., 2009), were processed in the palynolgy laboratory of the British Geological Survey. Preparation of the samples followed standard techniques involving hydrochloric and hydrofluoric acids (Faegri and Iversen, 1989). In order to identify a potential bias on fluorescence colours of palynomorphs by different preparation techniques, selected samples were processed using both acid -and non-acid techniques (Riding and Kyffin-Hughes, 2010). The paper focuses on 12 samples from nine locations (Fig. 1a) with good palynomorph preservation, allowing a total count of ca 130-280 pollen and spores and 10-360 dinoflagellate cysts and acritarchs per sample (Fig. 3, Table 2). Palynomorphs were identified using literature (e.g. Duane et al., 1992; Williams et al., 2004; Raine et al., 2008).

For the three sample locations, Hamilton Point, Roundel Point and the Watchtower, palynomorph assemblages were analysed from both the Pliocene and underlying Cretaceous successions to facilitate the identification of *in-situ* and reworked material. The Cretaceous samples are D6.212.1, D6.221.1 and D6.209.2. Colour and preservation were noted for each grain and compared between samples.

## 3.2. Fluorescence Microscopy

Fluorescence microscopy has become a widely used technique in palynology to identify reworked and *in-situ* material because the fluorescence colour of palynomorphs changes with increasing age (e.g. van Gijzel, 1967; Waterhouse, 1998; Yeloff and Hunt, 2005). For this study, grains were examined with 100x and 400x magnification under UV-fluorescence using an OlympusBH-2 microscope with high pressure mercury burner and 330-385 exciter filter following techniques described by Yellof and Hunt (2005). The fluorescence colour of palynomorphs in the Cretaceous samples was used as a "calibration standard" to identify potentially younger, *in-situ* palynomorphs in the Upper Neogene diamictites. Particular attention has been paid to the comparison of *Nothofagidites* pollen which occurs in both the Cretaceous and Neogene of the Antarctic Peninsula.

## 3.3. Statistical Analysis

In order to identify groups of palynomorphs and their provenance, non-parametric multivariate tests were applied using the statistical software package PRIMER (release 6, Primer-E Ltd). Similarity matrices were constructed employing the Bray Curtis similarity measure after square root-transforming data to reduce the weight of highly dominant taxa. The relationship between samples was based on two dimensional ordination of similarity matrices calculated from mean values and is presented by two-dimensional non-metric multidimensional scaling plots (nMDS, Fig. 4). The extent to which the rank order of distance between samples in the ordination agrees with the rank from the similarity matrices is indicated by the stress coefficient (i.e. the lower the stress, the better the agreement).

#### 4. Results

### 4.1 Palynomorphs

The palynomorph assemblages of the samples from James Ross Island are dominated by the pollen *Nothofagidites lachlaniae*, *N. flemingii*, *Podocarpidites* and *Peninsulapollis*, and the spores *Cyatheacidites*, *Lycopodiumsporites* and *Laevigatosporites ovatus* (Fig. 3). Other abundant taxa which regularly occur in most samples are *Alisporites*, *Proteacidites* and *Microcachryidites antarcticus*. Sample 5.30.11 from Lachman Crags, located on northwest James Ross Island (Fig. 1), differs from all other samples in showing low percentages of *Nothofagidites* and a higher diversity of Pteridophyta. The *Nothofagidites lachlaniae/N. flemingii* ratio is slightly higher in samples located in southeast James Ross Island (e.g. Hobbs Glacier D6.204, Hamilton Point D6.212, Roundel Point D6.209). Pairs of Cretaceous and Neogene samples collected at the same locations show very similar pollen and spore assemblages. This is in particular true for Roundel Point (D6.209), which is characterised by abundant *Podocarpidites*.

With the exception of sample D5.30.11 from Lachman Crags, all samples yielded a relatively rich and diverse assemblage of dinoflagellate cysts and acritarchs (Table 2). The most abundant dinoflagellate cysts are *Exochosphaeridium* and *Isabelidinium*, with the latter particularly frequent in samples from southeast James Ross Island (e.g. Hobbs Glacier D6.204, Hamilton Point D6.212, Roundel Point D6.209). Again, the palynomorph assemblages from pairs of Cretaceous and Neogene samples collected at the same location are very similar. *Trichodinium castanea* occurs in high numbers in both Cretaceous and Neogene samples from Roundel Point (D6.209). Most of the dinoflagellate cysts in the Neogene samples can be confidentially attributed to the Late Cretaceous (e.g. *Isabelidinium pellucidum, Odontochitina porifera, Nelsoniella tuberculata*), whereas only two samples at Cascade Cliffs (D5.8.9) and Hobbs Glaciers (D6.204.9) contain dinoflagellate cysts indicative of Eocene to Oligocene (*Deflandrea heterophlycta, Enneadocysta partridgei*) and Miocene (*Bitectatodinium tepikiense*) ages (Harland, 1978; Stover et al., 1996; Williams et al., 2004). The dinoflagellate cyst *Impletosphaeridium* spp. is abundant in all samples, whereas *Leiosphaeridia* was recorded in the Pliocene samples only.

## 4.2 Preservation, fluorescence colour and thermal maturity

The palynomorphs exhibit variable preservation in all samples independent of their age. Different stages of deterioration, caused by oxidation and mechanical damage, were recorded for the same taxa (e.g. *Nothofagus lachlaniae*) in both Cretaceous and Pliocene

samples. High, age-independent variability was also observed of the colour of palynomorphs which allowed no reliable estimates of thermal maturity.

Fluorescence strongly varied with taxon, ranging from yellow-orange to red-brownish colours. No differences in fluorescence colour was recorded between the Cretaceous and Pliocene samples. *Nothofagidites* pollen, for example, gave a generally weak red-brownish fluorescence signal which varied with preservation, independent of the age. The age-diagnostic Paleogene dinoflagellate cysts *Deflandrea heterophlycta* and *Enneadocysta partridgei* are the only exception, showing a very bright, almost bluish colour. The difference in fluorescence colour may indicate that they are younger (van Gijzel, 1967). However, the Miocene to Holocene dinoflagellate cyst *Bitectatodinium tepikiense* (see Harland, 1978; Stover el al., 1996; Pirrie et al, 1997), gave a surprisingly weak fluorescence signal which did not significantly differ from those of the Cretaceous reworked taxa.

## 4.3 Multidimensional scaling: patterns of similarities

The two dimensional non-metric multidimensional scaling (nMDS) shows very similar relationship patterns of samples for pollen/spores and dinoflagellate cysts/acritarchs, suggesting that both assemblages were controlled by similar environmental forcing (Fig. 4). The nMDS plot also shows that the Neogene palynomorphs strongly resemble their Cretaceous counterparts. Similarities between the Neogene and the underlying Cretaceous samples are often greater than between samples of similar age. The nMDS plot also indicates that the samples with the largest geographical distance often show the greatest dissimilarities in palynomorph assemblages. Samples from the northeastern part of James Ross Island (Lachman Crags D5.30.1, Cascade Cliffs D5.8.9, Forsters Cliff D5.10.2) closely resemble each other and are distinctively different from samples from the southeast (e.g. Hamilton Point D6.212, Watchtower D6.221 and Roundel Point D6.209).

## 5. Discussion

# 5.1. Identification of *in-situ* and reworked palynomorphs in the Pliocene diamictites

Well-preserved fossil pectinid shells of *Austrochlamys* sp. and cheilostome bryozoans suggest that the pollen-bearing sediments have not been transported over long distances and were probably deposited during relatively warm Pliocene periods (Nelson et al., 2009). However, whilst macrofossils are deposited *in-situ*, many organic-walled microfossils are

transported or reworked from older sediments. Our palynological analysis of pollen and spores has identified *Nothofagus*-dominated forest vegetation with *Podocarpus*, Proteaceae and ferns growing in a moist warm-temperate climate. Similar vegetation communities have been reconstructed for the Antarctic Peninsula from the Late Cretaceous to the Paleogene (e.g. Askin, 1997; Dettmann and Thomson, 1987; Troedson and Riding, 2002). Reworking from older sediments is also shown by age-indicative dinoflagellate cysts such as Isabelidinium pellucidum, Odontochitina porifera and Nelsoniella tuberculata. Detailed descriptions of Late Cretaceous dinoflagellate cyst assemblages from James Ross Island were given by Dettman and Thomson (1987), Dolding (1992) and Keating (1992). However, although it is evident that most of the palynomorphs are reworked from the Cretaceous, the presence of Bitectatodinium tepikiense at Hobbs Glacier also indicates that Late Neogene insitu palynomorphs have also been preserved. This raises the question as to which pollen and spores could also originate from the local Neogene *Nothofagus*-dominated tundra on James Ross Island, which may have been similar to tundra shrub vegetation previously reconstructed for the Miocene and possibly Pliocene coastal regions of East Antarctica (Francis abd Hill, 1996; Lewis et al., 2008; Warny et al., 2009). In the following section, we will discuss and review various methods applied to identify *in-situ* palynomorphs. Their unambiguous identification is the first necessary step towards a robust reconstruction of Pliocene vegetation and climate on the West Antarctic Peninsula.

### 5.1.1. The Nothofagidites lachlaniae/N. flemingii pollen ratio

The question whether or not *Nothofagus* was present during warm Neogene interglacials is a focus of many palaeobotanical studies on Antarctica (e.g. Fleming and Baron, 1996; Francis and Hill, 1996). Whereas fossil leaves or woods provide direct evidence for its presence, vegetation reconstructions using fossil *Nothofagidites* pollen are strongly hampered by pollen morphological constraints. As the geographical distribution of the southern beech genus *Nothofagus* ranges from warm temperate forest to cold polar tundra shrub, any meaningful palaeoenvironmental reconstruction requires an unambiguous identification to species level. However, the most abundant southern beech pollen in samples from James Ross Island is classified as *Nothofagidites lachlaniae* and *N. flemingii*, both referring to pollen types rather than real plant species. Because of uncertainties in the identification of the nearest living analogue, indirect measures such as the dominance of morphotypes (Hill and Truswell, 1993) or ratio of pollen types (Fleming and Baron, 1996)

have been postulated to infer the presence of *in-situ Nothofagidites* pollen. In their study, Fleming and Baron (1996) interpreted the dominance of *N. lachlaniae* pollen and low abundances of *N. flemingii* in marine cores as an indicator for the presence of *Nothofagus* tundra on Antarctica during the Pliocene. Our systematic comparison of Pliocene and underlying Cretaceous samples from James Ross Island strongly questions this approach in showing a clear dominance of *N. lachlaniae* pollen types in all samples. Variations in the ratio of *N. lachlaniae* versus *N. flemingii* pollen occurs in samples from the northeast of James Ross Island (e.g. Lachman Crags, D5.30.11) and points to changes in sediment provenance rather than implying the presence of *in-situ* material and the existence of past *Nothofagus* tundra shrub.

### 5.1.2. Colour and Preservation

The colour and preservation of fossil palynomorphs is a widely-used indicator of thermal maturation and hence the relative age of palynomorphs. The method has been successfully applied to distinguish between *in-situ* and reworked material in Paleogene/Neogene samples from Antarctica (e.g. Askin and Raine, 2000; Prebble et al., 2006a). The absence of any notable difference in colour or preservation of palynomorphs between the Cretaceous and Neogene control samples from James Ross Island might indicate absence of *in-situ* material. However, the high variability of preservation and colour within one sample generally questions the suitability of this approach for the palynological analyses of sediments from the Antarctic Peninsula. It should be noted that the Cretaceous sediments underlying the Hobbs Glacier formation largely consist of relatively soft, unconsolidated sand and siltstones which contain generally well preserved palynomorphs with a low thermal maturity. Damage and corrosion appears to be primarily the result of post-sedimentary glacial transport during the Neogene, and does therefore not necessarily imply an older age.

## 5.1. 3. Fluorescence microscopy

The fluorescence colours of palynomorphs from Cretaceous and Neogene samples closely resemble each other, suggesting that they either have similar geological ages or similar taphonomic histories (van Gijzel, 1967). No difference in fluorescence colours could be recorded between the acid -and non-acid treated samples. Waterhouse (1998) noted that the fluorescence colour of palynomorphs can also change with increasing corrosion and oxidation. This could explain the surprisingly weak fluorescence colour signal of the much

younger, Late Neogene dinoflagellate cyst *Bitectatodinium tepikiense* occurring in samples from Hobbs Glacier. Post-sedimentary glacial transport and corrosion during the Neogene might have altered the fluorescence of both Cretaceous and Neogene palynomorphs from James Ross Island. The exceptionally strong fluorescence signal of the age-indicative Eocene to Oligocene dinoflagellate cysts *Deflandrea heterophlycta* and *Enneadocysta partridgei* might have been caused by a different taphonomic history and provenance rather than reflecting the younger geological age.

### 5.1.4. Age-diagnostic acritarchs

The acritarch *Leiosphaeridia* and the dinoflagellate cyst *Impletosphaeridium* are regularly abundant in marine sediments from polar regions. Warny et al. (2007) concluded from the presence of high numbers of *Impletosphaeridium* in marine cores taken offshore the Antarctic Peninsula that the samples are likely to have been derived from sediments of Eocene to Miocene age. Our direct comparison of Late Neogene and Cretaceous samples from James Ross Island which are both rich in *Impletosphaeridium* strongly questions the suitability of this taxon for estimating geological ages. Given the highly abundant occurrences in some Upper Cretaceous samples from James Ross Island (e.g. D6.221.1, Table2), it is also highly unlikely that this acritarch, which is now extinct and has no modern analogue, could indicate proximate sea ice cover, as suggested by Warny *et al.* (2007).

Through mapping the modern distribution of the acritarch *Leiosphaeridia* in surface samples from the Arctic Ocean, Mudie (1992) showed that this acritarch occurs in high numbers at the contact margin between pack ice and seasonal ice. Accordingly, *Leiosphaeridia* has also been used in marine sediments from Antarctica as an indicator to reconstruct past sea-ice presence (e.g. Hannah, 2006; Prebble et al., 2006b; Warny et al., 2006). Given the ecological range and modern distribution of *Leiosphaeridia* in cold polar regions, it is likely that this acritarch in samples from James Ross Island are *in-situ* and have not been reworked from the warmer, presumably ice-free Cretaceous into the Upper Neogene diamictites. The absence of *Leiosphaeridia* in all Cretaceous control samples strongly supports this interpretation.

## 5.2. Provenance and glacier flow directions

Pirrie et al. (1997) concluded from sedimentological and palynological studies on diamictites from the Hobbs Glacier Formation in southeast James Ross Island that the

sediments were deposited by glaciers flowing from the Antarctic Peninsula towards the southeast. The majority of dinoflagellate cysts were derived from the Upper Campanian to Lower Maastrichtian strata of the Marambio Group. Our new data from different northern and eastern locations on James Ross Island show similar palynomorph assemblages and confirm the Late Cretaceous age and origin of most reworked palynomorphs. However, in showing local patterns and a close resemblance between Neogene and underlying Cretaceous palynomorphs, our study suggests a more local provenance of the Hobbs Glacier diamictites (Fig. 4). This finding corroborates detailed provenance studies by Nelson et al. (2009), suggesting that the main ice centres for the debris flows were situated on James Ross Island itself. According to their conceptual ice model, most diamictites in the east and northeast originated from a central large ice cap positioned over Mount Haddington (Fig. 1). Ice sheets from the Antarctic Peninsula delivered material across Prince Gustav Channel to the western parts of James Ross Island only. Again our new palynological data support this model in showing significantly different palynomorph assemblages for northwest James Ross Island (Lachman Crags, D5.30.11).

The presence of Eocene/Oligocene dinoflagellate cysts in samples from Hobbs Glacier and Cascade Cliffs cannot be explained with existing glacier flow models. There is no evidence of Paleogene strata outcropping on James Ross Island and the closest Eocene formations are on Seymour and Cockburn islands (Pirrie et al., 1992), located about 15-20 km to the east, which is contrary to the main direction of reconstructed ice flows. Given the vicinity of Seymour Island, it appears likely that the Paleogene dinoflagellate cysts were reworked into the Neogene diamictites through drifting icebergs or long-distance aeolian transport. The strong fluorescence signal of the Eocene/Oligocene dinoflagellate cysts supports this interpretation in suggesting a different taphonomic history.

# 5.3. Pliocene environments: how likely was a vegetation cover?

Our systematic analyses of pollen and spores in Neogene samples from James Ross Island give no evidence for a substantial vegetation cover on James Ross Island during warm Pliocene interglacials. Strong similarities between the Neogene and underlying Cretaceous palynomorph assemblages and fluorescence signals point to the absence of any *in-situ* pollen and spores of Late Neogene age. Unfortunately the absence of fossils can never be taken as proof of non-existence because sedimentary and post-sedimentary processes could have prevented the preservation or deposition of Pliocene pollen and spores. However, instead of

stressing the absence of palynomorphs as proof, we use a number of indirect proxies to assess the likelihood of vegetation on James Ross Island during Pliocene warm interglacials.

The presence of Paleogene and *in-situ* Late Neogene dinoflagellate cysts demonstrates that sedimentary conditions were favourable to the preservation of organic-walled microsfossils. It seems therefore unlikely that pollen and spores originating from a potential Pliocene vegetation cover have been selectively destroyed. Late Neogene dinoflagellate cysts have been recorded at Hobbs Glacier in low numbers. However, Pirrie et al. (1997) also recorded coeval dinoflagellate cysts in samples from James Ross Island, indicating that, although in low numbers, *in-situ* palynomorphs appear to be a constant element of the palynological assemblage.

Smellie et al. (2009) presented evidence from sedimentology and modelling simulations, suggesting that the Antarctic Peninsula ice sheet (APIS) was present even during the warmest interglacials of the Pliocene. The study points out significant differences in the sensitivity of APIS to climate change compared to West and East Antarctica. It is likely that a permanent APIS, fed by high snow accumulation, could have also prevented the growth of vegetation on James Ross Island. The occurrence of *Leiosphaeridia* in samples from James Ross Island implies the presence of sea-ice with summer sea surface temperature ranging from ca. -2 to +5°C (Edwards et al., 1991; Mudie, 1992). This interpretation corroborates modelling simulations predicting sea-ice cover throughout the year during the Early Pliocene with seasonal temperature variation at the sea surface (0-5m) ranging from -1.69 °C in winter and +3.08 °C during summer (Lunt et al., 2008; Williams et. 2010). Percentages of *Leiosphaeridia* in the James Ross Island diamictites are comparatively low, and it is possible that sea ice cover was reduced during the Pliocene as suggested by Williams et al (2010) from growth increment analysis and stable isotopic data of fossil bivalves (*Austrochlamys anderssoni*) from Cockburn Island.

Whilst the presence of *Leiospharidia* suggests near-modern climate conditions in the West Antarctic Peninsula, recently published ZS-MART analyses of bryozoans in Late Neogene diamictites from Cascade Cliffs (Fig. 1a) points to much higher temperatures with increased seasonality (Clark et al. 2010). The palynomorph assemblage from Hobbs Glacier (D6.204.9) supports this interpretation in providing no evidence for the sea-ice indicator *Leiospharidia* coupled with the occurrence of *Bitectatodinium tepikiense*, indicative of warmer, sub-arctic to temperate sea surface temperatures (Edwards et al. 1991). The discrepancies in climate estimates from the Antarctic Peninsula diamictites result from

uncertainties in dating control. It is not clear from the isotope dates (Table 1) whether the palynomorphs analysed from different locations and different depths originate from the same warm interglacial period. It is highly likely that the James Ross Island diamictites cover various Late and Early Pliocene interglacials with different annual temperature ranges.

#### 6. Conclusions

Since the Oligocene, Antarctica has been shaped by ice sheets and glacial dynamics resulting in the reworking of sediments through post-sedimentary transport. Glacial dynamics, which increased with further cooling towards the end of the Miocene, strongly reduces the chances to find any *in-situ* microfossils and undisturbed fossil records which are required for an unambiguous reconstruction of Neogene Antarctic environments. Our new palynological analyses of Early Pliocene diamictites from James Ross Island failed to identify *in-situ* pollen and spores which could indicate the presence of substantial vegetation on the Antarctic Peninsula. Our systematic comparison of Pliocene and underlying Cretaceous assemblages using different palaeoecological methods furthermore revealed that some previously published palynological approaches to identify Late Neogene Antarctic environments cannot be applied to sediments from the Antarctic Peninsula. Such approaches include the use of colour of palynomorphs for estimating relative geological ages and pollen ratio to identify in-situ *Nothofagus* pollen. Our results highlight the importance of identifying the provenance of glacial sediments and their potential microfossil "contaminants" before attempting reconstructions of Neogene palaeoenvironments in Antarctica.

Whilst the absence of *in-situ* pollen and spores cannot be taken as a proof of non-existence of vegetation, we still conclude from this study, using indirect multiple evidence, that the presence of a substantial Pliocene vegetation cover on James Ross Island was rather unlikely, even during warm interglacials. Evidence which supports this conclusion includes: a) the presence of acritach *Leiosphaeridia*, indicating sea-ice and sea surface temperatures close to modern values.
b) the presence of *in-situ* dinoflagellate cysts which indicate sedimentary conditions favourable to the preservation of palynomorphs. Selective destruction of *in-situ* pollen and spores is unlikely.

c) sedimentological characteristics of fossiliferous, pollen bearing diamictites, signifying episodes of ice expansion during relatively warm interglacials of the Early Pliocene (see also Nelson et al., 2009).

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Insufficient dating control prevents our palynomorph assemblages, indicating near modern conditions, to be related to specific periods of the Early Pliocene. The fossil palynomoph assemblages might represent communities which lived during the same or different interglacial periods. The sporadic occurrence of the sub-arctic to temperate dinoflagellate cyst *Bitectatodinium tepikiense* in one sample points to the existence of particularly warmer interglacials which might have not been adequately covered herein. Further research is needed to decipher and increase age and resolution control of interglacial records to enable a robust reconstruction of Late Neogene environments of the Antarctic Peninsula.

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#### CAPTIONS FOR FIGURES AND TABLES

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Fig.1. a) The locations of the Cretaceous and Neogene sample sites on James Ross Island. b) 717 A climate-vegetation model simulation (BIOME4-HadAM3) of the Mid Pliocene (ca. 3 Ma) 718 vegetation on Antarctica (after Salzmann et al., 2008). 719 720 Fig. 2: A sample of diamictite from Ekelöf Point, southeast James Ross Island (D6.260.x) 721 with well-preserved *Pecten* shells (*Austrochlamys* sp.). 722 723 Fig 3: The relative abundances of selected pollen and spore taxa, expressed as percentages of 724 the total counts of palynomorphs. Cretaceous samples are highlighted in yellow. 725 726 Fig. 4: Two dimensional non-metric multidimensional scaling (nMDS) for a) pollen and 727 spores, and b) dinoflagellate cysts and acritarchs in the diamictites from James Ross Island. 728 729 The samples from the northern and western coasts respectively are indicated by the following symbols: (▲) and (▼) The other symbols indicate pairs of Cretaceous and Pliocene samples 730 from the west coast of James Ross Island. 731 732 Table 1. The ages of the Neogene palynomorph-bearing diamictites derived from the mean 733 <sup>87</sup>Sr/<sup>86</sup>Sr dates of pectinid shells (*Austrochlamys* sp.) and the <sup>40</sup>Ar/<sup>39</sup>Ar dates from the 734 underlying and overlying basaltic lavas (after Smellie et al., 2006, 2008 and unpublished 735 data). 736 737 Table 2: Total counts of dinoflagellate cyst and acritarch specimens. Age-diagnostic 738 Paleogene/Neogene taxa are framed. The Cretaceous samples are highlighted in yellow. 739