

PALYNOSTRATIGRAPHY OF THE CRETACEOUS–PALEOGENE IN THE AUSTRAL BASIN, SW SANTA CRUZ PROVINCE, ARGENTINA

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ABSTRACT – This paper analyses the palynological assemblages recovered from the Cerro Cazador and Monte Chico formations and is an exhaustive study of the spore-pollen assemblages from the Cerro Dorotea Formation; all units with outcrops out in southwestern Santa Cruz Province, Argentina. Six sections were sampled in two main areas: Estancia San José and Estancia Laguna Salada. The palynological assemblages yielded by the studied formations are integrated with elements of marine (dinoflagellate cysts and acritarchs) and continental (spores and pollen grains) origin, present in different proportions throughout these units. Forty one species of pollen grains and spores were recognized in the Cerro Cazador Formation, 74 genera and 127 species in the Monte Chico Formation and 64 genera and 107 species in the Cerro Dorotea Formation. On the basis of the stratigraphic distribution of the identified species, four palynological assemblages were recognized, which were defined by the exclusive presence of characteristic species and their similarities with other spore-pollen assemblages. The following ages were suggested: (i) Association 1, upper sections of the Cerro Cazador Formation: upper Campanian–lower Maastrichtian; (ii) Association 2, lower and middle levels of the Monte Chico Formation: Maastrichtian, probably upper Maastrichtian; (iii) Association 3, upper levels of the Monte Chico Formation: Maastrichtian–Danian; and (iv) Association 4, Cerro Dorotea Formation: Danian. Based on this analysis, the K/P boundary is located between Associations 2 and 3, within the Monte Chico Formation. These palynological assemblages indicate a near-shore marine depositional environment close to the coastline, with marginal conditions, and a progressive shallowing of the basin.

Key words: palynostratigraphy, pollen grains, spores, Cretaceous, Paleogene, Santa Cruz Province.

RESUMO – Nesta contribuição são apresentadas as associações palinológicas das formações Cerro Cazador e Monte Chico, associações esporopolínicas da Formação Cerro Dorotea, no sudoeste da Província de Santa Cruz, Argentina. Seis seções foram amostradas em duas áreas principais: Estancia San José e Estancia Laguna Salada. As associações palinológicas são constituídas por elementos de origem marinha (cistos de dinoflagelados e acritarcas) e continental (grãos de pólen e esporos), presentes em diferentes proporções ao longo destas unidades. Quarenta e uma espécies de grãos de pólen e de esporos foram identificadas na Formação Cerro Cazador, 74 gêneros e 127 espécies na Formação Monte Chico e 64 gêneros e 107 espécies na Formação Cerro Dorotea. Com base na distribuição estratigráfica das espécies identificadas, e semelhanças com outras associações esporopolínicas, foram reconhecidas quatro associações palinológicas, com as seguintes idades: (i) Associação 1, secção superior da Formação Cerro Cazador: Campaniano superior–Maastrichtiano inferior; (ii) Associação 2, níveis mais baixos e médios da Formação Monte Chico: Maastrichtiano, provavelmente Maastrichtiano superior; (iii) Associação 3, níveis superiores da Formação Monte Chico: Maastrichtiano–Daniano e (iv) Associação 4, Formação Cerro Dorotea: Daniano. Com base nesta análise, o limite K/P localiza-se entre Associações 2 e 3, dentro da Formação Monte Chico. Estas associações palinológicas sugerem paleoambiente de deposição marinho proximal, em condições marginais e continentalização progressiva da bacia.

Palavras-chaves: palinoestratigrafia, grãos de pólen, esporos, Cretáceo, Paleógeno, Província de Santa Cruz.

INTRODUCTION

In this contribution the spore-pollen associations of the Cerro Cazador, Monte Chico and Cerro Dorotea formations cropping out in the Estancia San José and Estancia Laguna Salada areas are studied. The study area is located at the most southwestern extreme of the Province of Santa Cruz (Figure 1). Palynological sampling was conducted in several sections exposed in outcrops near the area of Cancha Carrera ($51^{\circ}11'20.2''S$, $72^{\circ}20'55.5''W$), and in the area of Cerro de la Cruz, near Rio Turbio City ($51^{\circ}33'00.5''S$, $72^{\circ}25'43.2''W$), forming an integrated profile of the entire analyzed sequence (Figures 2, 3).

GEOLOGICAL FRAMEWORK

The Cerro Cazador Formation consists of fine to medium-grained sandstones with interbedded calcarenites, in part glauconitic, with claystones, conglomerates and limestone banks with pelecypod remains and gastropods (Riccardi & Rolleri, 1980).

This formation is restricted to the sequence of “green sandstones” of Hauthal (1898), the *Lahillia luisa* Layers of Wilckens (1907), the lower levels of the “middle section of the Green Sandstone” of Brandmayr (1945), the sediments that Feruglio (1938, 1949) recognized as the “strata of Monte Cazador” or “*Lahillia luisa* Layers”, the basal section and

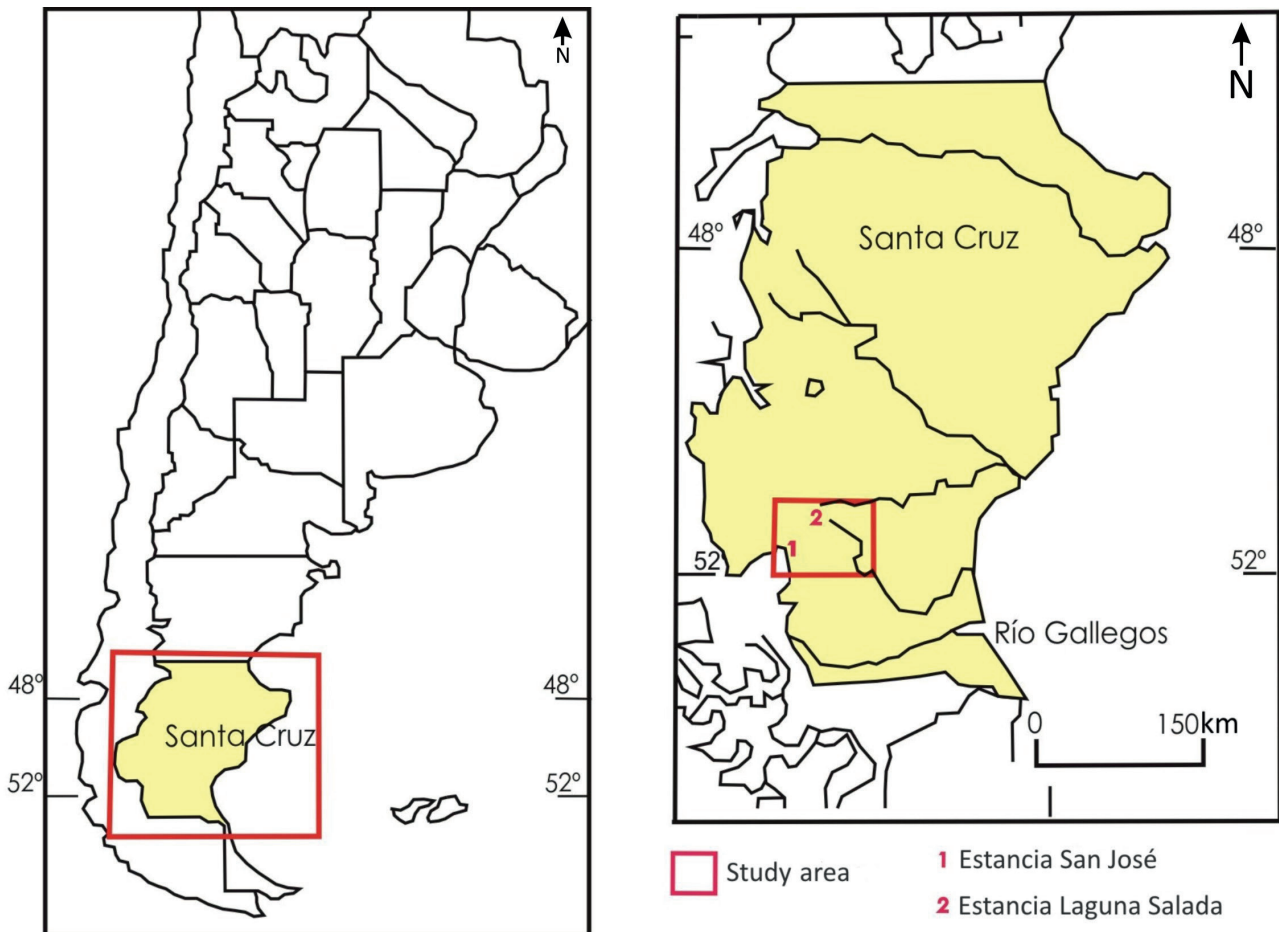


Figure 1. Location map of the study area.

the lower part of the middle of the “Cerro Cazador Strata or *Lahillia luisa* Layers” of Hünicken (1955), the Cerro Cazador Group of Borrello (1956), the Dorotea Formation of Katz (1963) and the Cerro Cazador Formation of Leanza (1972) (Table 1).

This unit conformably overlies the Cerro Toro Formation and is unconformably overlain by the Monte Chico Formation, so is assigned to the Campanian–Maastrichtian on the basis of its stratigraphic relationships (Malumián & Panza, 1996).

The Monte Chico Formation is composed of fine to medium-grained sandstone, brown, light brown to reddish and greenish gray, partly calcareous, with abundant dark gray concretions including invertebrate megafauna fossils, mudstone, limonite and conglomerates interbedded with coquinoid megafossil beds (Malumián & Panza, 1996) (Figure 3). The Monte Chico Formation corresponds to the upper levels of the “Middle Section of the Green Sandstone” of Brandmayr (1945), the strata of the Cerro Cazador or *Lahillia luisa* Layers of Feruglio (1938, 1949) and Hünicken (1955), the Cerro Cazador Formation of Leanza (1972) and the lower levels of the Dorotea Formation of Katz (1963). This paper follows the stratigraphic outline presented by Malumián & Panza (1996), who gave the formal description of the Monte Chico Formation, supporting the discordant relationship of the base of this unit, which shows a transgressive base overlying

sedimentary units of different ages. The upper boundary is concordant and gradual with the Cerro Dorotea Formation. The Monte Chico Formation correlates with the Calafate Formation; both are of Cretaceous age, the same lithology and show similar stratigraphic relationships.

The Cerro Dorotea Formation corresponds to the “strata of the Cerro Dorotea” of Feruglio (1938, 1949), and the “upper section of the Green Sandstone” or strata with *Ostrea rionegrensis* of Brandmayr (1945). The Dorotea Formation of Cecioni (*in Hoffstetter et al.*, 1957) or the Sierra Dorotea Group of Borrello (1956) do not include the same sequence as the Cerro Dorotea Formation as defined here, because the former included the strata of Monte Grande, the Cerro Cazador Layers and the Cerro Dorotea Layers since Cecioni considered it impossible to distinguish between these strata lithologically. Borrello (1956) divided the Sierra Dorotea Group into three sections: lower, middle and upper. The lower section corresponds to the Cerro Dorotea Formation (Figure 2). Some pelitic levels and coal strata have a gradual transitional relationship over the strata of the Monte Chico Formation (Malumián & Panza, 1996). To the SE of Estancia Laguna Salada and after about 500 meters of vegetation-covered rocks, there are outcrops of the top of the Cerro Dorotea Formation, consisting of shales, sandstones and conglomerate packages.

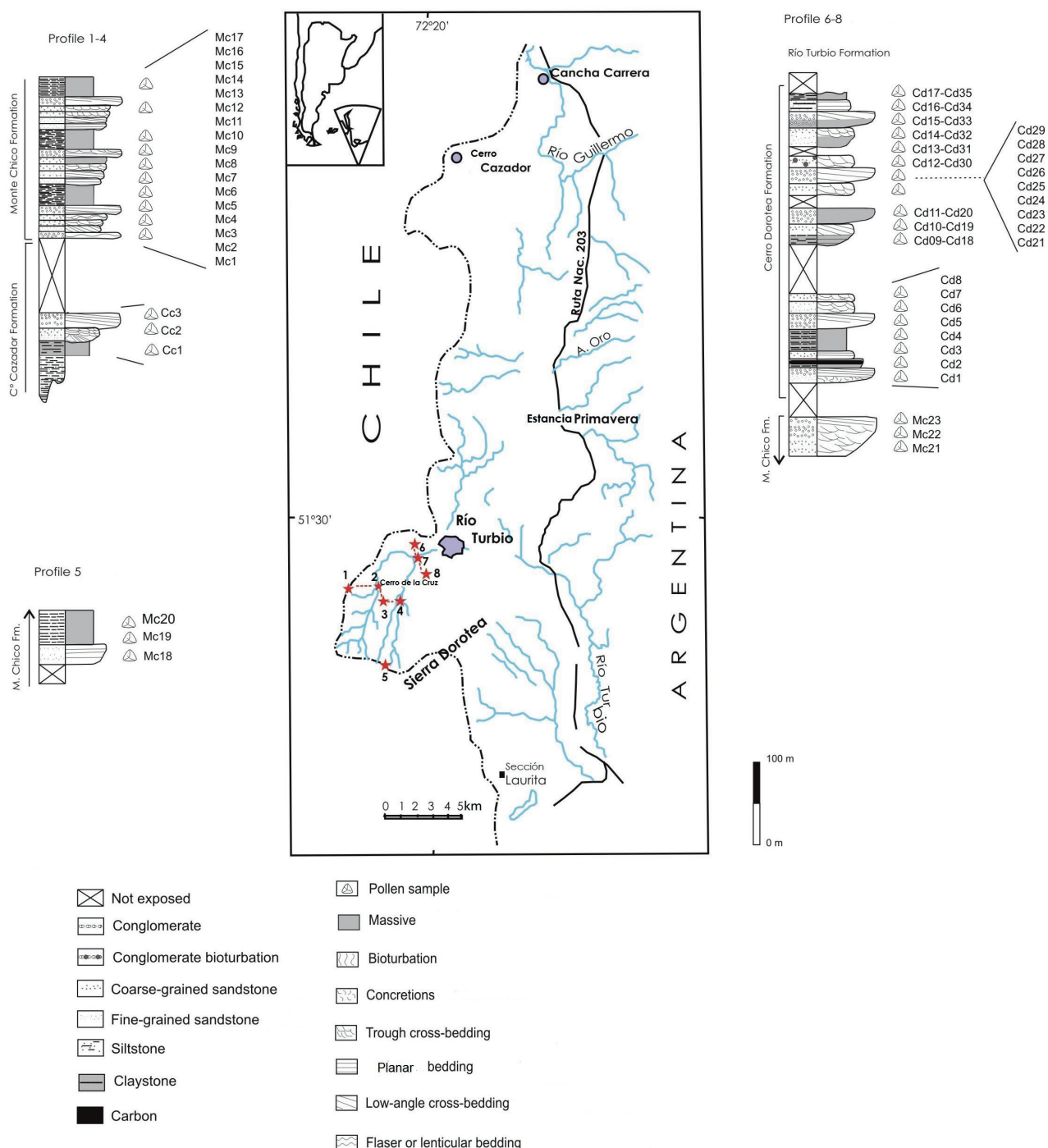


Figure 2. Stratigraphic section in Cerro de la Cruz area, near the city of Rio Turbio, Santa Cruz Province, Argentina.

The objective of this paper is to present the palynoflora from the Cerro Cazador, Monte Chico and Cerro Dorotea formations, with the aim of identifying one or several areas with restricted distributions of taxa, capable of being useful as a guide, to infer the relative age and to define the depositional paleoenvironment. All profiles were made in a west–east direction, perpendicular to the strike of the strata that has an inclination of 5° to 25° east. The exposed banks are conglomerate-bearing silicified masses, representing

a period of greater energy input, with related sections of pelitic and coarsening-upward sandstone sequences. It represents a marine depositional environment with little internal communication between the platform and the open sea (Malumíán & Panza, 1996). A preliminary palynological analysis of Cretaceous sequences near the study area (Povilauskas *et al.*, 2006; Povilauskas & Guler, 2008) suggested a coastal marine depositional environment close to the coastline, and an age of Maastrichtian–late Maastrichtian.

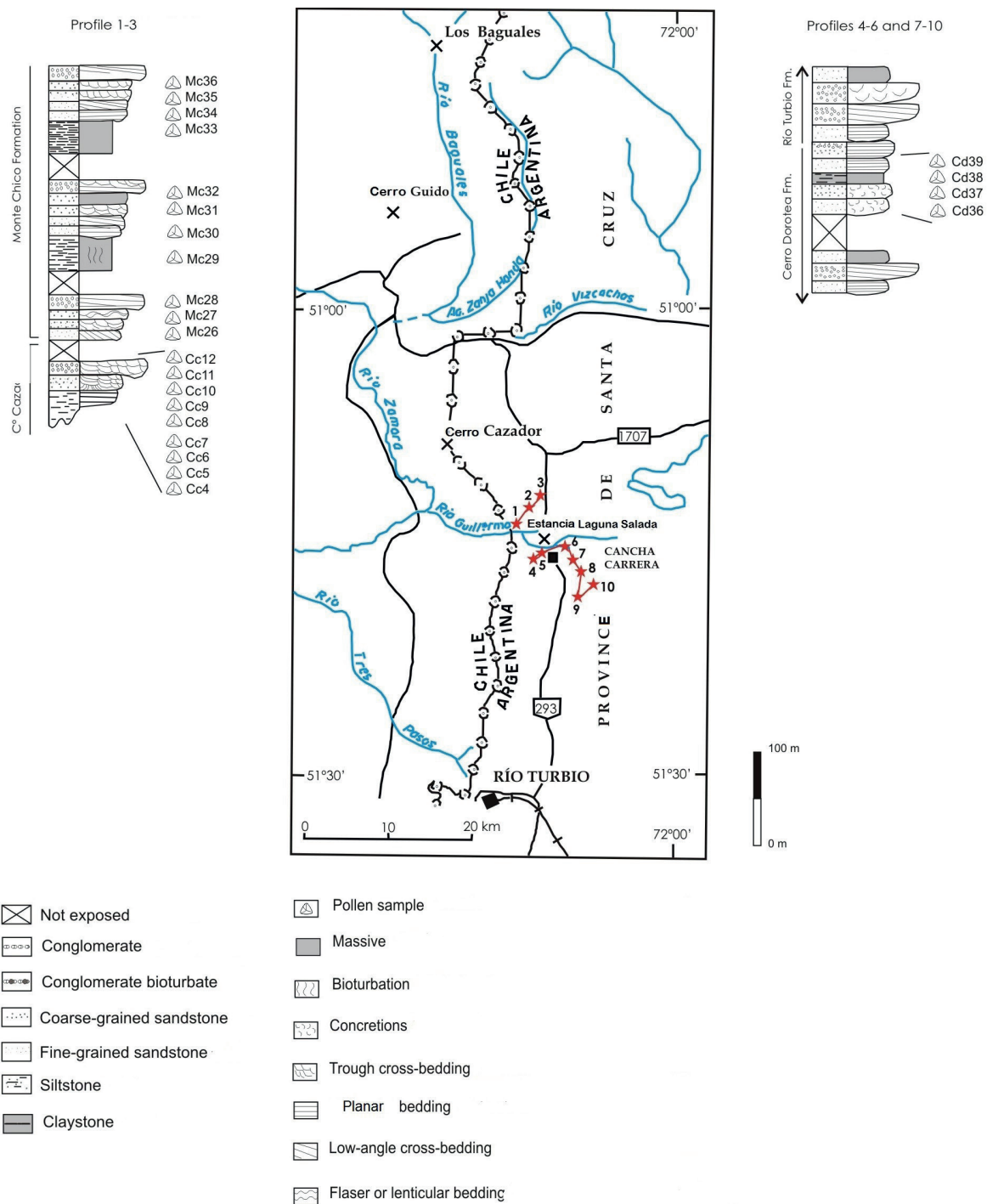


Figure 3. Stratigraphic section in Estancia Laguna Salada area, near of Cancha Carrera, Santa Cruz Province, Argentina.

MATERIAL AND METHODS

For the collection of the samples the best areas of exposure were chosen between the two areas of study (Estancia San José and Estancia Laguna Salada) (Figures 2, 3). A difficulty with sampling was that in several sections of interest a considerable percentage of the profiles were covered by vegetation, leaving only relatively small stratigraphic intervals for extraction

of samples. The distance between samples was irregular, depending on the lithology and the sections covered. At appropriate intervals sampling was conducted at a distance of 5 m. The collected material was bagged for subsequent laboratory analysis.

Palynological extraction was performed according to the conventional methods of physical and chemical extraction (HCl-HF) (Volkheimer & Melendi, 1976) and the residue

obtained was filtered through mesh (+10 and +25 microns). The final preparations were mounted in glycerine gelatin. The specimens were studied with a Leitz Wetzlar microscope and Olympus BX51 microscope (Germany/Japan) in the Paleopalynology Section and photographs were taken with a transparent optical microscope and scanning electron microscope (SEM) at the Argentinian Museum of Natural Sciences “Bernardino Rivadavia”. For photo documentation of the specimens a Nikon E4500 digital camera was used. Classification is considered semi-natural and identifies taxa to genus and species. The terminology used follows Punt *et al.* (2007).

The preparations are reposed in the Regional Provincial Museum “Padre Manuel Jesús Molina” in Rio Gallegos, Santa Cruz Province, Argentina, under the prefix MPM-MP with catalog numbers 1943 to 1978.

PALYNOSTRATIGRAPHY

On the basis of this analysis four spore-pollen associations characterized by groups of species with restricted distributions were recognized. Figure 4 shows the distribution of the species identified in the formations studied. In the associations groups of species common to the three formations were also identified, which, if they are not useful for the purposes of a biozone characterization, are useful for characterizing the whole association.

Among them *Arecipites minutiscabratus*, *Baculatisporites comaumensis*, *Clavifera triplex*, *Cyatheacidites annulatus*, *Gleicheniidites senonicus*, *Liliacidites kaitangataensis*, *Nothofagidites saraensis* and *Polypodiidites speciosus* are the most significant.

The four spore-pollen associations have characteristics that allow them to be clearly distinguished from each other. The transitions between these associations appear to be gradual. The changes were observed in both taxonomic composition and in the relative abundance of the palynomorphs.

Assemblage 1. Is characterized by the exclusive presence of *Baculatisporites* cf. *B. comaumensis*, *Biretisporites* cf. *B. potoniaei*, *Ischyosporites* sp. 1, *Trilites* cf. *T. fasolae*, *Verrucosisporites* sp. 2, *Podocarpidites* sp. 2 and *Triporopollenites* sp. 1, with the absence of the distinctive features of higher associations (from the Monte Chico and Cerro Dorotea formations). This association comes from the upper levels of the Cerro Cazador Formation (see appendix 1).

Assemblage 2. Is characterized by the exclusive presence of *Beupreadites elegansiformis*, *Camarozonosporites ohaiensis*, *Ceratosporites equalis*, *Forcipites sabulosus*, *Haloragacidites prioratus*, *Ilexpollenites salamanquensis*, *Liliacidites vermiculatus*, *Longapertites patagonicus*, *Ornamentifera echinata*, *Proteacidites subscabrous*, *Quadruplicatus brossus*, *Rhoipites baculatus*, *Rousea microreticulata*, *Senipites tercassata*, *Tricolpites bibaculatus* and *Tuberculatosporites parvus* and the absence of characteristic elements of the lower (1) and upper (3, 4) associations. Also in this association are the first records of *Baculatisporites turboensis*, *Biretisporites crassilabratus*,

Classopollis sp. 1, *Nothofagidites kaitangataensis*, *Peninsulapollis truswellidae*, *Periporopollenites demarcates*, *Peromonolites vellosus*, *Proteacidites beddoesii*, *Proteacidites tenuixinus*, *Psilatricolpites patagonicus*, *Psilatricolporites* cf. *P. salamanquensis*, *Rhoipites minusculus*, *Rousea patagonica*, *Sparganiaceapollenites barungensis*, *Spinizonocolpites hialinus*, *Triatriopollenites bertelsii*, *Trilites tuberculiformis* and *Triporopollenites* cf. *T. ambiguus*. This association comes from the basal levels and middle of the Monte Chico Formation (see appendix 1).

Assemblage 3. Is characterized by the exclusive presence of *Beupreadites* sp. 1, *Liliacidites* sp. 1, *Peninsulapollis askinia* and *Pseudowinterapollis couperi* and the absence of distinctive elements of associations 1, 2 and 4. This association comes from the upper levels of the Monte Chico Formation. In this association are the first records of *Ericipites scabratus*, *Gamerroites psilasaccus*, *Nothofagidites dorotensis* and *Nothofagidites nana* (see appendix 1).

Assemblage 4. Is characterized by the exclusive presence of *Bombacacidites* sp. 1, *Forcipites stipulatus*, *Nothofagidites waipawaensis*, *Propylipollis microverrucatus* and *Tetracolporites* sp. 1, and the absence of the distinctive elements of the lower associations (1, 2 and 3). This association comes from the Cerro Dorotea Formation (see Appendix 1).

AGE OF ASSOCIATIONS

The age of the associations identified is inferred on the basis of: (i) known temporal ranges of the species present in each of the associations, and (ii) the similarities to other previously studied palynological associations, especially those from Campanian, Maastrichtian and Paleocene sequences of Argentina and Antarctica (Table 2). The associations that were used for comparison in this analysis were the Pedro Luro Formation, Maastrichtian–Danian, Buenos Aires Province (Ruiz & Quattroccio, 1997); the Loncoche Formation, Maastrichtian, Mendoza Province (Papú, 2002); the Los Alamitos Formation, upper Campanian, Río Negro Province (Papú & Sepúlveda, 1995); the Paso del Sapo Formation, Maastrichtian, Chubut Province (Papú, 1988a,b, 1989); the Lefipán Formation, Maastrichtian, Chubut Province (Baldoni, 1992; Baldoni & Askin, 1993); the Salamanca Formation, Danian, Chubut Province (Archangelsky, 1973; Archangelsky & Zamalloa, 1986); the Lopez de Bertodano Formation, Maastrichtian–Danian, Antarctic Peninsula (Baldoni & Barreda, 1986; Askin, 1988, 1990) and the La Irene Formation, Maastrichtian–Danian, Santa Cruz Province (Povilauskas *et al.*, 2008).

Assemblage 1

Species exclusive to this association have greater stratigraphic significance as they have been identified in other basins in Argentina within a bounded time range. Only the records of *Nothofagidites saraensis* and *Peninsulapollis gillii*, also present in higher associations (from the Monte Chico Formation), constrain the maximum age limit of the

Table 1. Scheme of correlation of the units studied according to different authors.

	Hauthal, 1898	Wilkens, 1907	Brandmayr, 1945	Feruglio, 1949	Hünicken, 1955	Borrello, 1956	Leanza, 1972	Malumán & Panza, 1996
3º Green sandstones		Layers with <i>Laillia luisa</i>	Upper section of Green sandstones with <i>Ostrea riogrensis</i>	Strata of the Cerro Dorotea or Layers with <i>O. riogrensis</i>	Strata of the Cerro Dorotea or Layers with <i>O. riogrensis</i>	Cerro Dorotea Group-Lower sect	Dorotea Formation	Cerro Dorotea Formation
					↗ Upper sect ↗ Lower sect			Monte Chico Formation
Middle section of Green sandstones with <i>L. luisa</i>				Strata of Monte Cazador or Layers with <i>L. luisa</i>	Strata of the Cerro Cazador or Layers with Middle sect	Cerro Cazador Group	Cerro Cazador Formation	Cerro Cazador Formation
					→ Middle sect → Lower sect			
Lower section of Green sandstones with <i>Ostrea riogrensis</i>				Strata of Monte Grande	<i>L. luisa</i>			

2º Conglomerate								
1º Archeschistosis with <i>I. steinmanni</i>			Strata with <i>Inoceramus</i> <i>steinmanni</i>					

Table 2. Number of species identified in common with the four associations studied and their similarities to other sequences of Argentina and Antarctica.

Association	Species in common	Similarities with other sequences of Argentina and Antarctica
1	19	Los Alamitos Formation, Río Negro Province (upper Campanian) (Papú & Sepúlveda, 1995)
2	35	Lefipán Formation, Chubut Province (Upper Cretaceous) (Baldoni, 1992; Baldoni & Askin, 1993)
3	30	La Irene Formation Santa Cruz Province (upper Campanian–lower Maastrichtian) (Povilauskas <i>et al.</i> , 2008)
4	36	Paso del Sapo and Lefipán formations, Chubut Province (upper Maastrichtian–lowermost Danian) (Papú, 1988; 1990; Baldoni, 1992; Baldoni & Askin, 1993)
	22	Salamanca Formation, Chubut Province (Paleocene) (Archangelsky, 1973)

Figure 4. Stratigraphic distribution of selected species and recognized in the Cerro Cazador, Monte Chico and Cerro Dorotea formations.

Selected species	Cerro Cazador Fm.	Monte Chico Formation			Cerro Dorotea Fm.
	Association 1	Association 2	Association 3	Association 4	
<i>Baculatisporites cf. B. comaumensis</i>					
<i>Biretisporites cf. B. potonaei</i>					
<i>Ischyosporites sp. 1</i>					
<i>Trilites cf. Trilites fasolae</i>					
<i>Verrucosporites sp. 2</i>					
<i>Podocarpidites sp. 2</i>					
<i>Triplopollenites sp. 1</i>					
<i>Rousea microreticulata</i>					
<i>Neorastrickia sp. 1</i>					
<i>Reticulodisporites tenellus</i>					
<i>Liliacidites cf. L. regularis</i>					
<i>Liliacidites variegatus</i>					
<i>Forcipites sp. "A"</i>					
<i>Polypodiidites speciosus</i>					
<i>Baculatisporites comaumensis</i>					
<i>Clavifera triplex</i>					
<i>Cyatheacidites annulatus</i>					
<i>Gleicheniidites senonicus</i>					
<i>Ischyosporites volkheimeri</i>					
<i>Araucariacites australis</i>					
<i>Lygistepollenites florinii</i>					
<i>Microcachryidites antarcticus</i>					
<i>Phyllocladidites mawsonii</i>					
<i>Podocarpidites elegans</i>					
<i>Arecipites minutiscabrus</i>					
<i>Liliacidites kaitangataensis</i>					
<i>Nothofagidites saraensis</i>					
<i>Peninsulapollis gillii</i>					
<i>Psilatricolporites sp. 1</i>					
<i>Triatriopollenites lateflexus</i>					
<i>Tricolpites reticulatus</i>					
<i>Longapertites patagonicus</i>					
<i>Forcipites sabulosus</i>					
<i>Ceratosporites equalis</i>					
<i>Proteacidites subscabrus</i>					
<i>Senipites tercassata</i>					
<i>Ornamentifera echinata</i>					
<i>Tuberculatosporites parvus</i>					
<i>Beupreadidites elegansiformis</i>					
<i>Haloragacidites trioratus</i>					
<i>Ilexpollenites salamanquensis</i>					
<i>Liliacidites vermireticulatus</i>					
<i>Rousea microreticulata</i>					
<i>Tricolpites bibaculatus</i>					
<i>Quadruplicatus brossus</i>					
<i>Camarozonosporites ohaiensis</i>					
<i>Rhoipites baculatus</i>					
<i>Baculatisporites turboensis</i>					
<i>Peromonolites vellosus</i>					
<i>Rhoipites minusculus</i>					
<i>Psilatricolpites patagonicus</i>					
<i>Spinizonocolpites hialinus</i>					
<i>Classopolis sp. 1</i>					
<i>Biretisporites crassilabrus</i>					
<i>Triatriopollenites bertelsii</i>					
<i>Trilites tuberculiformis</i>					
<i>Sparganiaceapollenites barungensis</i>					
<i>Psilatricolporites cf. P. salamanquensis</i>					
<i>Proteacidites beddoesi</i>					
<i>Nothofagidites kaitangataensis</i>					
<i>Peninsulapollis truswelliae</i>					
<i>Proteacidites tenuixenus</i>					
<i>Rousea patagonica</i>					
<i>Triplopollenites cf. T. ambiguus</i>					
<i>Periporopollenites demarcatus</i>					
<i>Peninsulapollis askiniae</i>					
<i>Beupreadidites sp. 1</i>					
<i>Pseudowinterapollis couperi</i>					
<i>Liliacidites sp. 1</i>					
<i>Ericipites scabrus</i>					
<i>Gamerroites psilasaccus</i>					
<i>Nothofagidites dorotensis</i>					
<i>Nothofagidites nana</i>					
<i>Tetracolporites sp. 1</i>					
<i>Forcipites stipulatus</i>					
<i>Propylipollis microverrucatus</i>					
<i>Bombacacidites sp. 1</i>					
<i>Nothofagidites waipawaensis</i>					

association. These two species do not have records prior to the late Campanian in Patagonia and Antarctica or Australia (Baldoni & Barreda, 1986; Askin, 1988, 1990; Papú, 1990; Baldoni & Askin, 1993; Dettmann & Thompson, 1987; Dettmann & Jarzen, 1988).

The greatest similarities of Association 1 (Cerro Cazador Formation) occur with an association from the Los Alamitos Formation, upper Campanian, Río Negro Province (Papú & Sepúlveda, 1995), which shares many of the species present, such as *Araucariacites australis*, *Clavifera triplex*, *Cyatheacidites annulatus*, *Cyathidites minor*, *Gleicheniidites senonicus*, *Liliacidites* spp., *Microcachryidites antarcticus*, *Neoraistrickia* sp., *Peninsulapollis gillii*, *Podocarpidites* spp., *Stereisporites antiquasporites*, *Tricolpites reticulatus* and freshwater algae such as *Botryococcus* sp., among others (Table 3). Other recovered associations from Patagonia, from the Loncoche Formation, Maastrichtian of Mendoza Province (Papú, 2002), also have significant numbers of common species: 14 species. Antarctic associations, however, dominated by Nothofagaceae and Podocarpaceae (Baldoni & Barreda, 1986; Dettmann & Thomson, 1987; Askin, 1990), have low overall similarities. In the Cerro Cazador Formation Nothofagaceae (*Nothofagidites saraensis*) are recognized but in very low proportions.

Also dinoflagellate cysts present in the lower levels of the studied section (CC4, CC5, CC6 and CC7), represented by the family Peridiniaceae including *Cerodinium* sp., *Diconodinium* sp., *Isabelidinium* sp. cf. *I. pellucidum*, *Nelsoniella* sp., *Odontochitina spinosa*, *Odontochitina* spp., *Palaeocystodinium australinum*, *P. granulatum*, *P. lidiae*, *Spinidinium* sp., with lower proportions of the Family Goniaulacoideae such as *Exochosphaeridium* sp. and *Spiniferites ramosus* (Povilauskas & Guler, 2008), suggest an age of late Campanian–early Maastrichtian. This age would be consistent with that suggested by the spore-pollen associations.

Assemblage 2

In this group, some of the exclusive Association 2 taxa (lower levels of the Monte Chico Formation) are important from a chronostratigraphic standpoint. Among the most significant species are *Longapertites patagonicus*, in Argentina related to Danian deposits of Chubut Province (Archangelsky, 1973) and the oldest records in the Maastrichtian of the same province (Baldoni, 1992; Baldoni & Askin, 1993); *Proteacidites subscabrinus*, defined in New Zealand for the Oligocene, and recognized in Argentina and the Antarctic in the Maastrichtian–Danian; *Senipites tercassata*, defined for the Paleocene of Argentina (Archangelsky, 1973); *Beaupreaidites elegansiformis*, also distributed in the Campanian–Maastrichtian of Australia, the Antarctic and New Zealand (Dettmann & Jarzen, 1988; 1990; Cookson, 1950); *Ilexpollenites salamanquensis*, recognized from the Upper Cretaceous of New Zealand (Mc Intyre, 1968) and recorded from the Paleocene of Argentina (Archangelsky & Zamaloa, 1986); *Liliacidites vermiculatus*, defined in Argentina and distributed from the lower Paleocene

(Archangelsky & Zamaloa, 1986; Mautino & Anzótegui, 2002); *Tricolpites bibaculatus*, defined and distributed in the Paleocene of Argentina (Archangelsky & Zamaloa, 1986; Quattrocchio *et al.*, 1997); and *Quadraplanus brossus*, recognized primarily in Australia with a very restricted acme to the upper Maastrichtian–basal Danian? (Stover & Partridge, 1973; Helby *et al.*, 1987).

Nevertheless, other stratigraphic taxa listed in Association 2 are also important and continue to be recorded towards the top of the Monte Chico Formation (Association 3). Among them, the most significant are *Psilatricolporites* cf. *P. salamanquensis*, *Rhoipites minusculus*, *Rousea patagonica*, *Spinizonocolpites hialinus* and *Triporopollenites* cf. *T. ambiguus*, all features of Maastrichtian and Danian associations of Argentina (Archangelsky, 1973; Archangelsky & Zamaloa, 1986; Baldoni, 1992; Baldoni & Askin, 1993).

Association 2 presents the greatest similarities with associations from the Lefipán Formation, Upper Cretaceous of Chubut Province (Baldoni, 1992; Baldoni & Askin, 1993) and the La Irene Formation, upper Campanian–lower Maastrichtian of Santa Cruz Province (Povilauskas *et al.*, 2008), with whom it shares species such as *Liliacidites variegatus*, *Liliacidites kaitangataensis*, *Longapertites patagonicus*, *Spinizonocolpites hialinus*, *Tricolpites reticulatus*, *Peninsulapollis gillii*, *Rousea patagonica*, *Rhoipites minusculus*, *Triporopollenites ambiguus*, *Proteacidites tenuiexinus* and *Triatriopollenites lateflexus*, among others (Table 4).

Given the similarities to known biochrons, an age related to the Maastrichtian, probably late Maastrichtian, for the Association 2 (lower levels of the Monte Chico Formation) is suggested (Figure 5).

Assemblage 3

Among the taxa represented exclusively in this association that have stratigraphic significance is *Peninsulapollis askinia*; this species was defined for the Upper Cretaceous of Australia and Antarctica (Dettmann & Jarzen, 1988; Dettmann & Thomson, 1987; Truswell, 1983), but in Argentina is recognized from the Paleocene (Archangelsky & Seoane, 1994).

The greatest similarities of Association 3 occur with those from the Paso del Sapo and Lefipán formations (Papú, 1988a, 1988b, 1990; Baldoni, 1992; Baldoni & Askin, 1993), assigned to the upper Maastrichtian–basal Danian.

Based on this analysis, Association 3 is referred to the higher levels of the Monte Chico Formation around Maastrichtian–Danian in age. The increased diversity of *Nothofagidites* spp., together with the first records of *Nothofagidites dorotensis* and the extinction of *Quadraplanus brossus* might suggest a greater time restriction (basal Danian?) (Table 4). *Nothofagidites dorotensis* was defined in Argentina and recorded only from the Paleocene (Romero, 1973; Menéndez & Caccavari, 1975); meanwhile *Quadraplanus brossus* is an important guide fossil not documented beyond the base of the Danian.

Table 3. Stratigraphic distribution of the presents species and/or genera in the Cerro Cazador Formation and other Maastrichtian and Paleocene units in Southern South America and Antarctic Peninsula. Pedro Luro (1), Loncoche (2), Los Alamitos (3), Paso del Sapo (4), Lefipán (5), Salamanca (6), Cerro Dorotea (7), López de Bertodano (8), La Irene (9) formations. **Symbols:** ● shared species; ○ shared genera

Spores

- 1 *Bac.* cf. *B. conaumensis* (Cookson) Potonié, 1956
- 2 *Birretisp.* cf. *B. potoniaei* Delcourt & Sprumont, 1955
- 3 *Clavifera triplex* (Bolkhovitina) Bolkhovitina, 1966
- 4 *Cyatheaecidites annulatus* Cookson 1947, ex Potonié, 1956
- 5 *Cyatellidites minor* Couper, 1953
- 6 *Deltoidospora australis* (Couper) Pocock, 1970
- 7 *Echinospores* sp.
- 8 *Gleichenioidites apitanus* Llorens, 2008
- 9 *G. senonicus* Ross, 1949
- 10 *Ischyrosporites volkheimeri* Filatoff, 1975
- 11 *Ischyrosporites* sp.
- 12 *Laevigatosporites ovatus* Wilson & Webster, 1946
- 13 *Neoraistrickia* sp.
- 14 *Polyopidioidites speciosus* (Harris) Archangelsky, 1972
- 15 *Reticuloidosporites tenellus* Krutzsch, 1959
- 16 *Trilites parvullatus* Krutzsch, 1959
- 17 *Trilites* cf. *T. fasolae* Archangelsky, 1972
- 18 *Verrucosporites* sp. 1
- 19 *Verrucosporites antiquasporites* Dettmann, 1963
- 20 *Stereisporites* sp. 2

Gymnosperms pollen

- 21 *Anarcuriacites australis* Cookson, 1947
- 22 *Lygistopollenites florinii* Stover & Evans, 1973
- 23 *Microcachrytidites antarcticus* Cookson, 1947
- 24 *Phyllocladites mawsonii* Cookson, 1947 ex Couper, 1953
- 25 *Fodocarpidites elegans* Romero, 1977
- 26 *P. ellipticus* Cookson, 1947
- 27 *P. microreticuloidata* Cookson, 1947
- 28 *P.* sp.

Angiosperms pollen

- 29 *Arecipites minutiscabranus* (McIntyre, 1968) Milne, 1988
- 30 *Liliacidites kaianganensis* Couper, 1953
- 31 *L. cf. L. regularis* Archangelsky, 1973
- 32 *L. variegatus* Couper, 1953
- 33 *Nothofagidites saraensis* Menéndez & Caccavari de Filice, 1975
- 34 *Forcipites* sp.
- 35 *Peniculapollis gillii* (Cookson) Dettmann & Jarzen, 1988
- 36 *P.* sp.
- 37 *Psilariocolporites* sp.
- 38 *Rousea microreticulata* Archangelsky, 1986
- 39 *Triatriopollenites lateflexus* Archangelsky, 1973
- 40 *Tricolpites reticulatus* Cookson, 1947
- 41 *Triporopollenites* sp.

Assemblage 4

Association 4 (Cerro Dorotea Formation) is characterized by the exclusive presence of, among other species, *Bombacacidites* sp. 1, *Forcipites stipulatus*, *Nothofagidites waiapaeensis* and *Propylipollis microverrucatus* (Table 5).

From a chronostratigraphic point of view, among the most significant is *Forcipites stipulates*, defined in Australia and recognized from the Maastrichtian in Australia and Antarctica (Dettmann & Jarzen, 1988), without previous records in Argentina, and *Nothofagidites waipawaensis*, defined in Argentina, and recognized from the lower Paleocene (Romero, 1973; Romero & Zamaloa, 1997; Carrillo-Berumen *et al.*, 2013).

For its part, the greatest similarities of Association 4 (Cerro Dorotea Formation) occur with those from the Salamanca Formation (Paleocene) (Archangelsky, 1973). They share the presence of *Clavifera triplex*, *Cyatheacidites annulatus*, *Arecipites minutiscabrus*, *Liliacidites variegatus*, *Nothofagidites* spp., *Rhoipites minusculus*, *Rousea patagonica*, *Spinizonocolpites hialinus*, *Triatriopollenites lateflexus*, *Tricolpites reticulatus*, *Liliacidites regularis*, *Psilatricolporites salamanquensis* and *Ericipites scabrus* among the most significant taxa.

In comparison to the known biochrons spanning the stratigraphic position of the Cerro Dorotea Formation, some of the species found, and with special regard to the similarities found with other formations, suggest a similar age in the Danian. This temporal assignment is consistent with that previously indicated by Freile (1972) based on a preliminary palynological study of the Cerro Dorotea Formation.

PALEOECOLOGICAL IMPLICATIONS

Fossil evidence indicates that different groups experienced a global extinction event across the Cretaceous/Paleogene boundary. It eliminated 80% of marine invertebrates, the extinction of the dinosaurs occurred and there was a drastic reduction of many species of mammals (Pascual *et al.*, 1985; Pascual & Jaureguizar, 1990). The evidence of a comparable extinction to that of the fauna in the flora is ambiguous at least in the Southern Hemisphere (Diéguez, 2003). In the Northern Hemisphere in various sections of North America, and also in Japan and Europe there was a violent and rapid decline in the abundance and diversity of various plant groups across the area (Orth *et al.*, 1981). This process was followed by a significant increase in the concentration of spores of ferns, an event known as the “fern spike”. The latter was interpreted as a response of vegetation to strong ecological trauma. The vegetation diversity is later restored, but with a different composition to that presented before the K/T boundary. In the Southern Hemisphere the marine biota was as affected as in the Northern Hemisphere, but until recently there was no evidence of substantial changes in plant communities across the area. Palynological studies conducted in Australia and Antarctica showed very little change through the Cretaceous–Paleogene (Askin, 1988; Macphail, 1994). However, recent studies in New Zealand and Argentina showed a disturbance

Table 4. Relative abundance of the species in the selected levels of Monte Chico Formation.

Table 4. Cont.

Monte Chico Formation	MC3	MC5	MC7	MC10	MC11	MC12	MC13	MC14	MC15	MC17	MC22
61 <i>Cycadopites</i> sp. 1	0	0	0	0	0	0	0	0	1	0,67	0
62 <i>Dacrycarpites australiensis</i>	0	0	6	2,39	2	0,64	0	0	0	0,85	0
63 <i>Gameroites psilasaccus</i>	0	0	0	2	0,64	0	0	0	0	0	0
64 <i>Gameroites</i> sp. 1	0	0	0	1	0,32	0	0	0	0	0	0
65 <i>Lygisteppollenites florinii</i>	0	0	4	1,60	7	2,26	0	0	4	1,97	0
66 <i>Lygisteppollenites</i> sp. 1	0	0	1	0,39	1	0,32	0	0	1	0,49	0
67 <i>Microcachrylrites antarcticus</i>	7	3,04	0	10	3,23	17	8,94	8	3,44	9	4,43
68 <i>Phyllocladites mawsonii</i>	9	3,91	9	3,58	2	0,64	0	0	0	17	6,15
69 <i>Podocarpites elegans</i>	4	1,73	3	1,19	3	0,97	12	6,31	0	0	5,2,63
70 <i>Podocarpites ellipticus</i>	8	3,47	Tr	0	9	2,91	0	10	4,31	10	4,92
71 <i>Podocarpites marwickii</i>	3	1,30	1	0,39	5	1,61	0	0	0	0	3,15
72 <i>Podocarpites</i> cf. <i>P. microreticuloidata</i>	2	0,87	3	1,19	3	0,97	0	0	0	0	0
73 <i>Podocarpites microreticuloidata</i>	0	0	0	4	1,29	0	0	0	4	1,97	1
74 <i>Trisaccites microsaccatum</i>	0	0	0	5	1,61	0	0	0	0	1	0,36
Subtotals	33	14,3	31	12,3	61	19,7	31	16,3	19	8,19	28
Pollen Angiosperms											
75 <i>Arecipites minitiscabritus</i>	12	5,21	0	0	8	2,58	0	0	3	1,29	9
76 <i>Beaupreidites elegansiformis</i>	0	0	0	0	0	0	0	0	1	0,36	Tr
77 <i>Beaupreidites</i> sp. 1	0	0	0	0	0	0	0	0	1	0,36	Tr
78 <i>Clavamonocolpites</i> sp. 1	0	0	4	1,60	3	0,97	0	0	0	0	0
79 <i>Clavatricolpites</i> sp. 1	0	0	1	0,39	1	0,32	0	0	0	0	0
80 <i>Ericipites scabrius</i>	0	0	0	2	0,64	0	0	0	4	1,97	4
81 <i>Forcipites</i> sp. "A" en Dettmann y Jarzen 1988	1	0,43	0	0	0	0	0	0	1	0,36	10
82 <i>Forcipites sabulosus</i>	4	1,73	0	0	1	0,32	1	0,52	0	0	2,17
83 <i>Haloragacitides trioratus</i>	0	0	Tr	0	1	0,32	0	0	0	0	3,15
84 <i>Ilexpollenites salmanticensis</i>	0	0	0	0	0	0	2	0,86	0	0	0
85 <i>Liliacitides</i> sp. cf. <i>L. crassilabratius</i>	0	0	0	0	0	0	0	2	0,98	0	0
86 <i>Liliacitides kaitangataensis</i>	4	1,73	11	4,38	7	2,26	0	0	0	0	3,157
87 <i>Liliacitides</i> sp. cf. <i>L. regularis</i>	0	0	0	4	1,29	0	0	4	1,72	0	0
88 <i>Liliacitides variegatus</i>	6	2,61	6	2,39	4	1,29	5	2,63	2	0,86	0
89 <i>Liliacitides vermireticulatus</i>	2	0,87	5	1,99	3	0,97	3	1,57	0	0	4
90 <i>Liliacitides</i> sp. 1	0	0	0	4	1,29	0	0	0	0	0	0
91 <i>Longapertites patagonicus</i>	0	0	3	1,19	0	0	0	0	0	0	0
92 <i>Nothofagidites kaitangataensis</i>	0	0	0	0	0	0	0	0	4	1,08	4
93 <i>Nothofagidites dorotensis</i>	0	0	0	0	0	0	0	1	0,97	2	1,05
94 <i>Nothofagidites nama</i>	0	0	0	0	0	0	0	1	0,43	0	0
95 <i>Nothofagigites saraensis</i>	0	0	0	0	0	0	0	3	1,47	3	1,08
96 <i>Peninsulapollis astikmae</i>	0	0	0	0	0	2	1,05	0	0	3,26	2
97 <i>Peninsulapollis gilli</i>	4	1,73	15	5,97	13	4,20	0	0	7	3,01	0
98 <i>Peninsulapollis triswelliae</i>	2	0,87	6	2,39	3	0,97	6	3,15	4	1,72	0
99 <i>Peninsulapollis</i> sp. cf. <i>P. triswelliae</i>	0	0	2	0,79	2	0,64	0	0	0	1	0,52
100 <i>Peninsulapollis</i> sp. 1	0	0	7	2,78	0	0	0	0	0	9	3,26
101 <i>Peripropollenites demarcatus</i>	0	0	3	1,19	0	0	0	0	0	0	4

Table 4. Cont.

Monte Chico Formation	MC3	MC5	MC7	MC10	MC11	MC12	MC13	MC14	MC15	MC17	MC22
102 <i>Proteacidites beddoesii</i>	3 1,30	3 1,19	Tr 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
103 <i>Proteacidites parvus</i>	0 0	Tr 0	1 0,32	0 0	0 0	0 0	1 0,36	0 0	0 0	0 0	0 0
104 <i>Proteacidites subscabratus</i>	2 0,87	2 0,79	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
105 <i>Proteacidites tenuiximus</i>	Tr 0	1 0,39	Tr 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
106 <i>Pseudowinterapollis couperi</i>	0 0	1 0,39	Tr 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
107 <i>Psilatricolpites patagonicus</i>	5 2,17	5 1,99	4 1,29	0 0	1 0,43	0 0	0 0	0 0	0 0	0 0	0 0
108 <i>Psilatricolpites sp. 1</i>	0 0	3 1,19	0 0	0 0	0 0	0 0	0 0	0 0	1 0,67	Tr 0	1 0,72
109 <i>Psilatricolporites cf. P. salamanquensis</i>	3 1,30	2 0,79	0 0	0 0	0 0	0 0	0 0	1 0,52	1 0,67	0 0	2 1,43
110 <i>Psilatricolporites sp. 1</i>	0 0	0 0	0 0	0 0	0 0	0 0	4 1,44	2 1,05	0 0	2 1,71	0 0
111 <i>Quadraplanus brossus</i>	1 0,43	2 0,79	0 0	0 0	2 0,86	1 0,49	0 0	0 0	0 0	0 0	0 0
112 <i>Rhoipites baculatus</i>	0 0	2 0,79	3 0,97	0 0	2 0,86	0 0	0 0	1 0,52	1 0,67	0 0	0 0
113 <i>Rhoipites minusculus</i>	1 0,43	3 1,19	2 0,64	0 0	1 0,43	0 0	2 0,72	2 1,05	2 1,35	Tr 0	0 0
114 <i>Rhoipites sp. 1</i>	0 0	Tr 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
115 <i>Rousea microreticulata</i>	2 0,87	1 0,39	0 0	0 0	Tr 0	0 0	0 0	0 0	0 0	0 0	0 0
116 <i>Rousea patagonica</i>	0 0	0 0	0 0	0 0	2 0,86	0 0	0 0	0 0	0 0	0 0	0 0
117 <i>Senipites terrassata</i>	2 0,87	2 0,79	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
118 <i>Sparganiaceapollenites barungensis</i>	3 1,30	4 1,60	5 1,61	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
119 <i>Spinizonocolpites hialimus</i>	2 0,87	3 1,19	3 0,97	0 0	0 0	5 2,46	0 0	0 0	3 2,02	0 0	0 0
120 <i>Spinizonocolpites sp. 1</i>	0 0	0 0	0 0	0 0	0 0	0 0	0 0	1 0,52	1 0,67	0 0	0 0
121 <i>Triatriopollenites bertelsii</i>	2 0,87	2 0,79	0 0	3 1,57	3 1,29	0 0	7 2,53	2 1,05	0 0	Tr 0	0 0
122 <i>Triatriopollenites latiflexus</i>	4 1,73	5 1,99	4 1,29	0 0	0 0	0 0	0 0	0 0	0 0	0 0	2 1,43
123 <i>Tricolpites bibaculatus</i>	0 0	2 0,79	2 0,64	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
124 <i>Tricolpites reticulatus</i>	8 3,47	4 1,60	6 1,94	0 0	8 3,44	8 3,94	8 2,89	4 2,10	4 2,70	1 0,85	4 2,87
125 <i>Tricolpites sp. cf. T. reticulatus</i>	1 0,43	1 0,39	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0	0 0
126 <i>Tricolpites sp. 1</i>	2 0,87	1 0,39	1 0,32	0 0	0 0	0 0	2 0,72	0 0	0 0	0 0	0 0
127 <i>Tricolpites sp. 2</i>	1 0,43	Tr 0	0 0	0 0	Tr 0	0 0	0 0	0 0	3 2,02	0 0	0 0
128 <i>Triplopollenites sp. cf. T. ambiguus</i>	1 0,43	Tr 0	0 0	0 0	0 0	0 0	1 0,36	0 0	0 0	0 0	0 0
Subtotals	78 33,9	112 44,6	92 29,7	20 10,5	42 18,1	39 19,2	92 33,3	62 32,6	52 35,1	17 14,5	25 18
Totals of continental items	201 87,4	230 91,6	275 88,9	127 66,8	135 58,2	136 66,9	237 85,8	156 82,1	119 80,4	59 50,4	67 48,2
Green algae											
129 <i>Borycococcus</i> sp.	tr 0	tr 0	0 0	0 0	1 0,43	1 0,49	2 0,72	0 0	1 0,67	0 0	1 0,72
130 <i>Carinipollis gelsetensis</i>	2 0,87	3 1,19	1 0,32	3 1,57	3 1,29	2 0,98	tr 0	1 0,52	tr 0	1 0,85	2 1,43
Subtotals of fresh water	7 3,04	4 1,60	4 1,29	8 4,21	4 1,72	4 1,97	5 1,81	3 1,57	3 2,02	2 1,71	4 2,87
131 Acritars	5 2,16	1 0,39	4 1,29	5 2,63	1 0,43	1 0,49	3 1,08	3 1,57	2 1,35	1 0,85	1 0,72
132 Dinocysts	29 12,5	30 11,9	33 10,6	63 33,1	96 41,3	67 33	39 14,1	33 17,3	29 19,6	58 49,5	72 51,8
Totals of marine items	34 14,6	31 12,3	37 11,9	68 35,7	97 41,7	68 33,5	42 15,1	36 18,8	31 20,9	59 50,3	73 52,5
Totals of palynomorphs (continental and marine)	230 100	260 100	309 100	190 232	100 203	100 276	100 190	100 148	100 117	100 139	100

Table 5. Table relative abundance of the species in the selected levels of Cerro Dorotea Formation.

Cerro Dorotea Formation	CD1	CD3	CD7	CD17	CD21	CD23	CD24
	Cont.	%	Cont.	%	Cont.	%	Cont.
Spores							
1 <i>Baculatisporites conaumensis</i>	3	2,54	0	0	5	7,7	3
2 <i>Baculatisporites turbioensis</i>	0	0	0	0	0	0	0
3 <i>Baculatisporites sp. 1</i>	0	0	0	0	0	0	0
4 <i>Biretisporites crassilabrus</i>	1	0,84	0	0	1	1,54	1
5 <i>Biretisporites</i> sp. 1	0	0	0	0	0	0	0,72
6 <i>Biretisporites</i> sp. III	0	0	0	0	0	0	Tr
7 <i>Clavifera triplex</i>	0	0	0	0	1	1,54	0
8 <i>Concavissimisporites</i> sp. 1	0	0	0	0	0	0	0
9 <i>Convolutisporites</i> sp. 1	0	0	0	0	0	0	0
10 <i>Cyathacidites annulus</i>	6	5,10	0	0	0	10	7,19
11 <i>Cyathidites asper</i>	2	1,70	0	0	0	0	0
12 <i>Cyathidites australis</i>	1	0,84	6	15,8	3	4,61	2
13 <i>Cyathidites concavus</i>	0	0	0	0	0	2	1,43
14 <i>Cyathidites minor</i>	5	4,23	2	5,26	1	1,54	4
15 <i>Cyathidites punctatus</i>	0	0	0	0	0	Tr	0
16 <i>Deltoidospora australis</i>	4	3,39	0	0	0	3	2,15
17 <i>Echinosporis</i> sp. 1	0	0	0	0	0	0	4
18 <i>Foveosporites canalis</i>	0	0	0	0	0	0	2,68
19 <i>Gabonisporis</i> sp. 1	0	0	0	0	0	0	2,87
20 <i>Gleicheniidites apitanus</i>	0	0	0	0	0	0	4
21 <i>Gleicheniidites senonicus</i>	1	0,84	0	0	4	6,15	1
22 <i>Gleicheniidites</i> sp. 1	1	0,84	0	0	0	0,72	1
23 <i>Interlobites intraverrucatus</i>	0	0	0	0	0	0	0
24 <i>Ischyrosporites gremius</i>	0	0	0	0	0	0	0
25 <i>Ischyrosporites punctatus</i>	0	0	0	0	0	0	0
26 <i>Ischyrosporites volkheimeri</i>	0	0	0	0	1	0,72	0
27 <i>Klhkisporites</i> sp. 1	0	0	0	0	1	0,72	0
28 <i>Laevigatosporites ovatus</i>	3	2,54	0	0	3	4,61	4
29 <i>Leiotriletes regularis</i>	0	0	0	0	1	1,54	0
30 <i>Leptolepidites verrucatus</i>	0	0	0	0	0	2	1,43
31 <i>Osmundacitides wellmanii</i>	0	0	0	0	0	Tr	0
32 <i>Peromylonites vellosum</i>	1	0,84	0	0	0	0	1
33 <i>Punctatosporites scabratus</i>	0	0	0	0	0	0	0,67
34 <i>Retinilites austroclavatidites</i>	1	0,84	0	0	0	0	0
35 <i>Retinilites</i> sp. 1	0	0	0	0	0	0	0,67
36 <i>Rouseispores reticulatus</i>	0	0	0	0	0	0	Tr
37 <i>Rugulatisporites nequenensis</i>	0	0	0	0	0	0	0
38 <i>Rugulatisporites</i> sp. 1	0	0	0	0	0	0	0
39 <i>Rugulatisporites micraulaxus</i>	0	0	0	0	0	0	0
40 <i>Stereisporites antiquasporites</i>	3	2,54	0	0	1	0,72	3
41 <i>Trilites parvallatus</i>	3	2,54	0	0	0	0	2

Table 5. Cont.

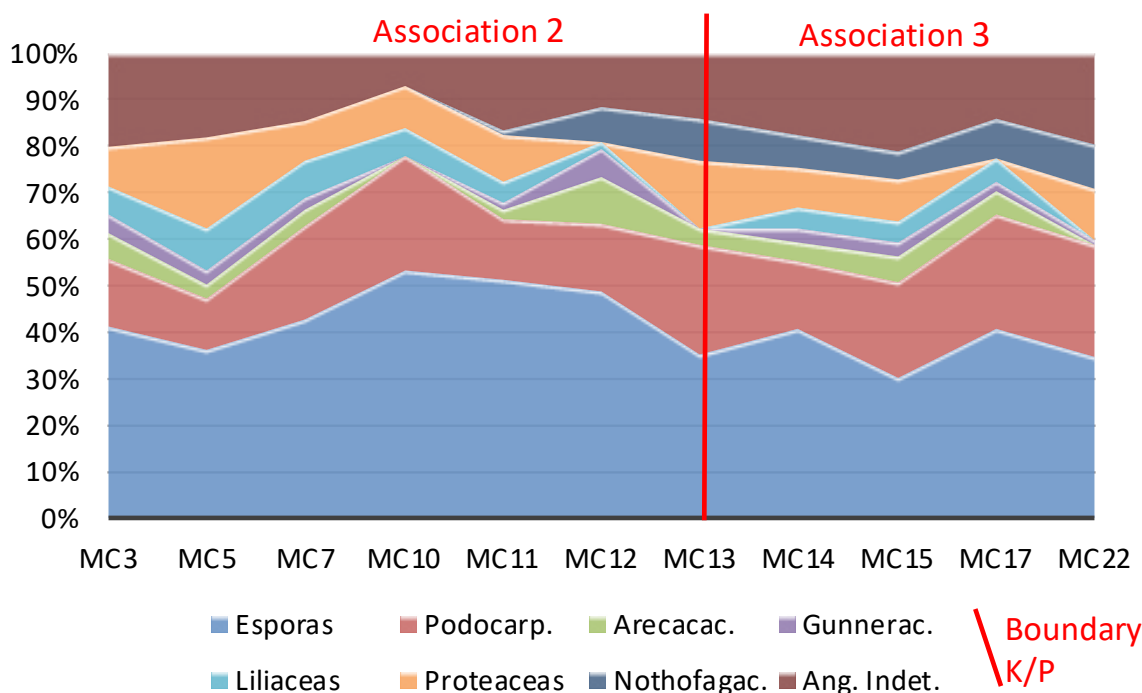


Figure 5. Graph showing the Cretaceous/Paleogene boundary suggested for Monte Chico Formation.

in the vegetation through the boundary, with a temporary loss of angiosperms and a sharp reduction in several groups of gymnosperms and spores (Pocock, 1962; Vajda *et al.*, 2001; Vajda & Raine, 2003; Barreda *et al.*, 2004; Cúneo *et al.*, 2008). According to the observations made in this paper it could be determined with a degree of certainty that the position of the Cretaceous–Paleogene boundary is located between Associations 2 and 3, as suggested in previous studies (Malumíán & Panza, 1996), in the Monte Chico Formation. Its location between these two associations is suggested mainly by the temporal ranges of some species of restricted distributions, especially *Quadrupланus brossus* and *Nothofагidites dorotensis*. *Quadrupланus brossus* is a characteristic species of the *Tricolpites longus* Zone defined for SE Australia (Helby *et al.*, 1987) assigned to the upper Maastrichtian–basal Danian? and the Monte Chico Formation is virtually restricted to Association 2. On the other hand, *Nothofагidites dorotensis*, which has its first appearance in Association 3, has no record prior to the Danian.

At the moment, however, no significant changes in diversity and/or abundance of species between these two associations appear, as was documented for other basins in Argentina (Vajda & Raine, 2003; Barreda *et al.*, 2004; Cúneo *et al.*, 2008). This result may be due on the one hand to (i) there having been no disturbance in the vegetation across the boundary in the southern sector of Patagonia, and/or (ii) that the level of detail of sampling is insufficient to document it.

DISCUSSION AND CONCLUSIONS

An analysis of the distribution of pollen and spore species in the units recognized four palynological associations with unique characteristics: Association 1, was recognized in the upper levels of the Cerro Cazador Formation and lacks the characteristic taxa of younger associations; Association 2, was recognized in the lower and middle levels of the Monte Chico Formation and lacks the characteristic elements of the lower (1) and upper (3, 4) associations; Association 3, was recognized in the upper section of Monte Chico Formation; and Association 4, was recognized in the Cerro Dorotea Formation (Figure 4).

Based on the known stratigraphic distribution of the species and observed affinities, it follows that Association 1 (upper levels of the Cerro Cazador Formation) has an inferred age of late Campanian–early Maastrichtian; Association 2 (lower and middle levels of the Monte Chico Formation) has an age in the region of the Maastrichtian, probably late Maastrichtian; Association 3 (higher levels of the Monte Chico Formation) has an inferred age limited to the proximity of the Maastrichtian–Danian; and Association 4 (Cerro Dorotea Formation) has a Danian age taking into account previous records (Archangelsky, 1973).

According to this analysis the position of the K/P boundary would be located between Associations 2 and 3, within the Monte Chico Formation (Figure 5). However, no significant changes in the diversity and/or abundance of species between

these two associations were seen, as was documented for other basins in Argentina. This result may be due on the one hand because (i) there has been no disturbance in the vegetation across the boundary in the southern sector of Patagonia and/or (ii) the level of sampling detail is insufficient to document it. Further studies on these and other sections, with a greater level of detail, may provide new information to answer this question.

From the point of view of the depositional environment of the three units (Cerro Cazador, Monte Chico and Cerro Dorotea formations), they would have evolved in a marine environment with progressively more marginal conditions that would indicate a progressive shallowing of the basin.

From a paleoclimatic perspective, associations recovered from the Cerro Cazador Formation suggest the development of vegetation with a high participation of herbaceous elements, especially ferns and plants of a terrestrial habit such as Liliaceae and Gunneraceae. Among the dominant elements of palm trees and the Proteaceae, the Podocarpaceae may have evolved away from the depositional environment, judging by their low relative frequencies in relation to the high pollen productivity of the group. The prevailing paleoclimatic conditions would have been warm and wet.

The associations from the Monte Chico Formation suggest the development of plant communities dominated by Proteaceae and Arecaceae with a dense cover of ferns under a hot and wet climate. The abundance of ferns indicates the presence of flooded or wet soil.

Spore-pollen associations recovered from the Cerro Dorotea Formation show no significant differences to those from the underlying Monte Chico Formation. Both units are dominated by fern spores, followed by Arecaceae, Liliaceae and Proteaceae pollen. Besides, the frequency of Nothofagaceae and Podocarpaceae elements in the Cerro Dorotea Formation suggest a forest close to marginal marine paleoenvironment where they were deposited. The vegetation developed under temperate warm and humid conditions.

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Appendix 1. List of species found.

Assemblage 1 (Cerro Cazador Formation)

- Baculatisporites* cf. *B. comaumensis* (Cookson) Potonié, 1956
Biretisporites cf. *B. potoniaei* Delcourt & Sprumont, 1955
Ischyosporites sp. 1
Podocarpidites sp. 2
Trilites cf. *T. fasolae* Archangelsky, 1972
Triporopollenites sp. 1
Verrucosporites sp. 2

Assemblage 2 (basal levels and middle of Monte Chico Formation)

- Baculatisporites turbioensis* Archangelsky, 1972
Beupreadites elegansiformis Cookson, 1950
Biretisporites crassilabratus Archangelsky, 1972
Camarozonosporites ohaiensis (Couper, 1953) Dettmann & Playford, 1968
Ceratosporites equalis Cookson & Dettmann, 1958
Classopollis sp. 1
Forcipites sabulosus (Dettmann & Playford) Dettmann & Jarzen, 1988
Haloragacidites trioratus Couper, 1953
Ilexpollenites salamanquensis Archangelsky, 1986
Liliacidites vermiculatus Archangelsky y Zamaloa, 1986
Longapertites patagonicus Archangelsky, 1973
Nothofagidites kaitangataensis (Te Punga) Romero, 1973
Ornamentifera echinata (Bolkhovitina) Bolkhovitina, 1966
Peninsulapollis truswelliae Dettmann & Jarzen, 1988
Periporopollenites demarcatus Stover, 1973
Peromonolites vellosus Partridge, 1973
Proteacidites beddoesii Stover, 1973
Proteacidites subscabratus Couper, 1960
Proteacidites tenuixinus Stover in Stover & Partridge, 1973
Psilatricolpites patagonicus Freile, 1972
Psilatricolporites cf. *P. salamanquensis* Archangelsky & Zamaloa, 1986
Quadruplicatus brossus (Stover) Stover & Partridge, 1973
Rhoipites baculatus Archangelsky, 1973
Rhoipites minusculus Archangelsky, 1983
Rousea microreticulata Archangelsky, 1986
Rousea patagonica Archangelsky, 1973
Senipites tercristata Archangelsky, 1973
Sparganiaceapollenites barungensis Harris, 1972
Spinizonocolpites hialinus Zamaloa & Archangelsky, 1986
Triatriopollenites bertelsii Archangelsky, 1973
Trilites tuberculiformis Cookson, 1947
Tricolpites bibaculatus Archangelsky & Zamaloa, 1986
Triporopollenites cf. *T. ambiguus* (Stover) Stover & Partridge, 1973
Tuberculatosporites parvus Archangelsky, 1972

Assemblage 3 (upper levels of Monte Chico Formation)

- Beupreadites* sp. 1
Ericipites scabratus Harris, 1965
Gamerroites psilasaccus (Archangelsky & Romero, 1974) Archangelsky, 1988
Liliacidites sp. 1
Nothofagidites dorotensis Romero, 1973
Nothofagidites nana Romero, 1977
Peninsulapollis askiniai Dettmann & Jarzen, 1988
Pseudowinterapollis couperi Krutzsch, 1970 emend. Mildenhall, 1979

Assemblage 4 (Cerro Dorotea Formation)*Bombacacidites* sp. 1*Forcipites stipulatus* (Stover & Evans) Dettmann & Jarzen 1988*Nothofagidites waipawaensis* (Couper 1960) Fasola 1969,*Propylipollis microverrucatus* Truswell & Owen, 1988*Tetracolporites* sp. 1