

Antarctic Science page 1 of 20 (2024) © The Author(s), 2024. Published by Cambridge University Press on behalf of Antarctic Science Ltd. This is an Open Access article, distributed under the terms of the Creative Commons Attribution licence (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted re-use, distribution and reproduction, provided the original article is properly cited.

doi:10.1017/S0954102024000129

Evidence for a floristically diverse rainforest on the Falkland archipelago in the remote South Atlantic during the mid- to late Cenozoic

ZOË A. THOMAS (1,2,3), MICHAEL MACPHAIL (1,4), HAIDEE CADD^{3,5}, DAVID J. CANTRILL (1,6,7), DAVID K. HUTCHINSON⁸, HEATHER A. HAINES^{2,3,9}, KAREN PRIVAT^{2,10}, CHRIS TURNEY^{2,3,11}, STEFANIE CARTER^{12,13} and PAUL BRICKLE^{12,14}

¹School of Geography and Environmental Science, University of Southampton, Southampton, UK
²Earth and Sustainability Science Research Centre (ESSRC), School of Biological, Earth and Environmental Sciences, University of New South Wales, Sydney, NSW, Australia

³Chronos ¹⁴Carbon-Cycle Facility, Mark Wainwright Analytical Centre, University of New South Wales, NSW, Australia

⁴Australian National University, Canberra, ACT, Australia

⁵ARC Centre of Excellence for Australian Biodiversity and Heritage, University of Wollongong, Wollongong, NSW, Australia
⁶Royal Botanic Gardens Victoria, Melbourne, VIC, Australia

⁷School of BioSciences, University of Melbourne, Parkville, VIC, Australia

⁸Climate Change Research Centre, School of Biological, Earth and Environmental Sciences, University of New South Wales, Sydney, NSW, Australia

⁹University of Nevada, Reno, NV, USA

¹⁰Electron Microscope Unit, Mark Wainwright Analytical Centre, University of New South Wales, Sydney, NSW, Australia
¹¹Division of Research, University of Technology Sydney, Sydney, NSW, Australia
¹²South Atlantic Environmental Research Institute (SAERI), Stanley, Falkland Islands
¹³UK Centre for Ecology and Hydrology, Bangor, UK
¹⁴School of Biological Sciences (Zoology), University of Aberdeen, Aberdeen, UK
2.thomas@soton.ac.uk

Abstract: We report the discovery of an ancient forest bed near Stanley, on the Falkland Islands, the second such ancient deposit identified on the South Atlantic island archipelago that is today marked by the absence of native tree species. Fossil pollen, spores and wood fragments preserved in this buried deposit at *Tussac House* show that the source vegetation was characterized by a floristically diverse rainforest dominated by *Nothofagus*-Podocarpaceae communities, similar to cool temperate *Nothofagus* forests/woodlands and Magellanic evergreen *Nothofagus* rainforests. The age limit of the deposit is inferred from the stratigraphic distribution of fossil pollen species transported by wind, birds or ocean currents from southern Patagonia, as well as similar vegetation types observed across the broader region. The deposit is suggested to be between Late Oligocene and Early Miocene, making it slightly older than the previously analysed Neogene West Point Island forest bed (200 km west of *Tussac House*). The combined evidence adds to our current knowledge of the role of climate change and transoceanic dispersal of plant propagules in shaping high-latitude ecosystems in the Southern Hemisphere during the late Palaeogene and Neogene.

Received 26 July 2023, accepted 25 February 2024

Keywords: biodiversity, forest bed, fossil pollen, lignite deposit, Miocene, Neogene, Oligocene, Palaeogene

Introduction

The Falkland Islands (Malvinas) are some of a number of remote islands and archipelagos at high to polar latitudes in the Atlantic, Indian and Southern oceans that once supported rainforest and rainforest scrub (woody spp. <5 m tall) in the pre-Quaternary period (Cantrill & Poole 2012). Evidence for the former presence of

tree-dominated vegetation includes *in situ* tree stumps, wood, leaves and fossil pollen and spores (Edwards 1921, Cookson 1947, Birnie & Roberts 1986, Poole & Cantrill 2007, Seward & Conway 1934, Stone *et al.* 2005, Truswell *et al.* 2005). In almost all cases, the palaeofloras are assumed to have colonized the islands via transoceanic dispersal of seeds and spores by wind, ocean currents and birds from adjacent larger

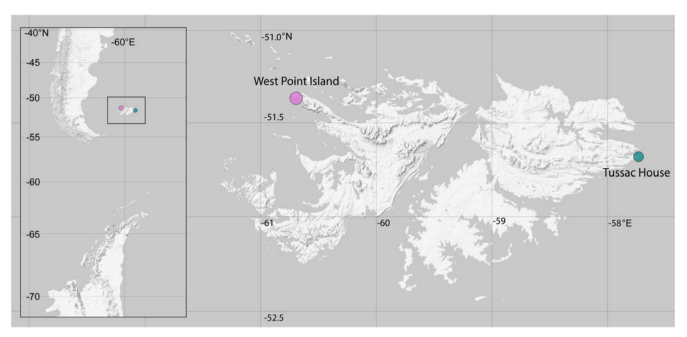


Figure 1. Location of *Tussac House* (green dot) and West Point Island (purple dot) on the Falkland Islands (inset, map of the South Atlantic region). Map produced using Generic Mapping Tools (GMT).

landmasses during globally warm periods (e.g. Late Cretaceous or Eocene; Zachos *et al.* 2001, Macphail & Cantrill 2006, Ortiz-Jaureguizar & Cladera 2006).

Here, we report the discovery of a buried lignitic deposit during excavations for the foundations of the Tussac House aged care centre on the foreshore at Stanley, the capital city on the eastern island of the Falkland Islands (Fig. 1). This is the second of two lignitic deposits (forest beds) found in the Falkland Islands that preserve evidence of forest or woody plant communities growing on this now-treeless archipelago (Macphail et al. 1994). The first discovery in 1899 comprised a lignitic deposit with tree trunks > 1.5 m in diameter found outcropping on the foreshore of West Point Island, a small island on the western side of the archipelago (Halle 1911). While it was initially proposed that this deposit may have been an accumulation of driftwood (Baker 1924), analysis of its stratigraphic relationships confirmed that the lignite was in situ, while the associated fossil pollen and spores suggested that the deposit was Late Neogene in age, representative of a time period when climate conditions allowed the growth of forests (Birnie & Roberts 1986). More recent palynological investigation by Macphail & Cantrill (2006) and an investigation of the wood assemblage (Poole & Cantrill 2007) of the West Point Island deposit narrowed its age limits to Middle Miocene to Early Pliocene. Here, we document the microfloras and wood fragments preserved in the Tussac House deposit. We infer the age limits by a combination of stratigraphic distribution of fossil pollen species transported by wind/birds from southern Patagonia and by comparing coeval floras present in Patagonia-Tierra del Fuego, the South Shetland Islands and West Antarctica.

Setting

The Falkland Islands lie at 52°S, 540 km east of the coast of South America. Unlike many volcanic islands off the Chilean coast, the Falkland Plateau is considered to be a rifted part of the African Plate associated with the break-up of Gondwana and the opening of the Atlantic Ocean during the Mesozoic (Stanca et al. 2022). The modern climate is made of up two main zones: subpolar oceanic climate (Köppen classification: Cfc) in lowland areas and polar tundra climate (Köppen classification: ET) in upland areas (Beck et al. 2018), being highly influenced by the surrounding cool South Atlantic waters. There are some isolated areas of cold desert (BWk) and cold steppe (BSk) in lowland areas influenced by rain shadows. The mean annual temperature is 5.5°C, with high mean monthly and annual wind speeds of ~8.5 m s⁻¹ (with prevailing westerly winds) and relatively low annual precipitation of ~600 mm, distributed uniformly throughout the year (Lister & Jones 2014). Although the Falkland archipelago is ~8° latitude north of the southern limits of tree growth in southern South America, trees do not now grow naturally on the islands, and vegetation is dominated by acid grasslands, including whitegrass (Cortaderia pilosa) and diddle-dee (Empetrum rubrum), with tussac grass (Poa flabellata) found in coastal areas.

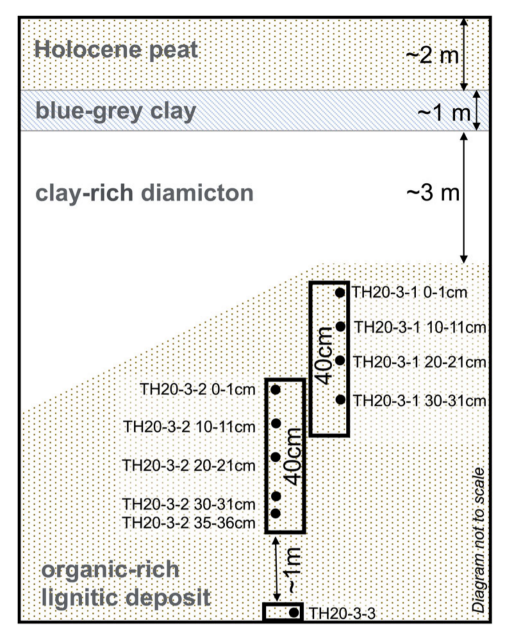


Figure 2. Lithostratigraphy of the Tussac House site.

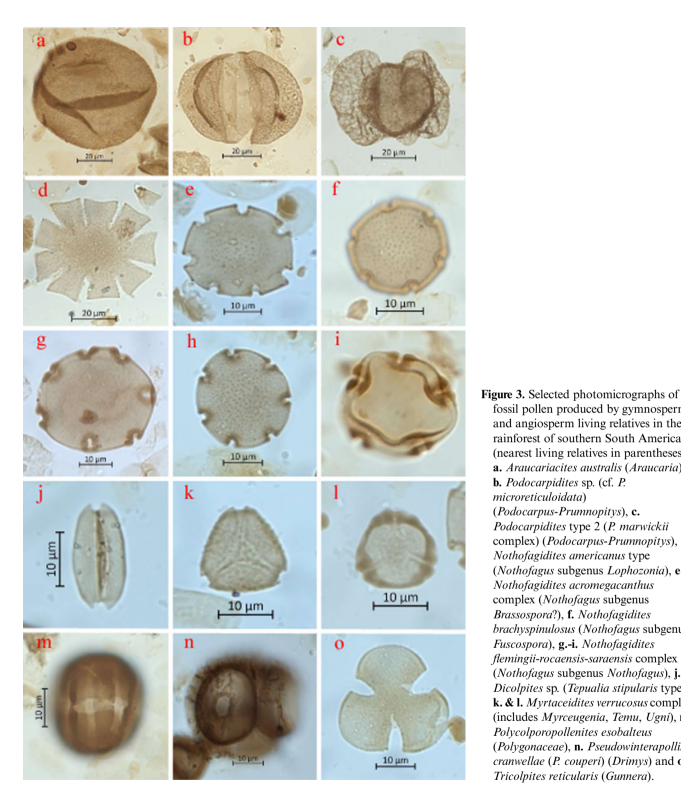
Woody shrubs, such as native boxwood (*Veronica elliptica*), are relatively uncommon and confined to topographically sheltered areas. Reasons for the lack of timber-sized trees (and tree species) include the subdued topography, very extensive periglacial activity but limited glacier formation during Pleistocene glacial stages and prevalence of gale-force winds (Clark & Wilson 1992, Rosenbaum 1996, Hamson *et al.* 2008).

The *Tussac House* site is located on the foreshore of Stanley Harbour (51.6949°S, 57.8315°W). Geotechnical boreholes drilled on the site indicate that the subsurface geology consists of ~2 m of Holocene peats sequentially underlain by periglacial sediments comprising a 1 m-thick blue-grey clay and ~3 m thick clay-rich

diamicton, which unconformably overlie the lignitic organic deposit. The total thickness of this organic deposit is unknown due to it meeting water table depth, but it is likely to be > 2 m thick.

Methods

A geotechnical excavation in March 2020 uncovered a lignitic wood-rich matrix at ~6 m depth. During a subsequent geotechnical dig in May 2020, the section was sampled, with the recovery comprising a 60 cm lignitic section (Fig. 2) and several large wood fragments (Supplemental Table 1a), probably representing tree-sized parent plants.



fossil pollen produced by gymnosperm and angiosperm living relatives in the rainforest of southern South America (nearest living relatives in parentheses): a. Araucariacites australis (Araucaria), b. Podocarpidites sp. (cf. P. microreticuloidata) (Podocarpus-Prumnopitys), c. Podocarpidites type 2 (P. marwickii complex) (Podocarpus-Prumnopitys), d. Nothofagidites americanus type (Nothofagus subgenus Lophozonia), e. Nothofagidites acromegacanthus complex (Nothofagus subgenus Brassospora?), f. Nothofagidites brachyspinulosus (Nothofagus subgenus Fuscospora), g.-i. Nothofagidites *flemingii-rocaensis-saraensis* complex (Nothofagus subgenus Nothofagus), j. Dicolpites sp. (Tepualia stipularis type), k. & l. Myrtaceidites verrucosus complex (includes Myrceugenia, Temu, Ugni), m. Polycolporopollenites esobalteus (Polygonaceae), n. Pseudowinterapollis cranwellae (P. couperi) (Drimys) and o. Tricolpites reticularis (Gunnera).



Figure 4. Selected photomicrographs of pollen of age-diagnostic and phytosociologically significant taxa (nearest living relatives in parentheses; all morphogenera and species are now extinct in southernmost South America and on the Falkland Islands: a. Canthiumidites cf. bellus (Randia type), b. Clavatipollenites glarius (Hedyosum), c. Thymelaepollis sp. of Macphail & Cantrill (Ovidia type), d.-f. Cupanieidites reticularis (Sapindaceae tribe Cupanieae), g. Dacrycarpidites australiensis (Dacrycarpus), h. Dacrydiumites florinii (Dacrydium), i. Microalatidites palaeogenicus (Phyllocladus), j. Phyllocladidites mawsonii (Lagarostrobos franklinii), k. Phyllocladidites elongatus (extinct Lagarostrobos sp.), l. Trichotomosulcites subgranulosus complex (extinct Podocarpaceae) and m.-o. Gen. et sp. nov. of Macphail & Cantrill (2006).

Ten samples of 1 cc of sediment spaced along the 60 cm palynostratigraphic for section were prepared Concentrating investigation (Macphail 2021). oxidation-resistant fossil pollen and spores (miospores) and fungal spores preserved in cultural deposits and soils requires the removal of the inorganic matrix and organic matrix (see Wood et al. 1996). Sample process included 1) treatment with 10% potassium hydroxide, 2) sieving residue through a 100 um mesh sieve, 3) addition of a lycopodium tablet and 10% hydrochloric acid, 4) sodium polytungstate heavy liquid separation at a specific density of 1.8 g/cm² to remove mineral matrix and finally 5) treatment with acetolysis solution (sulphuric acid and acetic anhydride). Aliquots of the organic extracts were set on permanent strew mounts using Petropoxy 154.

The plant microfossil component in each sample was counted using a ZeissTM Photomicroscope II fitted with Plan-NeofluarTM and PlanapoTM objectives, which provide magnifications up to 2000×. Specimens of fossil pollen and spores were counted until statistically robust numbers of fossil pollen and spores were recorded (> 250 specimens), and then the remainder of the strew mounts and additional strew mounts for each sample were scanned for additional rare morphospecies; in the case of low-yield samples, the pollen count comprises the total numbers of fossil pollen and spores present on the microscope slide.

Samples of wood from the section were taken for radiocarbon measurements at the University of New South Wales Chronos ¹⁴Carbon-Cycle Facility to determine whether the material returned infinite radiocarbon ages (i.e. > 50 000 years). Each sample type was pre-treated following the protocols outlined in Turney et al. (2021). The wood specimens were also examined by light microscopy to determine their To investigate taxonomic class. the anatomical characteristics, the material was fractured to expose the three planes of section (transverse and radial and tangential longitudinal) mounted on aluminium stubs. Specimens were not treated/coated prior to imaging. Backscattered electron imaging was conducted using a Hitachi TM4000Plus scanning electron microscope at 15 kV in standard vacuum mode (~30 Pa). Wood specimens are described using the terminology of the IAWA Committee (1989, 2004) wherever possible. We compared samples with the published literature on wood samples from West Point Island (Poole & Cantrill 2007) to facilitate identification.

To provide further context for our plant proxy evidence, we performed a brief comparison of the Falklands climate with palaeoclimate modelling scenarios of the Miocene and Oligocene. Here, we used the simulations of Farnsworth *et al.* (2019), who ran palaeoclimate simulations for each of the periods of interest: namely,

the Langhian (13.8–16.0 Ma), Burdigalian (16.0–20.4 Ma), Aguitanian (20.4–23.0 Ma), Chattian (23.0–27.8 Ma) and Rupelian (27.8-33.9 Ma), as well as a pre-industrial control simulation. These simulations were performed using the HadCM3BL model, palaeogeography reconstructions generated by Getech. The model simulations were coarse resolution (2.50° latitude × 3.75° longitude) to cover many geological periods within the computational resources available. This means that each simulation typically had no land grid cell assigned to the Falkland Islands, or in some cases only a single grid cell, depending on the configuration. Thus, the model can produce a good representation of South American topography and climate through time, whereas the Falkland Islands generally fall below the model resolution.

To overcome the resolution limits, we used the palaeoclimate simulations to reconstruct anomalies of temperature and precipitation at the Falkland Islands geographical coordinates rather than absolute values. These anomalies were then used to adjust a modern high-resolution (30 arc-seconds) climatology of temperature and precipitation (WorldClim version 2; Fick & Hijmans 2017) to create palaeoclimatologies. These palaeoclimatologies were reconstructed using an additive anomaly for temperature and a multiplicative anomaly for precipitation (following Beck *et al.* 2018).

Results

Palynostratigraphic analysis

All samples yielded large to very large numbers of mostly well-preserved terrestrial palynomorphs in organic extracts dominated by cuticle fragments and organic fines. Cysts of marine algae (dinoflagellates) and miospores of freshwater aquatic or semi-aquatic plants are absent except for occasional *Botryococcus* and a single specimen of *Sparganiaceaepollenites* (*Sparganium*, *Typha*) at 20–21 cm in TH20-3-1. Whether or not frequent to common 'amorphous spheres' (cf. *Leiosphaeridia*) in the samples represent freshwater algae is unknown.

Taxonomy

The majority of commonly occurring and some rare miospores in the *Tussac House* samples (Figs 3 & 4) can be assigned to fossil morphospecies formally described from Palaeogene-Neogene deposits in southern Australia, New Zealand and South America (references in Macphail *et al.* 1994, Barreda & Palazzesi 2007, Palazzesi & Barreda 2007, Raine *et al.* 2011). However, the morphological diversity exhibited by some morphospecies and genera is unusually high, particularly within

Table I. Modern affinity of selected fossil morphospecies and genera with Eocene to Miocene time distributions in Patagonia.

Family	Tribe/genus	Fossil morphospecies	Age range (Patagonia)
Amaranthaceae	-	Chenopodipollis chenopodiaceoides	Late Oligocene to Miocene
Anacardiaceae	-	Striatotricolporites gamerroi	Palaeocene to Late Miocene
Aquifoliaceae	Ilex	Ilexpollenites spp.	Eocene
Arecaceae	-	Arecipites spp.	Palaeocene to Late Oligocene
Asteraceae	Tribe Mutisieae	Mutisiapollis spp.	Late Oligocene to Early Miocene
	Subfamily Tubuliflorae	Tubulifloridites spp.	Miocene
Casuarinaceae	Allocasuarina/Casuarina	Haloragacidites harrisii	Palaeocene to Eocene
	Gymnostoma?	Haloragacidites trioratus	Middle Eocene to Early Miocene
Caesalpiniaceae	-	Margocolporites van wijhei	Late Oligocene to Miocene
Chloranthaceae	Hedyosum	Clavatipollenites glarius	Oligocene to Early Miocene
Convolvulaceae	Cressa, Wilsonia	Tricolpites trioblatus	Miocene
Cyperaceae	-	Cyperaceaepollis neogenicus	Miocene
Ericaceae	Empetrum, Gautlheria, Pernettya	Ericipites spp.	Palaeocene to Miocene
Euphorbiaceae	Alchornea	Psilatricolporites operculatus	Late Oligocene to Middle Miocene
·· F	(Austrobuxus/Dissilaria)	Malvacipollis diversus	Eocene
	Cf. Corynabutilon	Malvacipollis robustus ms.	Oligocene to Miocene?
Goodeniaceae	(CoprosmalOpercularia)	Poluspissusites spp.	Late Oligocene to Early Miocene
Grossulariaceae	Cf. Quintinia	Quintiniapollis	Late Oligocene to Early Miocene
Gunneraceae	Gunnera	Tricolpites reticulatus	Palaeocene to Middle Miocene
Loranthaceae	-	Gothanipollis bassensis	Eocene
Malvaceae	-	Baumannipollis, Malvacearumpollis	Late Oligocene to Middle Miocene
Menyanthaceae	(Liparophyllum type)	Striasyncolpites laxus	Early to Middle Miocene
Mimosaceae	Acacia	Acaciapollenites myriosporites	Early Miocene
14111105accac	(Adenanthera type)	Polyadopollenites	Early Miocene
Myrtaceae	(Includes <i>Metrosideros</i> type)	Myrtaceidites parvus-mesonesus	Palaeocene to Miocene
111,111100110	Myrceugenia, Temu, Ugni, etc.	Myrtaceidites verrucosus	Eocene to Miocene
Nothofagaceae	Nothofagus (Lophozonia) spp.	Nothofagidites americanus/tehuelchesii	Eocene to Miocene
Tromoragaeeae	Nothofagus (Brassospora) spp.	Nothofagidites acromegacanthus	Eocene to Miocene
	Nothofagus (Nothofagus) spp.	Nothofagidites flemingii	Eocene to Miocene
	Nothofagus (Nothofagus) spp.	Nothofagidites rocaensis/Saraensis	Oligocene to Miocene
Onagraceae	(Epilobium type)	Corsinipollenites atlantica.	Early to Middle Miocene
Onagraceae	Fuchsia	Diporites aspis	Late Oligocene
Poaceae	-	Graminidites spp.	Late Oligocene to Miocene
Polygonaceae	Persicaria	Glencopollis ornatus	Miocene
Proteaceae	Embothrium	Granodiporites nebulosus	Late Oligocene to Early Miocene
1 Totedecae	-	Peninsulapollis gillii	Palaeocene to Eocene
	Proteacidites cf. Telopea	Triporopollenites ambiguus	Middle Eocene
Rubiaceae	Randia?	Rubipollis oblatus	Late Oligocene
Radiacac	Randia type	Canthiumidites cf. bellus	Late Oligocene to Early Miocene
Sapindaceae	Tribe Cupanieae	Cupanieidites reticularis	Late Oligocene to Middle Miocene
Sparganiaceae	Sparganium	Sparganiaceaepollenites sphericus.	Middle Miocene
Thymelaeaceae	Ovida	Thymelaepollis sp.	Late Palaeogene-Neogene?
Trimeniaceae	-	Periporopollenites demarcatus	Eocene Eocene
Winteraceae	Drimys	Pseudowinterapollis couperi	Oligocene to Middle Miocene
· · · · · · · · · · · · · · · · · · ·	Drunys	1 вешонинстиронь сопрен	ongottic to middle miotelic

morphotypes assigned to *Podocarpidites* (nearest living relative (NLR) *Podocarpus-Prumnopitys*) and *Nothofagidites* (*Nothofagus*). In both cases, individual pollen grains can vary markedly in size and shape. Reasons for this include deposition of both immature and mature pollen grains representing plants whose remains form the lignitic deposit or were growing close to the site (M.K. Macphail, personal observation 2023). In addition, curtailed pollen development may have occurred due to the short growing season at high latitudes and the associated climate stress on flower development.

In this study, the *Podocarpidites* component has been assigned to two broad groups based on the size and

mode of attachment of the sacci to the corpus of the pollen grain, not to species identified in the Neogene flora of Tierra del Fuego by Zamaloa & Romero (2005). The Nothofagidites component is assigned to three broad morphospecies groups comprising **Nothofagidites** acromegacanthus (NLR Nothofagus subgenus Brassospora), Nothofagidites americanus/tehuelchesii (Nothofagidites subgenus Lophozonia) and Nothofagidites flemingiil rocaensis/saraensis (Nothofagidites subgenus Nothofagus). Unidentified triporate, tricolpate and tricolporate pollen grains have been grouped into Proteacidites spp. (Proteaceae), Tricolpites spp. and Tricolporites spp., respectively.

Table II. Relative abundances of commonly occurring and selected uncommon to rare morphospecies recovered from the *Tussac House* samples. Rare taxa (< 1%) within the spore-pollen count are indicated by '+'; very rare taxa recorded outside the spore-pollen count are indicated by 'x'.

Fossil taxon	,	TH20-3-1 (depths in c	m)		TH20	-3-2 (depth	s in cm)		TH20-3-3
	0–1	10–11	20–21	30–31	0-1	10–11	20-21	30-31	35–36	-
			Cr	ptogams						
Baculatisporites turboensis	+	×	×	+	+	1%		+	×	
Cyathidites australis/minor	×	×	×	×	×	×	×	×	×	×
Deltoidospora sp.	×				×		×	+	+	×
Foveotriletes palaequetrus		×	×	×						
Ischyosporites areapunctatis	+	×		+		+	×	+	×	×
Polypodiisporites inanguahuensis			×	×	×	×				
Unassigned trilete spores	+	×		×	+		+		×	
Total count	1%	1%	+	2%	2%	1%	+	2%	1%	+
			-	nnosperms						
Araucariacites australis	+	+	+	+	×	+	+	+	×	+
Cupressacites					+		+			
Dacrycarpites australiensis	+	×	+				+			
Dacrydiumites florinii	3%	4%	6%	3%	4%	2%	5%	4%	2%	1%
Equisetosporites sp.	×				+					
Microalatidites palaeogenicus	1%		+	+	+	+	+		+	+
Phyllocladidites mawsonii complex	42%	44%	45%	45%	34%	45%	42%	45%	48%	48%
Phyllocladidites verrucosus	+	+					×		×	+
Podocarpidites type 1 complex	12%	6%	8%	10%	6%	7%	11%	9%	7%	10%
Podocarpidites type 2 complex	9%	4%	8%	5%	3%	6%	4%	6%	5%	6%
Podosporites parvus				+	×	×	×			×
Trichotomosulcites subgranulosus	3%	2%	1%	2%	6%	1%	1%	1%	+	+
Unassigned gymnosperms	+	+	+	1%	+				+	+
Total count	72%	61%	69%	65%	55%	62%	63%	65%	62%	66%
	/ -		ngiosperms			~=, ~			/-	
Sparganiaceaepollenites			×	(monocot,	,1040110)					
Liliacidites type 2	+	×	+	×		×	+	+	+	+
Luminidites cf. phormoides			×	×	×		·		·	·
Total count	2%	1%	1%	+	1%	1%	2%	+	+	
Total Count	2/0		Angiosperi			1/0	2/0	'	'	
Clavatipollenites glarius		×	×	×	+		×	×		
Compositoipollenites cf. tarragoensis	×	×	+	^			1%	×	+	
Cupanieidites orthoteichus	^	×	Т			×	1 / 0 ×	^	Т	×
=		×	×	+	+	^	+	×	1	×
Ericipites spp.		×	^	т	Τ		т	^	+	^
Gen. et sp. nov.		*		V		V				
Gothanipollis cf. bassensis	.,		+	×		×	+	+	+	+
Granodiporites nebulosus	×									
Malvacipollis robustus ms	207	X	407	C 0 /	70/	C 0 /	607	507	407	50/
Myrtaceidites verrucosus type 1	3%	1%	4%	6%	7%	6%	6%	5%	4%	5%
Myrtaceidites verrucosus type 2	+	+	×				×	×		
Myrtaceidites verrucosus type 3	×				×				×	+
Myrtaceidites sp. (dicolpate)	×	+		+	+	+	+	+	+	
Nothofagidites americanus/tehuelchesii	+	2%	+	+	1%	2%	1%	+	+	+
Nothofagus acromegacanthus	7%	9%	4%	2%	5%	6%	6%	6%	6%	4%
Nothofagidites flemingiillsaraensis	10%	19%	14%	18%	20%	16%	14%	14%	15%	15%
Total Nothofagidites count	17%	30%	18%	20%	26%	24%	21%	20%	23%	19%
Polycolporopollenites esobalteus		×	×	×		×	×	×	×	×
Proteacidites cf. symphyonemoides		+	×	×						
Proteacidites type 1	+	×	+	+	+	+		×	+	+
Proteacidites type 2	×	×	2%	+	+	×		×	+	+
Total Proteacidites count	1%	1%	3%	3%	4%	2%	2%	3%	2%	2%
Pseudowinterapollis couperi	×	+	×	×	×	×	×	+	×	×
Unassigned <i>Rhoipites</i> spp.	+	×	+	+	+	+	+	×	×	+
Striatotricolporites gammeroi			×				cf.			
Thymelaepollis sp. (Ovidia type)	×	×	×	+	×	×	×	×	+	×
Tricolpites reticulatus	+			×	+	+	×	+	+	×

(Continued)

Table II. (continued).

Fossil taxon	,	TH20-3-1 (depths in c	m)		TH20)-3-2 (depth	s in cm)		TH20-3-3
	0–1	10–11	20–21	30–31	0-1	10–11	20–21	30–31	35–36	-
Canthiumidites cf. bellus						×				
Unassigned Triporopollenites spp.	+	×				+				
Inassigned angiosperms	+	+	+	+	+		+	+	+	+
Total dicotyledon count	26%	36%	31%	33%	42%	36%	34%	33%	36%	31%
Pollen sum	591	513	590	609	495	613	634	507	556	555
				Fungi						
Unassigned spores	+	+	1%	+	+	1%	2%	5%	59%	23%

Table III. Tussac House wood anatomical characteristics.

	TH20-3	TH20-5a	TH20-5b	TH20-5c	TH20-6	TH20-8
Anatomical affinity Growth rings • Number of rings	Podocarpaceae xylotype 3 Distinct	Podocarpaceae xylotype Indistinct	Distinct 101	Distinct 110	Xylotype 7 Distinct 56	Podocarpaceae xylotype 3 Distinct
Average growth rate	0.369 mm/year	-	0.407 mm/ year	0.516 mm/year (high variability)	0.514 mm/year	0.256 mm/year
Resin canals Parenchyma • Contents • Strand length	Not observed	Present (rare)	Not observed	Not observed	Present (rare)	Not observed
Tracheids • Spiral thickenings	? absent	? absent	? absent	? absent	? absent	? absent
• Pitting tangential longitudinal section	Uniseriate, contiguous, 15–20 µm diameter	Uniseriate, non-contiguous, 10–15 µm diameter	?	?	Rare	Uniseriate, non-contiguous, 10–15 µm diameter
• Pitting radial longitudinal section	Uniseriate, contiguous	Uniseriate, non-contiguous, 10–15 μm diameter	?	?	rare	Uniseriate, non-contiguous
Rays • Width; height (cells)	Homocellular Uniseriate	Homocellular Uniseriate (number of cells between 2 and 12, $n = 72$)	Homocellular ?	Homocellular Uniseriate (number of cells between 2 and 4, <i>n</i> = 12)	Homocellular Uniseriate (number of cells between 2 and 3, $n = 14$)	Homocellular Uniseriate (number of cells between 3 and 12, $n = 33$)
• Indentures	? absent	? absent	? absent	? absent	? absent	? absent
Cross-field pits	1–4 Cupressoid	? Cupressoid	?	?	1–2 5–10 μm Cupressoid	1–2 5–12 μm Cupressoid
Ray tracheids	? absent	? absent	? absent	? absent	? absent	? absent

More generally, the usage of Australian and New Zealand names for morphologically similar pollen and spore morphotypes in South America has led to a *de*

facto broadening of the original fossil species diagnosis as used in the South American literature, and it is probable that the fossil taxon represents more than one

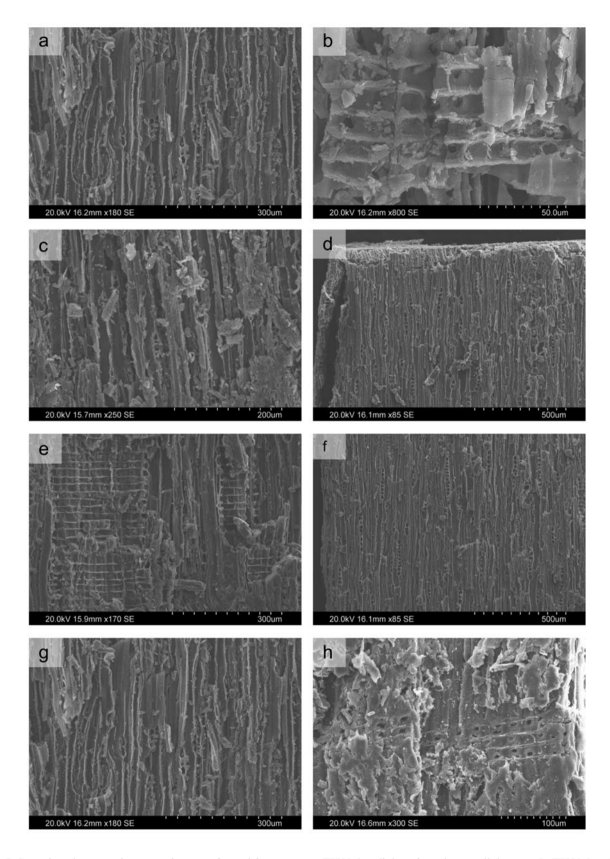


Figure 5. Scanning electron microscopy images of wood fragments. **a.** TH20-3 radial section - homocellular rays, **b.** TH20-3 radial section - smooth cell end wall, **c.** TH20-3 tangential longitudinal section - low-height uniseriate rays, **d.** TH20-5a tangential longitudinal section - smooth end walls, **f.** TH20-5a tangential longitudinal section, **g.** TH20-6 tangential longitudinal section - low-height uniseriate rays and **h.** TH20-6 radial section - cross-field pits.

ecotype, species or genus, some of which are extinct and potentially had different ecological tolerances to their NLRs (Macphail *et al.* 1994, Macphail & Cantrill 2006).

Many of these and less commonly occurring taxa have NLRs in Argentina and/or Chile (Table I), although the taxonomic resolution is mostly low (to genus and family levels), or they are ancestral members of the living clades in South America at least to the family level (cf. Heusser 1971, Markgraf & D'Antoni 1978, Palazzesi & Barreda 2007). At least one novel morphotype (Gen. et sp. nov., this study) appears to be unique to the Falkland archipelago (Macphail & Cantrill 2006).

Dominance and diversity

All fossil assemblages are dominated by fossil species of Podocarpaceae, Nothofagaceae and Myrtaceae (NLRs in parentheses) in which Phyllocladidites mawsonii (Lagarostrobos franklinii) is the dominant morphospecies **Podocarpidites** (Podocarpus/Prumnopitys/ and Nothofagidites Retrophyllum), acromegacanthus, Nothofagidites flemingii/rocaensis/saraensis (Nothofagidites subgenus Nothofagus), **M**vrtaceidites verrucosus (numerous South American Myrtaceae genera) and Proteacidites (Proteaceae) occur in frequent to common numbers (Table II). Estimates of relative abundance are based on statistically robust counts of > 450 fossil pollen and spores (excluding fungal spores) for each sample. Percentages < 1% and morphospecies recorded outside the pollen sum are indicated by '+' and 'x' in Table II, respectively.

Wood analysis

Samples of wood from the section were taken for radiocarbon dating at the University of New South Wales Chronos ¹⁴Carbon-Cycle Facility. All gave infinite radiocarbon ages of > 50 ka (Table III & Supplemental Table 1b), confirming the geological antiquity of the deposit.

Widespread compression made it difficult to diagnose anatomical characteristics of many of the wood fragments (Table III & Supplemental Tables 1a & 1b). Where possible, xylotype was inferred from available diagnostic characteristics and compared directly to the xylotype classification of Poole & Cantrill (2007).

The material observed is assigned to xylotype 3 (Podocarpaceae) documented from the West Point Island deposit (Poole & Cantrill 2007). Although widespread compression makes identification of certain features difficult, some anatomical characteristics are observable (Fig. 5). Samples TH20-3, TH20-5 and TH20-8 appear to have consistent anatomical characteristics where these are observable. Rays are exclusively uniseriate and range between 2 and 12 cells. Cell end walls, when observed, tend to be smooth. In the radial section view, cross-field

pits are generally cupressoid. This xylotype appears to have anatomical similarities to podocarpaceous wood. Xylotype 7 appears to be represented by only one sample: TH20-6. This xylotype is characterized by rare or absent pitting and the presence of resin canals. More material with better preservation is needed for further taxonomic assessment.

Discussion

Most of the fossil pollen and spores preserved in the Tussac House forest bed (Macphail 2021) closely resemble pollen (gymnosperms, angiosperms) and spores (ferns, fern allies) produced by plants in the modern floras of South America and other landmasses in the Southern Hemisphere, although time distributions in southern South America (cf. Tables I & II; Macphail 1999, Partridge 1999, Raine et al. 2011) may vary from, for example, New Zealand and Australia. Some herbs and shrubs remain extant on the Falkland Islands (e.g. Huperzia fuegianum, Amaranthaceae, Asteraceae, Gunnera, Ericaceae (Empetrum, Gaultheria) Myrtaceae (Myrteola); Broughton & McAdam 2005). Other taxa almost certainly are ancestral members of the living clades at least to the family level. In many instances, it is probable that the fossil taxon represents more than ecotype, species or genus, some of which are extinct and potentially had different ecological tolerances to their NLRs (Macphail et al. 1994).

Absolute and probable age limits of the Tussac House forest bed

Unlike Argentina (see Barreda 1996a,b, Barreda & Palamarczuk 2000a,b,c, b, Barreda & Palamarczuk 2000a,b,Barreda et al. 2009, Amenabar et al. 2023), no formal pollen and spore-based palynostratigraphy has been developed to date and correlate Palaeogene and Neogene sediments on the Falkland Islands independently dated marine sediments on the submerged Falkland Plateau or Malvinas Basin (Amenabar et al. 2023 and references therein). Accordingly, the age limits of the West Point Island forest bed and the Tussac House forest bed (Macphail & Cantrill 2006, Macphail 2021) have been inferred using the time distribution of fossil pollen species (morphospecies) whose NLRs occur in mega- and meso-thermal rainforest in Patagonia, commonly defined as encompassing the Argentinean provinces of Neuquén, Río Negro, Chubut, Santa Cruz and Tierra del Fuego (Quattrocchio et al. 2011, Palazzesi et al. 2021 and references therein).

The present study focuses on the time distribution of fossil pollen of three rainforest thermophiles preserved in marine sedimentary formations in Patagonia, viz. *Canthiumidites* cf. *bellus Clavatipollenites* and

Cupanieidites, whose NLRs, respectively, are Randia (now found in dry forest communities in northern South America), Hedyosmum (Chloranthaceae; now found in lowland to montane rainforest, including disturbed areas at low elevations, from 18°N to 25°S) and Cupania (Sapindaceae; now found north of 30°S in southern South America; see Marchant et al. 2002). Isotopic (86Sr/87Sr, Ar/Ar, U-Pb) foraminifer and marine dinoflagellate dinocysts of these formations provide independent age control for the time distributions of the morphospecies in Patagonia (Guler et al. 2021 and references therein, Parras & Cuitino 2021).

Supporting evidence is provided by 1) objective (Jaccard coefficient) comparisons of Oligocene-Miocene palynofloras in Patagonia and Tierra del Fuego, 2) the presence or absence of pollen from wind-pollinated plants dominating Patagonian steppe vegetation (e.g. Nanez et al. 2009, Cornu et al. 2012, 2014) and 3) stratigraphic distributions of taxa that are now prominent in Patagonian steppe (Amaranthaceae, Asteraceae, Poaceae) and Randia type (Canthiumidites cf. bellus), which may be evidence for sourcing of pollen from northern Argentina in the Oligocene-Miocene.

Complicating the palynostratigraphic record of the establishments, extirpations and extinctions of these and other mesotherm taxa are (Barreda & Palazzesi 2007, Barreda et al. 2008, Quattrocchio et al. 2011, Vizcaíno et al. 2013, Estebenet et al. 2014, Sachse et al. 2015, Warny et al. 2019, Parras et al. 2020, Palazzesi et al. 2021, Amenabar et al. 2023): 1) geographical, geomorphic and climatic factors such as the progressive tectonic uplift of the Andes during the Palaeogene-Neogene,) opening of Drake Passage and development of the Antarctic Circumpolar Current in the Late Eocene and earliest Oligocene c. 34-23 Ma, 3) two marine inundation events (c. 22–15 Ma) during the latest Oligocene and Early to Middle Miocene represented by marginal and shallow- to deep-water marine deposits, 4) extinction in Patagonia by the mid-Eocene of previously established mega- and meso-therm taxa followed by the associated emergence of *Nothofagus*-Podocarpaceae communities in the Middle Eocene to Early Oligocene and by the expansion of arid-adapted Patagonian steppe communities in the Late Oligocene and Miocene and 5) marked differences between the palaeofloras occupying coastal and hinterland regions.

Much of the published age-range data from Argentina come from outcrops deposited during the Late Oligocene to Early Miocene *Patagoniense* transgression. Representative sections occur from the Andean Foothills in the north-west to Tierra del Fuego in the extreme south of Argentina, although these represent significantly different periods within the Oligocene to Middle Miocene time (Guerstein *et al.* 2004, Blisniuk *et al.* 2005, Parras *et al.* 2012, Cornou *et al.* 2014,

Estebenet et al. 2014, Kohn et al. 2015, Guler et al. 2021, Parras & Cuitino 2021). For example, the palynostratigraphically well-studied Santa Cruz Formation (Parras et al. 2020) began accumulating at c. 17.5 Ma in the south-west, c. 19.8 Ma in the north-west and c. 19. 3 Ma in the south-east in Santa Cruz Province and at c. 16 Ma in the Golfo San George Basin in the adjacent Chubut Province but is absent further to the north and west in the Valdes and Neuguén basins. Underlying Eocene to Early Miocene formations are mostly confined to specific basins in Patagonia (e.g. the Monte Leon, Río Leona, Río Turbio and San Julien formations (Austral-Magallanes Basin), the Chenque and Centinela formations (Golfo San Jorge Basin) and Slogget Formation (Tierra de Fuego); cf. Guerstein et al. 2004, Parras et al. 2020, Parras & Cuitino 2021, Amenabar et al. 2023). Palynostratigraphic data from marine cores on the Argentine continental shelf are limited to two wells, as are data from confirmed Middle and Late Miocene and Pliocene continental outcrops in Patagonia.

For this reason, and due to the possibility that other relevant records are unpublished, the age limits of samples from the *Tussac House* forest bed are assigned to broad periods of geological time only. The inferred age limits assume that pollen of mesotherm taxa have been transported long distances by wind, birds or insects prior to their extinction or extirpation in continental Patagonia (see Macphail & Cantrill 2006, Macphail 2021). This approach includes caveats that are difficult to test given that the morphospecies are mostly preserved in lignitic sediments where much of the associated palynoflora represents plants growing in or around the site. The reasons for this are as follows:

1) The Falklands flora show a strong affinity with Patagonian South America and Tierra de Fuego and more limited links with the sub-Atlantic Islands, New Zealand and, especially in the past, Australia (see McDowell 2005). However, any colonization of the Falkland archipelago by mega- and meso-thermal plants in the early Palaeogene is most likely to have occurred during periods of global warmth, particularly the Early Eocene hyperthermals. Nevertheless, recognize that rainforest we thermophiles with broad ecological tolerances might have survived after they became extinct in Patagonia (due to increasing aridity) or West Antarctica (due to cold conditions; cf. Zachos et al. 2008, Hyland & Seldon 2018, Fernandez et al. 2021, Srivastava et al. 2023). Nevertheless, given the biologically profound impact of the major global cooling event at the Eocene-Oligocene boundary (Mi-1) elsewhere at midto high latitudes in the Southern Hemisphere (cf. Macphail et al. 1994, 2014), we consider it improbable that any thermophiles survived this event

Table IV. Chronostratigraphic distribution of trace records of fossil *Cupania*-type, *Hedyosmum*-type and *Randia*-type pollen in Patagonia, Argentina. (+) indicates species is present; (-) indicates species was not recorded, (cf.) indicates comparable pollen.

Basin/province	Formation	Age li	mits		Mesothern	n taxa	Ste	ppe taxa		Reference
	(offshore well)	Published	Revised ^a	Randia type	Cupania type	Hedyosmum type	Amaranthaceae	Asteraceae	Poaceae	
Central Chile (Region V1,	(Lauca Basin	Mio-Pliocene boundary	NA	-	-	-	-	+	+	Feitl et al. (2019)
~33°S)	Subtropical Andes)	Latest Miocene	NA	-	cf.	-	-	-	-	Tronosco & Encinas (2006)
Salta Basin	Salto Group	Palaeocene-Early Eocene	NA	-	-	-	-	-	+	Quattrocchio & Volkheimer (1990)
	Metan Subgroup	Undifferentiated Miocene	NA		-	-	-		+	Quattrocchio <i>et al.</i> (2003)
	Cerro Morado	Early-Middle Miocene	NA	-	cf.	-	+	-	-	Barreda et al. (2003)
Neuquen and Nirihuau basins	Salto del Macho	Undifferentiated Palaeogene	NA	-	-	-	-	-		Cornou <i>et al.</i> (2012)
	Salto del Macho	Eocene-Early Oligocene	NA	-	-	-			+	Cornou et al. (2014)
	(YPE.SE.LH x-2)	Middle(?) Miocene	NA	-	-	-	+		+	Ottone et al. (2013)
Colorado Basin	(Cx-1)	Late Miocene	NA	-	-	-		+		Guler et al. (2001)
Catamarca	Chiquimil	Late Miocene	NA	-	-	-	-	+	+	Mautino & Anzotegui (1998, 2002a,b)
Chubut Basin	Chenque	Late Oligocene-Middle Miocene	Middle Miocene	+	+	-	+	+	+	Barreda (1996a)
	Porto Madryn	Middle-Late Miocene	Late Miocene(?)	-	-	-	+	+	+	Palazzesi & Barreda (2004)
Golfo San Jorge Basin	Chenque	Oligocene(?)-Miocene(?)	Middle Miocene	-	-	-	+	+	+	Barreda (1993)
J	Perfiles Mazarredo	Late Oligocene-Miocene	Miocene(?)	-	+	+	-	-	-	Barreda & Palamarczul (2000b)
	Chenque (top)	Early Miocene	Miocene(?)	-	+	+	+	-	-	Barreda & Palamarczul (2000b)
Austral-Magallanes Basin	Río Turbio	Early(?) Eocene	Middle Eocene	-	-	-	-	-	-	Romero & Zamaloa (1985)
	Río Turbio	Middle Eocene	Middle Eocene	-	-	+		-	-	Fernandez et al. (2021)
	Río Leona	Early Late Oligocene	Early Late Oligocene	-	-	+		+	+	Barreda et al. (2009)
	Continental shelf	Late Oligocene-Middle Miocene	NA	+	-	-	+	+	+	Barreda & Palamarczul (2000a)
	San Julian	Undifferentiated Oligocene	Oligocene	+	-	-		+		Barreda (1997)
	San Julian	Late Oligocene	Late Oligocene	-	+	+	+	+	+	Heredia et al. (2012)
	San Julian	Late Oligocene	Late Oligocene	-	-	_	+	-	+	Nanez et al. (2009)
	Continela	Late Oligocene	Late Oligocene-Early Miocene	-	-	-	+	-	+	Guerstein et al. (2004)
	Monte Léon	Early(?) Miocene	Early Miocene	-	-	-	+		+	Barreda & Palamarczul (2000c)

ABLE IV. (continued).

Basin/province	Formation	Ag	Age limits		Mesotherm taxa	n taxa	Ste	Steppe taxa		Reference
	(offshore well)			Randia	Cupania	Randia Cupania Hedyosmum				
		Published	Revised a	type	type type		type Amaranthaceae Asteraceae Poaceae	Asteraceae	Poaceae	
	Monte Léon	Early Miocene	Early Miocene		1	1	+	+	+	Barreda & Palamarczuk (2000c)
	Monte Léon	Early Miocene	Early Miocene		+		+	1	+	Nanez et al. (2009)
	Loreto	Middle Tertiary	NA							Fasola (1969)
Tierra del Fuego	Sloggett	Palaeogene (Late Eocene?)	NA			1				Olivero et al. (1998)
	(Drake Passage)	Mid-Late Eocene	NA							Amenabar et al. (2023)
	(Aries x-1)	Late Oligocene	NA		1		+	1	+	Palamarczuk & Barreda (2000)
	(Aries x-1)	Early Miocene	NA	ı			+	ı	+	Palamarczuk & Barreda (2000)

"After Parras & Cuintino (2021) and Amenábar et al. (2023) NA = not applicable. given the exposed habitats on the topographically subdued Falkland archipelago. Whether any recolonization of the Falklands by thermophiles growing in northern Argentina occurred during the Middle Miocene Climatic Optimum (MMCO) at *c.* 15–17 Ma is unknown.

- 2) Transoceanic dispersal of seeds by wind or water from South America into the Atlantic Ocean is well documented (cf. Renner 2004, Turney et al. 2016, Thomas et al. 2018, Zwier et al. 2022), but successful establishment on remote oceanic islands such as the Falklands is markedly reduced if the 'target' environment(s) differ significantly from those occupied by the parent plants (see McDowell 2005, Ali 2017). Most of these constraints do not apply to pollen or spores of temperate and subtropical plants in South America that are dispersed into the south-west Atlantic or Pacific oceans, although the marine depositional conditions tend to favour long-term preservation more than many terrestrial environments (Kappen & Straka 1988, Montade et al. 2011).
- 3) The rarity of *Canthiumidites* cf. *bellus*, *Clavatipollenites* and *Cupanieidites* and the absence of other South American thermophile pollen types in the *Tussac House* forest bed is likely to reflect deposition in an environment where much of each microflora is derived from locally growing plants (see Barreda *et al.* 2009).
- 4) There is an unknown likelihood that some earlier or later occurrences of *Canthiumidites* cf. *bellus*, *Clavatipollenites* and *Cupanieidites* in Patagonia are unpublished.

Records of *Cupania* type (*Cupanieidites*), *Hedyosmum* type (*Clavatipollenites*) and *Randia* type (*Canthiumidites* cf. *bellus*) obtained from review surveys of the published palynostratigraphic literature for Patagonia and adjacent regions in southern South America (see Barreda *et al.* 2008) are given in Table IV. The fossil record of *Randia* type (*Canthiumidites* cf. *bellus*) in Patagonia is notable as this thermophile is an important element in dry forests in northern South America (Marchant *et al.* 2002).

Maximum age limits of the Tussac House forest bed

On the data available, the parent source(s) of Clavatipollenites but apparently not Cupanieidites or Canthiumidites cf. bellus were present in Patagonia in or by the Middle Eocene. Assuming the former became extinct during the Mi-1 global cooling event, the maximum age limit of the Tussac House forest bed is inferred to be Late Oligocene based on records of Clavatipollenites, Cupanieidites and Canthiumidites cf. bellus in the Austral-Magallanes, Chubut and Golfo San Jorge basins in (or by) the Late Oligocene sediments

(Río Leona, Monte Léon and San Julian formations; Barreda 1997, Barreda *et al.* 2009, Heredia *et al.* 2012). Whether any of the parent sources had reached as far south as Tierra del Fuego by this time is uncertain.

Minimum age limit of the Tussac House forest bed

The minimum age limit of the Tussac House forest bed is no older than Middle Oligocene based on multiple records of Cupanieidites and Clavatipollenites in the Austral-Magallanes Basin (Monte Leon and San Julian formations) and, less certainly, Golfo San Jorge Basin (Perfiles Mazarredo Formation), but it might be as young as Middle Miocene (Cupanieidites, Canthiumidites cf. bellus) based on 86Sr/87SR dates for the Chenque Formation in the Golfo San Jorge (cf. Barreda 1996a, Parras et al. 2020). Whether or not the latter age limit is supported by the trace records of pollen representing the wind-pollinated/dispersed families Amaranthaceae (Chenopodipollis chenopodiaceoides) and Poaceae (Graminidites media) in the Tussac House microfloras is uncertain as the parent plants are recorded in Nothofagaceae-Podocarpaceae communities prior to the Late Miocene expansion of steppe vegetation in Patagonia. The absence of Asteraceae (Tubulifloridites spp.) pollen is less easily explained unless this is due to its lower pollen production and more limited dispersal by wind and animals.

Depositional environment

Relative level data indicate that the top depth of the *Tussac House* lignitic deposit is ~1.5 m above mean high water mark and that the base of the deposit extends below present-day mean sea level and therefore may have accumulated during a period of low relative sea level. The local depositional environment is more likely to have been a damp depression given the virtual absence of obligate or semi-obligate algal cysts or pollen of herbs such as Sparganiaceae (Sparganiaceaepollis) and Menyanthaceae (*Striasyncolpites laxus*), which occur in 'marsh' deposits in Patagonia during the Oligocene-Miocene. One possibility, suggested by the strong east to west alignment of the basement rocks and ongoing tectonic uplift of the archipelago (Stanca *et al.* 2022), is a depression on the back slope on a tilted fault block.

Palaeoflora and vegetation

As the microfloras represent essentially the same Podocarpaceae-*Nothofagus* rainforest or rainforest shrub community (Table II), the data more compelling suggests that the lignites were deposited over a short period of geological time under perhumid cool or, less likely, cold conditions. The closest modern analogues are

Magellanic evergreen *Nothofagus* rainforest (see table 10.1 in Veblen *et al.* 1996). Shared characteristics include the rarity of *Araucaria* and the prominence of *Nothofagus* spp., whose NLRs may be ancestral to extant species in the subgenus *Fuscospora* (e.g. *Nothofagus antarctica*, *Nothofagus betuloides* and *Nothofagus pumilo*) given the high-latitude position of the Falkland Islands throughout the Late Palaeogene and Neogene. The parent plants of *Nothofagidites americanus* and *Nothofagus teheulchesii* may be extinct, as subgenus *Lophozonia* spp. in South America (*Nothofagus alpina*, *Nothofagus obliqua*) are restricted to latitudes above ~40°S in southern Chile and Argentina.

This palaeo-community, however, has no clear modern analogue, as the NLRs of most Podocarpaceae morphospecies are endemic to Tasmania (Lagarostrobos franklinii), New Zealand and/or Southeast Asia (e.g. Dacrycarpus (Dacrycarpites australiensis), Dacrydium (Dacrydiumites florinii) and Phyllocladus (Microalatidites palaeogenicus)). Two podocarp clades are extinct (Podosporites parvus, Trichotomosulcites subgranulosus). The NLRs of most rare ferns and angiosperm shrubs represent humid-demanding species with geographical distributions in South America. Examples are the probable tree fern Cyathidites australis (Cyatheaceae), the club-moss Foveotriletes palaequetrus (Huperzia fuegianum) and Embothrium (Diporites granulosus), all of which occur as far south as Tierra del Fuego, and tree ferns in Andean 'cloud forests' at lower latitudes (Broughton & McAdam 2005). The same is likely to be true of the herb flora whether the unassigned Tricolpites/Tricolporites counts represent herbaceous or woody plants. The absence of Lophosoria (Cyatheacidites annulatus) is surprising given that this ground fern became established elsewhere on sub-Antarctic islands (e.g. the Kerguelen Islands; Cookson 1947) during the Miocene.

Although the scanning electron microscopy analysis of the wood assemblage was able to identify two distinct xylotypes, poor sample preservation precluded a species-level identification and as such is unable to inform on the probable age limits of or genera presented in the Tussac House forest bed. An interesting characteristic observed in the West Point Island deposit was the preponderance of Podocarpaceae pollen (60%) compared to the wood, which is largely cf. Austrocedrus. Although the wood assemblages analysed from the Tussac House deposit were significantly lower in number, a similar trend is apparent. This is probably a reflection of the pollination strategy in Podocarpaceae (wind pollinated) as well as of taphonomic effects. The pollen grains are saccate and tend to float on surface water, so different concentrations of grains from the same vegetation are found depending on whether the grains were deposited from water or onto some sort of

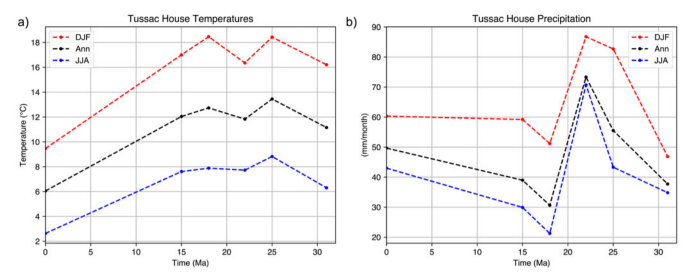


Figure 6. Reconstructed **a.** temperature and **b.** precipitation time series at *Tussac House* based on WorldClim v2 and anomaly-based temperature/precipitation reconstructions for the five palaeo-simulations of the Langhian, Burdigalian, Aquitanian, Chattian and Rupelian. Ann = annual; DJF = December, January, February; JJA = June, July, August.

peat surface (Traverse 2007). Even within standing water the deposition can fluctuate due to differential transport.

Comparison of the Tussac House and West Point Island forest beds

Comparisons between the West Point Island forest bed (Macphail & Cantrill 2006) and Tussac House forest bed microfloras suggest that the latter are slightly older than the former. With few exceptions, all common and most rare morphospecies recorded in the Tussac House forest bed also occur in the West Point Island forest bed. The only significant differences are that the relative abundance of Phyllocladidites mawsonii is lower in the West Point Island microfloras (15-25%) compared to the Tussac House microfloras (34-48%) and that Chenopodipollis chenopodiaceoides, Graminidites and Myrtaceidites eucalyptoides were not recorded in the Tussac House microfloras. Cupanieidites reticularis was not recorded in the West Point Island samples. Although trace occurrences of Myrtaceidites eucalyptoides in the West Point Island microfloras may represent contamination from drilling equipment, the significance of the other differences provides insights into probable age limits given the wide dispersal of Amaranthaceae and Poaceae pollen by wind around the Southern Hemisphere (Macphail 1979) vs the probable dispersal of Cupanieae pollen by insects.

On these data, the probable age limits of the West Point Island forest bed are more likely to be Late Oligocene to Early (Middle?) Miocene than the Middle(?) Miocene to Early Pliocene age limits (Table IV) based in part on *Sparganiaceaepollenites* and *Thymelaepollis* sp. (Macphail & Cantrill 2006). If correct, the revised age

limits for the West Point Island forest bed indicate that Lagarostrobos populations on the Falkland Islands had undergone a significant contraction during the Early to Middle Miocene (i.e. during the warming period leading up to the MMCO at c. 15 Ma). This seems ecologically improbable given the broad distribution of extant Lagarostrobos franklinii from the lowlands to the subalpine/alpine zone in perhumid western Tasmania. Possible explanations are that the sources Phyllocladidites mawsonii pollen on the Falkland Islands were now extinct, cold-demanding ecotypes or, less likely, that warming leading up to the MMCO was accompanied by a significant reduction in precipitation across the archipelago. In either case, the most probable period when Lagarostrobos and any other potentially tree-sized podocarps became extirpated from the Falkland Islands is the Late Miocene, a conclusion that is ecologically consistent with the earlier development of tundra on the South Shetland Islands during the Mi-1 glaciation (Warny et al. 2016).

Climate modelling and interpretation

Anomaly-based reconstructions of annual mean temperature and annual mean precipitation over the Falkland Islands were developed from palaeoclimate modelling scenarios of the Miocene and Oligocene generated by Farnsworth *et al.* (2019) using the HadCM3BL coupled climate model to provide further context for the palaeoflora evidence. Although the modern annual mean temperature of the Falklands is ~6°C, the reconstructed values are significantly warmer, ranging from ~12°C to 13°C in the Early to Middle Miocene (13.8–23.0 Ma), up to 14–15°C in the Late

Oligocene (23.0–27.8 Ma) and then cooler again (11–12°C) in the Early Oligocene (27.8–33.9 Ma; Supplemental Fig. 3). Precipitation varies significantly, from ~500 mm/year in the modern down to 300–400 mm/year for the Middle Miocene, becoming significantly wetter (~700–900 mm/year) in the Early Miocene (20.4–23.0 Ma) and Late Oligocene (23.0–27.8 Ma) and then trending lower again in the Early Oligocene (27.8–33.9 Ma; Supplemental Fig. 4). These patterns are consistent with a hypothesized decrease in precipitation from the Early to Middle Miocene, a possible cause of the changes in *Lagorostrobos* pollen discussed above.

We also use the temperature and precipitation climatologies to generate Köppen climate classifications for each of the Miocene and Oligocene time slices (Supplemental Fig. 5). Based on WorldClim version 2 data (Fick & Hijmans 2017), the modern Falklands climate is predominantly subpolar oceanic (Cfc), with areas of polar tundra (ET) at higher elevations. Going back in time, the Falklands are classified as predominantly cold steppe (BSk) for both the Langhian (13.8-16.0 Ma) and Burdigalian (16.0-20.4 Ma) due to relatively low precipitation and slightly warmer average temperatures. For the Aquitanian (20.4-23.0 Ma) and Chattian (23.0–27.8 Ma) the Falklands become predominantly temperate oceanic (Cfb; i.e. a similar zone to modern-day western Europe), in accordance with higher precipitation during those periods. Further back to the Rupelian (27.8-33.9 Ma) the Falklands are again dominated by cold steppe (BSk) climate.

Palaeoclimate time series for the closest grid point to Tussac House show summer (December-February), winter (June-August) and annual mean reconstructed temperatures and precipitation rates over the mid- to late Cenozoic (Fig. 6). These time series indicate the substantial variations in precipitation palaeo-reconstructions (with periods both wetter and drier than modern conditions), and the temperature reconstructions show that Tussac House was consistently 6-7°C warmer than modern in terms of the annual mean, and with greater seasonal variability, ranging from 6-8°C in winter to 16-18°C in summer. These climatic conditions would have been amenable for the development of the rainforest/scrub communities identified in the pollen spectra within the Tussac House lignite deposit.

Conclusion

The *Tussac House* deposit has a number of similarities to and differences from the West Point Island deposit that was first discovered in 1906. This study affirms that the South Atlantic region hosted diverse rainforests

during the mid- to late Cenozoic period, perhaps in several episodic phases. All fossil pollen and spore assemblages (microfloras) recovered from Tussac House samples represent the same palaeo-community Nothofagus-Podocarpaceae cool temperate rainforest or rainforest scrub - growing on the eastern Falkland sometime between the Late Oligocene Islands (maximum age limit) and the Middle Miocene (minimum age limit). In the absence of local independent age controls, these age limits rely on the time distribution of pollen from mesothermal rainforest plants found in southern Argentina, particularly Cupanieidites reticularis (NLR Sapindaceae tribe Cupanieae) and Clavatipollenites glarius (NLR Hedyosum). The minimum age limit is supported by palaeoclimate modelling and absence the wind-transported pollen types from the arid-adapted steppe vegetation that developed in Argentina during the late Palaeogene-Neogene.

Acknowledgements

Special thanks to the South Atlantic Environmental Research Institute (SAERI) for facilitating this work. Thanks also to Sammy Hirtle, Daniel Ford and the construction team on the *Tussac House* site, who kindly paused their excavation to allow sampling. The authors acknowledge the facilities and the scientific and technical assistance of Microscopy Australia at the Electron Microscope Unit (EMU) within the Mark Wainwright Analytical Centre (MWAC) at UNSW Sydney. We would like to express our sincere gratitude to the two reviewers for their valuable comments and suggestions, which greatly improved the quality of this manuscript.

Financial support

This study was funded by the Australian Research Council Fellowship (DE200100907) awarded to ZAT. DKH is supported by DE220100279.

Competing interests

The authors declare none.

Author contributions

ZAT led the field expedition. ZAT, HC, CT and SC collected the samples. MM and HC undertook the Palaeogene palynostratigraphic and Quaternary pollen analyses and interpretation, respectively. DJC, HAH and KP undertook the wood analysis. DKH undertook the climate modelling. ZAT and MM drafted the

manuscript, and all authors contributed to the preparation of the manuscript.

Climate data availability

The HadCM3BL climate data presented in this study were downloaded from an open access repository: https://www.paleo.bristol.ac.uk/ummodel/scripts/papers/Farnsworth_et_al_2019b.html. WorldClim version 2 climatologies were downloaded from https://www.worldclim.org/data/worldclim21.html.

Supplemental material

To view supplementary material for this article, please visit https://doi.org/10.1017/S0954102024000129.

References

- ALI, J.R. 2017. Islands as biological substrates: classification of the biological assemblage components and the physical island types. *Journal of Biogeography*, 44, 984–994.
- Amenabar, C.R., Guerstein, G.R., Alperin, M., Daners, G., Casadio, S. & Raising, M.R. 2023. Correlation of Middle to Upper Eocene units from high latitude southwestern Atlantic sites. *Ameghiniana*, **60**, 327–341.
- Baker, H.A. 1924. Final report on the geological investigations in the Falkland Islands 1920–1922. Stanley: CSO, 33 pp.
- Barreda, V. 1993. Late Oligocene?-Miocene pollen of the families Compositae, Malvaceae and Polygonaceae from the Chenque Formation, Golfo San Jorge Basin, southeastern Argentina. *Palynology*, 17, 169–186.
- BARREDA, V. 1996a. Biostratigrafía de polen y esporas de la Formación Chenque, Oligoceno tardío?-Mioceno de las provincias de Chubut y Santa Cruz, Patagonia, Argentina. *Ameghiniana*, 33, 35–56.
- BARREDA, V. 1996b. Palinoestratigrafía de la Formación San Julián en el área de Playa La Mina (provincia de Santa Cruz), Oligoceno de la cuenca Austral. *Ameghiniana*, 34, 283–294.
- BARREDA, V. 1997. Palinoestratigrafia de la Formacion San Julian en el area de Playa La Mina (Provincia Santa Cruz) Oligoceno de La Cuena Austral. *Ameghiniana*, 34, 283–294.
- Barreda, V. & Palamarczuk, S. 2000a. Estudio palinoestratigráfico del Oligoceno tardío-Mioceno en secciones de la costa patagónica y plataforma continental argentina. *INSUGEO, Serie Correlación Geológica*, **14**, 103–138.
- BARREDA, V. & PALAMARCZUK, S. 2000b. Palinoestratigrafía de depósitos del Oligoceno tardío-Mioceno, en el área sur del Golfo San Jorge, provincia de Santa Cruz, Argentina. *Ameghiniana*, 37, 103–117.
- Barreda, V. & Palamarczuk, S. 2000c. Palinomorfos continentales y marinos de la Formación Monte León en su área tipo, provincia de Santa Cruz, Argentina. *Ameghiniana*, 37, 3–12.
- BARREDA, V. & PALAZZESI, L. 2007. Patagonian vegetation turnovers during the Paleogene-early Neogene: origin of arid-adapted floras. *Botanical Review*, 73, 31–50.
- Barreda, V., Guler, V. & Palazzesi, L. 2008. Late Miocene continental and marine palynological assemblages from Patagonia. *In Rabassa*, J., ed., *The Late Cenozoic of Patagonia and Tierra del Fuego*. Amsterdam: Elsevier, vol. 16, 343–350.
- Barreda, V., Palazzesi, L. & Marenessi, S. 2009. Palynological record of the Paleogene Rio Leona Formation (southernmost South America: stratigraphical and paleoenvironmental implications. *Review of Palaeobotany and Palynology*, **154**, 22–33.

- Barreda, V., Limarino, C., Fauque, L., Tripaldi, A. & Net, L. 2003. Primer registro palinológico del miembro inferior de la Formación Cerro Morado (Mioceno), Precordillera de La Rioja. *Ameghiniana*, 40, 81–87.
- BECK, H.E., ZIMMERMANN, N.E., McVICAR, T.R., VERGOPOLAN, N., BERG, A. & WOOD, E.F. 2018. Present and future Köppen-Geiger climate classification maps at 1-km resolution. *Scientific Data*, 5, 1–12.
- BIRNIE, J.F. & ROBERTS, D.E. 1986. Evidence of tertiary forest in the Falkland Islands (Islas Malvinas). *Palaeogeography, Palaeoclimatology, Palaeoecology,* **55**, 45–53.
- BLISNIUK, P.M., STERN, L.A., CHAMBERLAIN, C.P., IDLEMAN, B. & ZEITLER, P.K. 2005. Climatic and ecologic changes during Miocene surface uplift in the southern Patagonian Andes. *Earth and Planetary Science Letters*, 230, 125–142.
- BROUGHTON, D.A. & McAdam, J.H. 2005. A checklist of the native vascular flora of the Falkland Islands (Islas Malvinas): new information on the species present, their ecology, status and distribution. *Journal of the Torrey Botanical Society*, 132, 115–148.
- CANTRILL, D. & POOLE, I. 2012. The vegetation of Antarctica through geological time. Cambridge: Cambridge University Press, 480 pp.
- CLARK, R. & WILSON, P. 1992. Occurrence and significance of ventifacts in the Falkland Islands, South Atlantic. Geografiska Annaler, 74, 35–46.
- COOKSON, I. 1947. Plant microfossils from the lignites of Kerguelen Archipelago. Reports (B.A.N.Z Antarctic Research Expedition (1929–1931). Series A, v. 2, Geology; pt 8, 142 pp.
- CORNOU, M.E., QUATTROCCHIO, M.E. & MARTINEZ, M.A. 2014. Palinoestratigrafía de la Formación Salto del Macho, Paleógeno de la Cuenca de Ñirihuau, Argentina. *Ameghiniana*, **51**, 566–571.
- CORNOU, M.E., MARTINEZ, M.A., QUATTROCCHIO, M.E. & ASENSIO, M.A. 2012. Estudio palinológico de esporas de la Formación Salto del Macho, Paleógeno de la Cuenca de Ñirihuau, Argentina. *Ameghiniana*, **49**, 26–37.
- ESTEBENET, M.S.G., GUERSTEIN, G.R. & RAISING, M.E.R. 2014. Middle Eocene dinoflagellate cysts from Santa Cruz Province, Argentina: biostratigraphy and paleoenvironment. *Review of Palaeobotany and Palynology*, **211**, 55–65.
- Farnsworth, A., Lunt, D.J., Robinson, S.A., Valdes, P.J., Roberts, W.H., Clift, P.D., *et al.* 2019. Past East Asian monsoon evolution controlled by paleogeography, not CO₂. *Science Advances*, **5**, eaax1697.
- Fasola, A. 1969. Estudio palinológico de la Formación Loreto (Terciario medio), provincia de Magallanes, Chile. *Ameghianian*, **6**, 3–49.
- Feitl, M., Kern, A.K., Jones, A., Fritz, S.C., Baker, P., Joeckel, R.M., et al. 2019. Paleoclimate of subtropical Andes during the latest Miocene, Lauca Basin, Chile. Palaeogeography, Palaeoclimatology, Palaeoecology, 534, 109336.
- Fernadez, D.A., Santamarina, P.E., Palazzesi, L., Telleria, M.C. & Barreda, V. 2021. Incursion of tropically-distributed plant taxa into high latitudes during the middle Eocene warming event: evidence from the Rio Turbio Fm, Santa Cruz, Argentina. *Review of Palaeobotany and Palynology*, 295, 104510.
- FICK, S.E. & HIJMANS, R.J. 2017. WorldClim 2: new 1-km spatial resolution climate surfaces for global land areas. *International Journal of Climatology*, **37**, 4302–4315.
- Guerstein, G.R., Guler, M.V. & Casadio, S. 2004. Palynostratigraphy and palaeoenvironments across the Oligocene-Miocene boundary within the Centinela Formation, southwestern Argentina. *Special Publication of the Geological Society of London*, No. 230, 325–343.
- GULER, M.V., GUERSTEIN, G.R. & QUATTROCCHIO, M.E. 2001. Palinología del Neógeno de la perforación Cx-1, Cuenca del Colorado, Argentina. *Revista española de micropaleontología*, 33, 183–204.
- Guler, M.V., Estebenet, M.S.G., Navarro, E.L., Fuentes, S., Cuitino, J.I., Palazzesi, L., et al. 2021. Miocene Atlantic transgressive-regressive events in northeastern and offshore Patagonia: a palynological perspective. *Journal of South American Earth Sciences*, 109, 103239.

- Halle, T.G. 1911. On the geological structure and history of the Falkland Islands. Uppsala: Almqvist & Wiksells, 117 pp. Reprinted from Bulletin of the Geological Institute, University of Uppsala 11, 115–229.
- HAMSON, J.D., EVANS, J.D.A., SANDERSON, D.C.W., BINGHAM, R.G. & BENTLEY, M.J. 2008. Constraining the age and formation of stone runs in the Falkland Islands using Optically Stimulated Luminescence. Geomorphology, 94, 117–130.
- HEREDIA, M.E., PAEZ, M.M., GUERSTEIN, G.R. & PARRAS, A. 2012. Palinología del miembro Gran Bajo de la Formación San Julián (Oligoceno tardío) en su localidad tipo, Santa Cruz, Argentina: consideraciones paleoambientales. *Ameghiniana*, 49, 473–496.
- HEUSSER, C.J. 1971. Pollen and spores of Chile. Tuscon, AZ: University of Arizona Press, 180 pp.
- HYLAND, E.G. & SHELDON, N.D. 2018. Paleosol-based paleoclimate reconstruction of the Paleocene-Eocene Thermal Maximum, northern Argentina. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 511, 639–642.
- IAWA COMMITTEE. 1989. IAWA list of microscopic features for hardwood identification. IAWA Bulletin, 10, 219–332.
- IAWA COMMITTEE. 2004. IAWA list of microscopic features for softwood identification. *IAWA Journal*, **25**, 1–70.
- KAPPEN, L. & STRAKA, H. 1988. Pollen and spores transport into the Antarctic. *Polar Biology*, **8**, 173–180.
- Kohn, M.J., Strömberg, C.A., Madden, R.H., Dunn, R.E., Evans, S., Palacios, A. & Carlini, A.A. 2015. Quasi-static Eocene–Oligocene climate in Patagonia promotes slow faunal evolution and mid-Cenozoic global cooling. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 435, 24–37.
- LISTER, D. & JONES, P. 2014. Long-term temperature and precipitation records from the Falkland Islands. *International Journal of Climatology*, 35, 1224–1231.
- MACPHAIL, M.K. 1979. Vegetation and climates in southern Tasmania since the last glaciation. *Quaternary Research*, 11, 10.1016/0033-5894 (79)90078-4.
- MACPHAIL, M.K. 1999. Palynostratigraphy of the Murray Basin, inland southeast Australia. *Palynology*, **23**, 199–242.
- MACPHAIL, M.K. 2021. The *Tussac House* 'forest bed' (Port Stanley, East Falkland Island). Palynostratigraphic report (unpublished).
- MACPHAIL, M. & CANTRILL, D.J. 2006. Age and implications of the Forest Bed, Falkland Islands, southwest Atlantic Ocean: evidence from fossil pollen and spores. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 240, 602–629.
- MACPHAIL, M., ALLEY, N. & TRUSWELL, E. 1994. Early Tertiary vegetation: evidence from spores and pollen. *In Hill, B., ed., History of the Australian vegetation: Cretaceous to recent.* Adelaide: University of Adelaide Press, 189–261.
- MACPHAIL, M., HILL, R.S., CARPENTER, R.J. & MCKELLAR, J. 2014. Cenozoic oil-shale deposits in southeastern-central Queensland: palynostratigraphic age determinations and correlations for the Biloela Formation (Biloela Basin) in GSQ Monto 5. Queensland Geological Record, 2014, 01.
- MARCHANT, R., ALMEIDA, L., BEHLING, H., BERRIO, J.C., BUSH, M., CLEEF, A., et al. 2002. Distribution and ecology of parent taxa of pollen lodged within the Latin American pollen data base. Review of Palaeobotany and Palynology, 121, 1–75.
- MARKGRAF, V. & D'ANTONI, H.L. 1978. *Pollen flora of Argentina*. Tucson, AZ: University of Arizona Press, 208 pp.
- MAUTINO, L.R. & ANZOTEGUI, L.M. 1998. Palinología de la Formación
 Chiquimil (Mioceno Superior), en Río Vallecito, provincia de Catamarca, Argentina. Parte 1. Esporas: espaceis nuevas.
 Ameghiniana, 39, 227–233.
- Mautino, L.R. & Anzotegui, L.M. 2002a. Palinología de la Formación Chiquimil (Mioceno Superior), en Río Vallecito, provincia de Catamarca, Argentina. Parte 2. Polen. *Ameghiniana*, **39**, 257–270.

- MAUTINO, L.R. & ANZOTEGUI, L.M. 2002b. Palinología de la Formación Chiquimil (Mioceno superior) Vallecito, provincia de Catamarca. Parte 3. Polen. *Ameghiniana*, **39**, 271–284.
- McDowell, R.M. 2005. Falkland Islands biogeography: converging trajectories in the South Atlantic Ocean. *Journal of Biogeography*, 32, 49–62.
- MONTADE, V., NEBOUT, N.C., KISSEL, C. & MULSOW, S. 2011. Pollen distribution in marine surface sediments from Chilean Patagonia. *Marine Geology*, 282, 161–168.
- Nanez, C., Quattrochio, M.E. & Ruiz, L. 2009. Palinología y micropaleontología de las Formaciones San Julián y Monte León (Oligoceno Mioceno temprano) en el subsuelo de cabo Curioso, provincia de Santa Cruz, Argentina. *Ameghiniana*, **46**, 669–693.
- OLIVERO, E.B., BARREDA, V., MARENSSI, S.A., SANTILLANA, S.N. & MARTINIONI, D.R. 1998. Estratigrafía sedimentología y palinología de la Formatión Sloggett (Paleogeno continental), Tierra del Fuego. *Revista de la Associacion Geologica, Argentina*, 53, 504–516.
- Ortiz-Jaureguizar, E. & Cladera, G.A. 2006. Paleoenvironmental evolution of southern South America during the Cenozoic. *Journal of Arid Environments*, **66**, 498–532.
- Ottone, E.C., Reinarte Mazurier, S.M.E. & Salinas A. 2013. Palinomorphos Miocenos del Subsuelo de Santiago del Estero Argentina. *Ameghiniana*, **50**, 509–521.
- Palamarczuk, S. & Barreda, V. 2000. Palinología del Paleogeno tardío-Neo geno temprano, pozo Aries x-l, Plataforma Continental Argentina, Tierra del Fuego. *Ameghiniana*, 37, 221–234.
- PALAZZESI, L. & BARREDA, V. 2004. Primer registro palinológico de la Formación Puerto Madryn, Mioceno de la provincia del Chubut, Argentina. Ameghiniana, 41, 355–362.
- PALAZZESI, L. & BARREDA, V. 2007. Major vegetation trends in the Tertiary of Patagonia (Argentina): a qualitative paleoclimatic approach based on palynological evidence. Flora - Morphology, Distribution, Functional Ecology of Plants, 202, 328–337.
- PALAZZESI, L. & BARREDA, V. 2021. Major vegetation trends in the Tertiary of Patagonia (Argentina): a qualitative paleoclimatic approach based on palynological evidence. Flora - Morphology, Distribution, Functional Ecology of Plants, 202, 328–337.
- PALAZZESI, L., VIZCAÍNO, S.F., BARREDA, V.D., CUITIÑO, J.I., DEL RÍO, C.J., GOIN, F., et al.. 2021. Reconstructing Cenozoic Patagonian biotas using multi-proxy fossil records, *Journal of South American Earth Sciences*, 112, 103513.
- Parras, A. & Cuttino, J.I. 2021. Revised chrono and lithostratigraphy of the Oligocene-Miocene *Patagoniense* marine deposits in Patagonia: implications for stratigraphic cycles, palaeogeography, and major drivers. *Journal of South American Earth Sciences*, 110, 103327
- Parras, A., Guerstein, G.R., Perez Panera, J.P., Griffin, M., Nanez, C., Cusminsky, G. & Quiroga, A. 2020. Integrated stratigraphy and paleontology of the lower Miocene Monte Leon Formation, southeastern Patagonia, Argentina: unravelling palaeoenvironmental changes and factors controlling sedimentation. *Palaeogeography, Palaeoeclimatology, Palaeoecology*, **556**, 109701.
- Partridge, A.D. 1999. Late Cretaceous to Tertiary geological evolution of the Gippsland Basin, Victoria, 2 vols. Unpublished PhD thesis. Melbourne: Latrobe University, 439 pp.
- POOLE, I. & CANTRILL, D. 2007. The arboreal component of the Neogene forest bed, West Point Island, Falkland Islands. *IAWA Journal*, 28, 423–444.
- QUATTROCCHIO, M.E. & VOLKHEIMER, W. 1990. Paleogene environmental trends as reflected in palynological types, Salta Basin, NWArgentina. *Neues Jahrbuch für Geologie und Paläontologie*, **181**, 377–396.
- QUATTROCCHIO, M.E., DE CABRERA, J. & GALLI, C. 2003. Formación Anta (Mioceno Temprano/Medio), Subgrupo Metán (Grupo Orán), en el río Piedras, Pcia. de Salta: Datos palinológicos. Revista de la Associación Geologica, Argentina, 58, 117–127.

- QUATTROCCHIO, M.E., VOLKHEIMER, W., BORROMEI, A.M. & MARTINEZ, M.A. 2011. Changes of the palynobiotas in the Mesozoic and Cenozoic of Patagonia: a review. *Biological Journal of the Linnean Society*, **103**, 380–396.
- RAINE, J.I., MILDENHALL, D.C. & KENNEDY, E. 2011. New Zealand fossil spores and pollen: an illustrated catalogue. Lower Hutt: GNS Science, vol. 4.
- RENNER, S. 2004. Plant dispersal across the tropical Atlantic by wind and sea currents. *International Journal of Plant Sciences*, 165, S23–S33.
- ROMERO, J. & ZAMALOA, M.C. 1985. Polen de angiospermas de la Formación Río Turbio (Eoceno), Provinca de Santa Cruz, República Argentina. *Ameghiniana*, **22**, 43–51.
- ROSENBAUM, M. 1996. Stone runs in the Falkland Islands. *Geology Today*, 12, 151–154.
- SACHSE, V.F., STROZYK, F., ANKA, Z., RODRIGUEZ, J.F. & DI PRIMO, R. 2015. Tectono-stratigraphic evolution of the Austral Basin and adjacent areas of Andean tectonics, southern Argentina, South America. *Basin Research*, 28, 462–482.
- Seward, A.C. & Conway, V. 1934. A phytogeographical problem: fossil plants from the Kerguelen. *Annals of Botany*, **48**, 715–741.
- Srivastava, G., Bhatia, H., Verma, P., Singh, Y.P., Utescher, T. & Mehrotra, R.C. 2023. High rainfall afforded resilience to tropical rainforests during Early Eocene Climatic Optimum. *Palaeogeography, Palaeoclimatology, Palaeoecology*, **628**, 111762.
- STANCA, R.M., McCarthy, D.J., Paton, D.A., Hodgson, D.M. & Mortimer, E.J. 2022. The tectono-stratigraphic architecture of the Falkland Plateau basin; implications for the evolution of the Falkland Islands Microplate. *Gondwana Research*, **105**, 320–342.
- STONE, P., ALDISS, D.T. & EDWARDS, E.J. 2005. Rocks and fossils of the Falkland Islands. Nottingham: British Geological Survey, 60 pp.
- THOMAS, Z.A., JONES, R.T., FOGWILL, C.J., HATTON, J., WILLIAMS, A., HOGG, A.G., *et al.* 2018. Evidence for increased expression of the Amundsen Sea Low over the South Atlantic during the late Holocene. *Climate of the Past*, **14**, 1727–1738.
- Traverse, A. 2007. Production, dispersal, sedimentation and taphonomy of spores/pollen in relation to the interpretation of palynofloras. *In* Traverse, A., *Paleopalynology*. Berlin: Springer, 497–542.
- TRONOSCO, A. & ENCINAS, A. 2006. La tafoflora de cerro Centinela (Chile, VI Región): vegetación y clima de Chile central a fines del Miocenocomienzos del Plioceno. *Ameghiniana*, 43, 171–180.
- TRUSWELL, E.M., QUILTY, P.G., McMINN, A., MACPHAIL, M.K. & WHELLER, G.E. 2005. Late Miocene vegetation and

- palaeoenvironments of the Drygalski Formation, Heard Island, Indian Ocean: evidence from palynology. *Antarctic Science*, **17**, 427–442.
- Turney, C.S.M., Becerra-Valdivia, L., Sookdeo, A., Thomas, Z.A., Palmer, J., Haines, H.A., *et al.* 2021. Radiocarbon protocols and first intercomparison results from the Chronos ¹⁴Carbon-Cycle Facility, University of New South Wales, Sydney, Australia. *Radiocarbon*, **63**, 1003–1023.
- Turney, C.S.M., Jones, R., Fogwill, C., Hatton, J., Williams, A.N., Hogg, A., *et al.* 2016. A 250 year periodicity in Southern Hemisphere westerly winds over the last 2600 years. *Climate of the Past*, **12**, 189–200.
- VEBLEN, T.T., DONOSO, C., KITZBERGER, T. & REBERTUS, A.J., 1996.
 Ecology of southern Chilean and Argentinean Nothofagus forests. In
 VEBLEN, T., HILL, R. & READ, J., eds, The ecology and biogeography of Nothofagus forests. New Haven, CT: Yale University Press, 93–353.
- VIZCAÍNO, S.F., BARGO, M.S. & FERNÍCOLA, J.C. 2013. Expediciones paleontológicas durante los Siglos XIX y XX a la Formación Santa Cruz (Mioceno Inferior, Patagonia) y destino de los fósiles. Presented at III Congreso Argentino de Historia de la Geología (CAHGEO), Salta, 29–31 August 2013.
- WARNY, S., KYMES, C.M., ASKIN, R.A., KRAJEWSKI, K.P. & BART, P.J. 2016. Remnants of Antarctic vegetation on King George Island during the early Miocene Melville Glaciation. *Palynology*, 40, 66–82.
- WARNY, S., KYMES, C.M., ASKIN, R.A., KRAJEWSKI, K.P. & TATUI, A. 2019. Terrestrial and marine floral response to latest Eocene and Oligocene events on the Antarctic Peninsula. *Palynology*, **43**, 4–21.
- WOOD, G.D. 1996. Palynological techniques-processing and microscopy. In JASONIUS, J. & McGregor, D.C., eds, Palynology: principles and application. American Association of Stratigraphic Palynologists Foundation, vol. 1, 29–50.
- ZACHOS, J.C., DICKENS, G.R. & ZEEBE, R.E. 2008. An Early Cenozoic perspective on greenhouse warming and carbon-cycle dynamics. *Nature*. 451, 279–283.
- ZACHOS, J.C., PAGANI, M., SLOAN, L., THOMAS, E. & BILLUPS, K. 2001. Trends, rhythms, and aberrations in global climate 65 Ma to present. *Science*, 292, 686–693.
- Zamaloa, M.D.C. & Romero, E.J. 2005. Neogene palynology of tierra del fuego, Argentina: conifers. *Alcheringa*, **29**, 113–121.
- ZWIER, M., VAN DER BILT, W.G., DE STIGTER, H. & BJUNE, A.E. 2022. Pollen evidence of variations in Holocene climate and Southern Hemisphere westerly wind strength on sub-Antarctic South Georgia. The Holocene, 32, 147–158.