A new Late Triassic dipteridacean fern from the Paso Flores Formation, Neuquén Basin, Argentina

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Sterile and fertile fronds of dipteridacean ferns from the Paso Flores Formation (late Norian–Rhaetian) at Cañadón de Pancho area, south of the Neuquén Province, Patagonia (Neuquén Basin), Argentina, are described. The Paso Flores Formation specimens comprise an interesting mixture of features showing a unique combination that does not fit in the diagnosis of any of the known fossil genera of Dipteridaceae. Characters such as the number and shape of primary segments in each rachial arm, the coalescence of the primary segments at the base of the frond, the distribution of sori and the number of sporangia per sori allow to differentiate the Paso Flores Formation specimens from the other genera of the family. The new genus and species, *Patagoniapteris artabeae* is proposed. The specimens share some frond morphological features with the fossil genera *Clathropteris*, *Digitopteris*, *Thaumatopteris*, *Sewardalea* and with some species of the *Dictyophyllum*, as well as the characteristic of the sori with the living species *Dipteris lobbiana*. The Paso Flores Formation environments developed on the western margin of Gondwana under seasonal temperate-warm and humid to sub-humid climates with a marine influence from the west. The Cañadón de Pancho assemblages are late Norian–Rhaetian in age, being the youngest fossil flora recorded from Argentinian Triassic basins to date.

**Key words:** Gleicheniales, Dipteridaceae, Norian, Rhaetian, Paso Flores Formation, Neuquén Basin, Patagonia, Gondwana.

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Received 5 December 2020, accepted 21 May 2021, available online 29 November 2021.

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**Introduction**

The Dipteridaceae (order Gleicheniales) comprises two extant genera, *Dipteris* Reinwardt, 1828, and *Cheiropleuria* Presl, 1851 (Smith et al. 2006; Taylor et al. 2009; Zhang et al. 2013). Currently, they are distributed in warm tropical to subtropical regions of Asia, Australia, and Polynesia (Corсин and Waterlot 1979; Kramer et al. 1990; Zhang et al. 2013), where they are generally found at higher altitudes in open exposed areas (Kramer et al. 1990; Choo et al. 2016). *Dipteris* includes a species *Dipteris lobbiana* (Hooker, 1853) Moore, 1857, which grows on stream banks in open canopy, and other species, such as *Dipteris conjugata* Reinwardt, 1828, a colonizer of disturbed sites and exposed ridges (Cantrill 1995).

The fossil records of this group of ferns are distributed worldwide and they were diverse in the Southern Hemisphere during the Late Triassic to Early Jurassic interval when they reached their greatest generic diversity (Corсин and Waterlot 1979; Tidwell and Ash 1994; Bomfleur and Kerp 2010; Zhou et al. 2015; Bodnar et al. 2018). The earliest fossils with the distinctive venation pattern of Dipteridaceae were recorded from Middle Triassic strata (Webb 1982; Tidwell and Ash 1994; Cantrill 1995; Kustatscher and van Konijnenburg-van Cittert 2011; Bodnar et al. 2018) suggesting that the origin of the family took place in the Early Triassic or even earlier, during the late Palaeozoic (Tidwell and Ash 1994; Choo et al. 2016; Choo and Escapa 2018). This fern family decreased its diversity during Late Jurassic and Cretaceous (Choo and Escapa 2018). Cretaceous and Palaeogene records of Dipteridaceae are scarce and consist only of various species of *Hausmannia* (Choo et al. 2016). In Argentina, the Dipteridaceae have been recorded in the late Anisian–early
Ladinian and late Norian–Rhaetian intervals (Bodnar et al. 2018).

Smith et al. (2006) and Zhang et al. (2013) described the morphology of the extant Dipteridaceae characterized by a long stipe that bears fan-shaped fronds with toothed margins, bilobated and often deeply dissected lamina with a typical reticulate venation with veinlets inside the meshes. Exindusiate, discrete, compital sori are scattered over the abaxial surface of fertile fronds; sporangia maturation simultaneous or maturation mixed, with vertical or slightly oblique annulus.

Fossil fronds assigned to the Dipteridaceae are represented by several species included in the genera *Clathropteris* Brongniart, 1828, *Dictyophyllum* Lindley and Hutton, 1834, *Digitopteris* Pott and Bomfleur in Pott et al., 2018, *Goepertella* Ôishi and Yamasita, 1936 emend. Arrondo and Petriella, 1982, *Hausmannia* Dunker, 1846, *Sewardalea* Choo and Escapa, 2018, and *Thaumatopteris* Goepertt, 1841 (Herbst 1992a, b; Choo and Escapa 2018).

Dipteridacean genera were established based on the gross morphology of fronds and their venation patterns (Ôishi and Yamasita 1936; Arrondo and Petriella 1982; Herbst 1992a, b; Rees 1993; Rees and Cleal 2004). In Argentina, Stipanicic and Menéndez (1949) described specimens of Dipteridaceae according to Ôishi and Yamasita’s concept (1936), although they remarked that there are controversial criteria in the dipteridacean classification among different authors. The taxonomic classification proposed by Herbst (1992a, b), in agreement with that of Ôishi and Yamasita (1936), considers morphological features, such as the dissection of the frond lamina, and the disposition and torsion of the primary veins originated from the division of the stipe. Based on these criteria, Herbst (1992a, b) established three subgenera within the genus *Dictyophyllum*: *Dictyophyllum*, *Thaumatopteris*, and *Clathropteris*.

Choo and Escapa (2018) contributed to the first verifiable phylegetic hypothesis of the evolution of the Dipteridaceae, analyzing extinct as well as extant taxa in a single cladistic study and concluded that the evolutionary trend in this family has been toward increasing complexity in the venation pattern and laminal fusion. They recognized only five fossil genera: *Goeperttella*, *Thaumatopteris*, *Clathropteris*, *Digitopteris*, and *Sewardalea*. Choo and Escapa (2018) considered that fossil genera such as *Dictyophyllum*, *Kenderlykia* Tururanova-Ketova, 1962, *Hausmannia*, and *Prototrichipis*Andreac, 1853, are ambiguously placed on the tree and are recognized as possibly unnatural morphogenea.

Three dipteridacean species from the Paso Flores Formation were previously described: *Dictyophyllum tenuifolium* (Stipanicic and Menéndez, 1949) Bonetti and Herbst, 1964, *Thaumatopteris rothi* Frengüelli, 1941, and *Goeperttella stipanicicici* Herbst, 1993 (Frengüelli 1941; Herbst 1964, 1993; Stipanicic and Menéndez 1949; Bonetti and Herbst 1964; Herbst 1992a, b) (Table 1).

The Triassic species with fertile fronds described in Argentina are: *Dictyophyllum tenuifolium* which has sori irregularly arranged along the primary veins and at the base of the secondary veins, as well as on the lamina occasionally, and *Thaumatopteris tenuiserrata* Menéndez, 1951, which has sori irregularly arranged among the secondary veins (Menéndez 1951; Herbst 1992a, b). In specimens reported from the Jurassic strata of Argentina sori are arranged on the lamina in quite a distinct manner. *Thaumatopteris rocablanquesis* Herbst, 1965, has the sori grouped over the whole abaxial surface; *Clathropteris obovata* Ôishi, 1932, has tetra-hexasporangiate sori located in each areole of third order, and *Clathropteris meniscioides* (Brongniart, 1825) Brongniart, 1828, has the sori scattered across the abaxial surface (Herbst 1965, 1966, 1992a, b; Choo et al. 2016) (Table 2).

In this contribution we describe a new taxon of Dipteridaceae fertile and sterile fronds, *Patagoniapteris artabeae* gen. and sp. nov. from the uppermost part of the Paso Flores Formation of late Late Triassic age, in the Cañadón de Pancho area, Neuquén Province, Patagonia.

**Institutional abbreviations.**—MCF-PBPB, Palaeontological Collection of the Museo Municipal “Carmen Funes”, Plaza Huincul city, Neuquén Province, Argentina.

**Nomenclatural acts.**—This published work and the nomenclatural acts it contains have been registered in Plant Fossil Names Registry (PFNR): urn:lsid:plantfossilnames.org:ref:972.

**Geological setting.**

The Neuquén Basin is located on the eastern side of the Andes and central Chile, between 30–41° S latitude (Fig. 1). It originated during the Late Triassic–Early Jurassic on the western edge of Gondwana by continental extension (D’Elía et al. 2012). Syn-rift depocenters developed during the initial stages of the basin formation as isolated troughs linked with profuse magmatic activity (Legarreta and Uliana 1996; Spalletti et al. 1999; Howell et al. 2005). They were filled by a complex variety of clastic and volcaniclastic deposits associated with extensive lava flows known as the Precuyano Cycle (Gulisano et al. 1984; Legarreta and Gulisano 1989). The southern part of the Neuquén Basin is characterized by reactivated normal faults and reverse faults related to the evolution of the Andes Cordillera resulting in well exposed Mesozoic successions of the initial stage of the basin (D’Elía et al. 2012).

One of the initial syn-rift Upper Triassic siliciclastic continental units of the basin, is the Paso Flores Formation that crops out as isolated sections at the south of the Neuquén Basin, northern Patagonia, Argentina (Figs. 1, 2). In this region the basement is constituted by metamorphic and igneous rocks of the Late Palaeozoic age and by the Choïyoi magmatic-volcanic complex (upper Permian–Middle Triassic). Locally, the formation is overlain unconformably by an Lower Jurassic coastal marine siliciclastic sequence of the Nestares Formation (Zavattieri and Volkheimer 2001;
The Paso Flores strata were studied by Nullo (1979), González Díaz (1982), Lapido et al. (1984), Spalletti et al. (1988), Ganuza et al. (1995), and references therein. The type section of the unit is exposed at Cerro Mariana and surrounding areas at the Estancia Manantiales de Paso Flores, on the south-eastern margin of the Limay river along the boundary between the Río Negro and Neuquén provinces (Fig. 2). The outcrops of the Paso Flores Formation on either side of the Limay River (more than 3 km east of the Alerca Dam), and those of the Lomas and Cañadón de Ranquel Huao area (Fig. 2) are the thickest sections, representing alluvial fan deposits and gravelly braided systems described in detail by Spalletti et al. (1988, 1990). These sections represent the lower to middle parts of the unit. The thinnest section of the Paso Flores Formation crops out at the Cañadón de Pancho area, on the western side of the Collón Curá and south of the Quemquemtreu rivers, in the south-west of the Neuquén Province (González Díaz 1982; Ganuza et al. 1995) (Figs. 2–3). That section has been considered to constitute the formation’s upper part (Zavattieri and Mego 2008).

Cañadón de Pancho section.—In this area, the Paso Flores Formation crops out in an extensive area between the Quemquemtreu and Caleufú rivers (Figs. 2, 3). González Díaz (1982), for the first time, referred to the outcrops of the western margin of the Collón Curá River as belonging to the Paso Flores Formation. Here, it lies unconformably on granitic rocks (Huechulafquen Formation) and on the metamorphic complex (Cushamen Formation and equivalents) of the late Palaeozoic age (Varela et al. 2005), and, in turn, is covered discordantly by light-grey tuffitic continental sedimentary strata of the Collón Curá Formation (middle Miocene age) (Fig. 3). The Paso Flores Formation at the Cañadón de Pancho section is composed of four successive sedimentary facies associations described in detail by Ganuza et al. (1995): (i) lenticular beds of medium to fine clast-supported conglomerate, interbedded with lenticular coarse sandstone and conglomeratic sandstone deposited in a braided fluvial system; (ii) laterally persistent laminated and rippled mudstone and siltstone with scattered thin coal layers (marginal lacustrine facies) that upwards change to coarse and thick sandstone deposits (progradational mouth bars of deltaic system); (iii) interbedded tabular mudstone and lenticular sandstone beds, interpreted as deposits of a low-sinuosity meandering fluvial system; and (iv) lenticular coarse-grained sandstone and fine conglomerate formed in a braided fluvial system revealing renewed higher energy. The well-preserved megafossil remains and microfossil assemblages studied were recovered from low-sinuosity meandering fluvial systems, deltaic and marginal lacustrine siltstone, mudstone, and fine-grained sandstone deposits.

Palaeobotanical records of the Paso Flores Formation.—The fossil plant content of the Paso Flores Formation has been studied by several authors (Frengüelli 1937; Bonetti and Herbst 1964; Spalletti et al. 1988; Arrondo et al. 1991; Morel et al. 1992, 1999; Ganuza et al. 1992; Herbst 1993; Artabe et al. 1994; Zamuner and Artabe 1994; Ganuza et al. 1995; Gnaedinger and Zavattieri 2017a, b), who recorded abundant and diverse megafauna assemblages. This fossil flora includes species of the following genera: Asterotheca, Marattiopsis, Cladophlebis, Coniopteris, Dictyophyllum, Thaumatopteris, Goeppertella, Dicrodium, Zuberia, Johnstonia, Xylopteris, Schleroteris, Pachydermophyllum, Hedydiphyllum, Rissikia, Ginkgoites, Báiæra, Sphenobaiera, Nilssonia, Pseudocentis, Yabeiella, Taeniopteris, Kurtziana, Solentes, Czekanowskia, Cycadocarpidium, Linguifolium, Cordaicarpus, Umkomasia, Lutanthus, Rissikistrobus, and Protocircoporoxylon.

Age of the Paso Flores Formation.—It is based on its stratigraphic relationships and its floristic content (megafauna and
microflora). Its plant assemblages are characterized by components of typical Southern Hemisphere Triassic “Dicroidium Flora”, together with the incoming morphotypes with strong Jurassic affinity. Thus, Spalletti et al. (1999, 2003), Artabe et al. (2003), and Morel et al. (1999, 2003) assigned the fossil flora of the Paso Flores Formation to the third Florian Stage (Florian floras) or Dictyophyllum tenuiserratum–Linguifolium arctum–Proto circoporoxylon marianensis Biozone of the Late Triassic (Norian–Rhaetian) age.

Zamuner et al. (2001), Zavattieri (2002), and Zavattieri and Volkheimer (2001) chronostratigraphically analyzed the distribution of the 25 species recorded for the first time in Paso Flores palynoflora at the Cañadón de Pancho locality. They assigned this microflora assemblage to the late Norian–Rhaetian age based on the co-occurrence of typical Triassic palynomorphs (Alisporites–Falci sporites microflora) together with Classopollis (= Corollina) simplex (Danzé-Corsin and Laveine, 1963) Reiser and Williams, 1969, and other Rhaetian species like Foveogleicheniidites atavus Raine in de Jersey and Raine, 1990, Foveosporites moretonensis de Jersey, 1964, Dictyophillidites atraktos Stevens, 1981 (Dipteridaceae dispersed spores), Retitriletes rosewoodensis (de Jersey, 1959) McKellar, 1974, among other species of Early Jurassic distribution (de Jersey and Raine 1990). The Classopollis (= Corollina) (Cheirolepidiaceae pollen) has been recorded previously in the type locality of the Paso Flores Formation at the Limay river (Zavattieri and Mego 2008) and in other Norian to Rhaetian units of Argentina (i.e., Tronquimalal Group, Malargüe depocenter). These Cheirolepidiaceae pollen grains are frequent to dominant in Jurassic strata of Argentina and Gondwana (Gnaedinger and Zavattieri 2017b, 2020, and references therein).
Therefore, the palaeobotanical content (macroflora and microflora) of the Paso Flores Formation is considered the youngest Triassic flora known in Argentina.

Material and methods

Eleven well-preserved medium to quite large adult frond fragment impressions were recovered. They were studied using a Leitz M50 stereoscopic microscope with an attached EC2 (LM) camera, and Nikon Coolpix P100 camera and scanning electron microscope (SEM Jeol 5800LV) at the Universidad Nacional del Nordeste (Corrientes Province, Argentina). For the description, we follow the terminology of Choo et al. (2016) and Choo and Escapa (2018).

The palaeobotanical material is housed in the Palaeontological Collection of the Museo Municipal “Carmen Funes”, Plaza Huincul city, Neuquén Province, Argentina, under the prefix MCF-PBPH.

**Systematic palaeontology**

Class Polypodiopsida Cronquist, Takhtajan, and Zimmerman, 1966

Order Gleicheniales Schimper, 1869

Family Dipteridaceae Seward and Dale, 1901

Genus *Patagoniapteris* nov.

*PFNR*: PFN002691

**Etymology**: In reference to Patagonia the geographical region situated in south Argentina and Chile to which the Neuquén Province belongs, and *pteris* refers to ferns.

**Type species**: *Patagoniapteris artabeae* sp. nov., monotypic.

**Diagnosis**.—Fan-shaped fronds dissected in two equal and opposite rachial arms have numerous primary segments (more than 18). Frond lamina between the primary segments fused at the base, the remaining part free, lanceolate, with undulated to deeply dissected margins. Primary and secondary veins simple, tertiary veins dichotomize and form irregularly polygonal meshes. Exindusiate sori arranged on either side of the primary veins, and of the base of secondary veins. Circular sori have more than 45 sporangia. Annulus oblique and complete.

*Patagoniapteris artabeae* sp. nov.

*PFNR*: PFN002692

**Etymology**: In honour of Analia B. Artabe, a recognized Argentinian palaeobotanist, for her important contributions mainly to the knowledge of Triassic and Jurassic floras of Argentina.

**Type material**: Holotype: MCF-PBPH 066, fertile frond fragment impression. Paratype: MCF-PBPH 415, fertile frond fragment impression; from the type locality and horizon.

**Type locality**: Cañadón de Pancho area, south-west of the Neuquén Province, Patagonia, Argentina (Figs. 2–3).

**Type horizon**: Upper part of the Paso Flores Formation, Late Triassic (late Norian–Rhaetian).

**Material**.—MCF-PBPH 067, 069, 074, 076, 077, 078, 079, 413, 416. Sterile and fertile frond fragment impressions from the type locality and horizon.

**Diagnosis**.—Fan-shaped fronds dissected in two equal and opposite rachial arms, each one having more than 18 primary segments. The primary segments fused up to 1/3 of the total length of the preserved lamina, and the remaining part free, lanceolate, showing undulated to deeply dissected margins (less than 2/3 length of the secondary veins). Primary and secondary veins simple, whereas tertiary veins dichotomized and forming irregular polygonal meshes. Exindusiate sori arranged in two regular rows on side of the primary veins,
Fig. 5. Drawing of the dipteridacean fern *Patagoniapteris artabeae* gen. and sp. nov. from the upper Norian–Rhaetian, Upper Triassic Paso Flores Formation, Neuquén Province, Argentina. A. MCF-PBPH 066, basal portion of a rachial arm with the primary segments fused lamina. B. MCF-PBPH 415, primary segments of fertile frond with deeply dissected lobes. C. Hypothetical reconstruction of the frond.

Fig. 4. Dipteridacean fern *Patagoniapteris artabeae* gen. and sp. nov. from the upper Norian–Rhaetian, Upper Triassic Paso Flores Formation, Neuquén Province, Argentina. A. MCF-PBPH 066, part of the frond showing fused and dissected portions. A1, A2, basal portion of a rachial arm with the lamina of the primary segments fused. A3, detail showing primary segments with the dissected portion of the lamina. B. MCF-PBPH 415, primary segments of fertile frond; B1, B2, deeply dissected lobes. C, D. Primary segments of sterile frond. MCF-PBPH 067d (C) and MCF-PBPH 069 (D). C, D1, deeply dissected lobes. D2, detail of a lobe of the primary segment, showing secondary and tertiary veins. Scale bars 10 mm.
and on both sides of the base of secondary veins. Circular sori with 45–60 sporangia. Annulus oblique and complete.

Description.—Sterile and fertile frond fragments bilaterally symmetrical, large in size, maximum preserved length ca. 20 cm. Stipe up to 7 mm in width, unknown length; apex divided into two equal rachial arms on both sides, each one is 3–4 mm wide and has at least 18 primary segments (Figs. 4A1, A2, 5A, C). Primary segments attach directly the stipe, which is twisted forming a fan-structure in conjunction with the veins of successive primary segments. The lamina frond is fused up to 1/3 of the total length preserved in an extension of at least 6.5 cm long. This fused part has polygonal venation meshes (Figs. 4A1, A2, 5B, C). The free part of the frond is dissected (Figs. 4A3, 5B). The primary segments are lanceolate, showing undulated to deeply dissected margins (less than 2/3 in length of the secondary veins) (Figs. 4B1, B2, C, D1, D2, 5C). The lobes are triangular up to 2–2.5 cm long × 1.5 cm wide with acute apex (Figs. 4B1, B2, C, D1, D2, 5C).

Fig. 7. Dipteridaceous fern *Patagoniapteris artabeae* gen. and sp. nov. from the upper Norian–Rhaetian, Upper Triassic Paso Flores Formation, Neuquén Province, Argentina. A. MCF-PBPH 066, view of the sori. A1–A5, details of the sori with and without sporangia (arrow). Scale bars 1.5 mm.

Fig. 6. Dipteridaceous fern *Patagoniapteris artabeae* gen. and sp. nov. from the upper Norian–Rhaetian, Upper Triassic Paso Flores Formation, Neuquén Province, Argentina. A. MCF-PBPH 074, detail of polygonal pattern venation of the primary segment fused lamina. B. MCF-PBPH 415, polygonal pattern of venation. B1, B2, details of the secondary and tertiary veins and polygonal meshes. B3, details of the dichotomized tertiary veins and polygonal meshes shown. C, D. Shape and distribution of sori. MCF-PBPH 076 (C) and MCF-PBPH 066 (D). View of the sori on both side of the primary veins, as well as at the base of the deep lobes on both sides of the secondary veins (arrows). Scale bars 5 mm.
Table 1. Comparison of the morphology fronds and reproductive organ features between fossil and current genera of the Dipteridaceae (data taken from Seward and Dale 1901; Herbst 1992a, b; Smith et al. 2006; Choo and Escapa 2018; Zhang et al. 2013). Grey areas indicate features of different genera shared with the Paso Flores Formation specimens.

|                      | Thaumatopteris | Sewardalea | Clathropteris | Digitopteris | Dictyophyllum | Hausmannia | Dipteris | Patagoniapteris gen. nov. |
|----------------------|----------------|------------|---------------|--------------|---------------|-----------|----------|--------------------------|
| Initial stipe dichotomy | anisotomous     |            |               |              |               |           |          | stipe with an isomomous initial dichotomy |
| Rachial arms          | two unequal     |            |               |              |               |           |          | two equal and opposite   |
| Frond lamina          |                |            |               |              |               |           |          |                          |
|                      | dissected; primary segments forming deep lobes between secondary veins (more than 2/3 length of the secondary veins) | dissected; primary segments linear and free | primary segments fused up to 1/3 of the total length of the primary veins; apical portion dissected | primary segments basally connate; apical portion free, linear and entire-margins | dissected; primary segments lanceolate or oblanceolate | entire or slightly dissected, lobed and/or crenate margins | deeply dissected primary segments | primary segments fused up to 1/3 of the total length of the primary veins, the apical portion is free; primary segment margins are undulated to deeply dissected between secondary veins (more than 2/3 length of the secondary veins) |
| Primary segments per rachial arm | 3–6 | more than 12, but up to over 100 | 3–12 | less than 10 | less than 15 | bilobulate | 4–5 (1 or more times dichotomized in 8–20 or more) | more than 18 |
| Primary and secondary veins | simple | simple | simple | simple | primary veins simple; secondary veins dichotomized | simple | primary veins dichotomized | primary veins dichotomized | simple |
| Venation              | irregularly polygonal | irregularly polygonal | regular orthogonal | polygon | irregularly polygonal, never regular orthogonal shape | quadrangular to polygonal | polygon | irregularly polygonal |
| Sori distribution     | scattered | no data | in the rectangular areole | the complete lamina is covered by sori, one in each areole | either side of primary and secondary veins | the complete lamina is covered by sori, grouped by 4 in each areole | the complete lamina is covered by sori, except D. lobbiana in a single row on each side of the primary veins | a single row on each side of the primary veins, as well as on both sides of the base of the secondary veins |
| Sporangia             | 1–4 | – | few | – | see Table 2 | 4–6 | see Table 2 | >45 |
| Age                   | Middle Triassic–Early Jurassic | Late Triassic–Early Jurassic | Late Triassic–Early Jurassic | Late Triassic | Middle Triassic–Early Jurassic | Middle Triassic–Early Cretaceous | Extant | Late Triassic |

5B). Primary and secondary veins are simple, tertiary veins are dichotomized forming irregular polygonal meshes (Figs. 4B–D, 5, 6A, B). Primary veins are up to 1.4 mm wide in the base and up to 0.5 mm wide in the preserved apical portion. The primary veins show scalariform pitting in the primary xylem tracheids (Fig. 8A1, A3). Secondary veins of the deep lobes are subopposite to alternate, and depart at angles between 45–70º and are 0.2–0.3 mm in wide. Between two successive secondary veins, there are other veins of the same thickness, reaching up to each interlobe sinus (Figs. 4B–D, 5, 6B, C). Tertiary veins are opposite, 0.1–0.3 mm, and departing from secondary veins at 50–60º, and dichotomize twice towards the margin of the lobes, and between them are horizontal veins forming a net (Fig. 6B). Sori are exindusiate and arranged on both sides of the primary veins, sometimes at the base on both sides of the secondary veins (Fig. 6C, D).
Table 2. Comparison of sori among Dipteridaceae taxa (updated from Ōishi and Yamasita 1936; Webb 1982; Herbst 1992a, b; Guignard et al. 2009).

| Species                        | Shape                      | Arrangement                                                                 | Number of sporangia |
|--------------------------------|----------------------------|-----------------------------------------------------------------------------|---------------------|
| *Dictyophyllum bremerense*     | elongated or circular      | elongated on either side of primary and secondary veins; circular well-spaced or crowded on abaxial surface | at least 30         |
| *Dictyophyllum davidii*        | elongated or circular      | elongated on either side of primary and secondary veins; circular scattered over abaxial surface | at least 25         |
| *Dictyophyllum ellenbergii*    | elongated or circular      | elongated on either side of primary veins; circular adjacent to secondary veins | –                   |
| *Dictyophyllum rugosum*        | circular                   | densely crowded over whole abaxial surface                                  | 1–4                 |
| *Dipteris lobbiana*            | circular                   | well-spaced over whole abaxial surface; rarely slightly elongated along primary veins | 3–30 (average 12–15) |
| *Thaumatopteris brauniana*     | circular                   | very crowded over whole abaxial surface                                    | 20–40               |
| *Thaumatopteris rocalanquensis*| circular                   | well-spaced, over whole abaxial surface and 2 to 3 between the secondary veins | –                   |
| *Sewardalea nathorstti*        | circular or variable       | well-spaced over whole abaxial surface predominantly near primary and secondary veins | 5–8                 |
| *Sewardalea exile*             | circular                   | very crowded over whole abaxial surface between tertiary veins             | 4–7 or 3–5          |
| *Sewardalea falcata*           | circular                   | crowded over whole abaxial surface                                          | 6–7                 |
| *Clathropteris menisoides*     | circular                   | scattered on the abaxial surface                                             | 8–15                |
| *Clathropteris obovata*        | circular                   | scattered on the abaxial surface                                             | 10–20               |
| *Dipteris conjugata*           | variable                   | crowded over whole abaxial surface                                           | 12–17               |
| *Dipteris lobbiana*            | circular                   | a single row on either side of the primary veins                            | average 30          |
| *Patagoniapteris arteae*       | circular                   | a single row on each side of the primary veins, as well as on both sides of the base of the secondary veins | 45–60 (average 52)  |

Sori are circular, 1.22–1.45 mm; in each sorus 45–60 sporangia (preserved), although larger amount of sporangia is not excluded (Figs. 6C, D, 7, 8A–8). The sporangia are 142–197 µm in diameter and have a ring-like annulus. The annulus is oblique, complete, and composed of more than 12 cells (Fig. 8A–8).

Remarks.—Comparisons with fossil genera: In the Dipteridaceae, the morphological and reproductive features commonly used for the diagnoses of genera and species include: the number, size, and shape of primary segments (Ōishi 1932; Frenguelli 1941; Ash 1970; Choo and Escapa 2018), tooth shape (Ōishi 1932), the angle of insertion of the secondary, tertiary and quaternary veins (Heer 1877; Ōishi 1940; Schweitzer et al. 2009), and the size and number of sporangia per sori (Ash 1970). The appearance of tertiary veins is an additional feature sporadically mentioned in the discussions of species boundaries, but rarely used as a diagnostic generic feature (Herbst 1966; Kon’no 1968; Choo et al. 2016).

Table 1 shows that the Paso Flores specimens share morphological features with those established for the Dipteridaceae: (i) with *Clathropteris* and *Digitopteris* they share frond lamina between primary segments fused up to 1/3 of the total length of the primary veins and the free apical portion; (ii) with *Thaumatopteris* and with some species of the *Dictyophyllum* they share the presence of deep lobes between secondary veins (more than 2/3 of the length of them); (iii) with *Sewardalea* they share the presence of numerous primary segments (more than 12); (iv) with *Thaumatopteris*, *Dictyophyllum*, and *Digitopteris* they share the polygonal venation meshes.

The presence of sori in the Dipteridaceae classification, has different considerations. Harris (1931) showed an evolutionary sequence from well-defined sori of acrosticoid disposition, whereas Ōishi and Yamasita (1936) considered that: (i) the shape of sori are basically the same, being circular or rounded, and the size of the sori decrease in the following order of the genera: *Hausmannia*–*Thaumatopteris*–*Dictyophyllum*; (ii) the number of sporangia is not constant, even in one species, however, it decreases according to the following approximate genera order: *Thaumatopteris*–*Clathropteris*–*Hausmannia*–*Dictyophyllum*–*Camptopteris*; (iii) the size of the sporangia decreases according to the increase in size of sori; (iv) the annulus is sometimes oblique and mostly complete and each annulus usually has up to 30 cells. On the other hand, in the systematic review of the family, Herbst (1992a, b) remarked that he does not use the characteristics of the sori (or sporangia) because they are similar in all Dipteridaceae. In genera such as...
Dictyophyllum and Thaumatopteris (see Tables 1 and 2) the distribution of the sporangia can be: in acrosticoid form (regularly or irregularly distributed) or grouped in sori (also regularly or irregularly) (Ôishi and Yamasita 1936).

So far, there are several species described in the literature that provide some descriptions of reproductive structures (Table 2). The species that have sori distributed in the primary veins are: Dictyophyllum davidii Walkom, 1917, and D. bremerense Shirley, 1898. The first mentioned species differs from the material of Argentina in the elongated shape of the sori, and from the second one by the number of sporangia per sori (Table 2).

Based on the morphological features of the fronds and the distribution of sori and number of sporangia, the Paso Flores Formation specimens are clearly distinct from other dipteridacean genera, justifying the establishment of Patagoniapteris artabeae gen. and sp. nov. (Tables 1, 2).

Dictyophyllum tenuifolium (Stipanicic and Menéndez, 1949) Bonetti and Herbst, 1964, and Dictyophyllum (Thaumatopteris) rothi (Freygülelli, 1941) Bonetti and Herbst, 1964, were previously described from the Paso Flores Formation. Patagoniapteris artabeae gen. and sp. nov., differs from the first mentioned species in the number of sporangia per sori (Table 2). It differs from the second species because this new taxon herein described has partially fused lamina fronds and more than 15 rachial arms (Table 1).

Comparisons with extant genera: Seward and Dale (1901) proposed that the genus Dipteris should be representative of the family Dipteridaceae. Morphologically, the leaf architecture of Dipteris is quite characteristic. The fronds branch dichotomously with veins that split forming a mesh pattern. Sori are exindusiate and arranged following the primary veins or distributed on abaxial surface on both sides of the primary veins. The annulus of the sporangia is oblique.

The distribution of the sori in Dipteris conjugata Reinwardt, 1828, covers the whole abaxial surface of the large frond, although they are frequently concentrated on either side of the primary veins with 12–17 sporangia (Table 2). In Dipteris lobbiana (Hooker, 1853) Moore, 1857, the sori are situated on the abaxial surface of the narrow lamina in two regular rows on either side of the primary veins, which contains an average of 35 sporangia (Armour 1907). The other species of Dipteris either resemble one of these two species or form intermediates between them (Armour 1907; Webb 1982) (Table 2).

Webb (1982) interpreted that in modern species, this division appears to be mainly a function of the width of the lamina on either side of the primary veins (wide in Dipteris conjugata, narrow in Dipteris lobbiana). This is not the case in Patagoniapteris artabeae gen. and sp. nov. because it has a broad lamina like Dipteris conjugata and sori distribution and number of sporangia similar to Dipteris lobbiana (Tables 1, 2).

Stratigraphic and geographic range.—Type horizon and locality only.

Discussion

In the recent phylogenetic analysis, Choo and Escapa (2018) provided an enlightening analysis into the evolution of Gleicheniales as a whole, and of Dipteridaceae in particular. Patagoniapteris artabeae gen. and sp. nov. could be considered a new transitional form among Dipteridaceae because it shares some morphological and reproductive features with several fossil genera and the current Dipteris (see Table 1 and 2): (i) with Thaumatopteris it has a common deep lobes between secondary veins, and with Sewardalea it has in common numerous primary segments; but neither genera have the fused base of the lamina; (ii) Clathropteris, many Dictyophyllum and Digitopteris species have primary segments up to 1/3 fused at the base. However, the first genus is noted for its orthogonal venation and the second has less than 15 rachial arms, while the third has apical portion segments with whole margins. Dipteris, on the other hand, has a sori distribution and a number of sporangia similar to Patagoniapteris artabeae gen. and sp. nov., but differs from this new taxon because the frond is deeply dissected and the primary veins are dichotomized (Fig. 5C, Tables 1, 2; Choo and Escapa 2018: fig. 3).

Bodnar et al. (2018) pointed out that during the Triassic, the Dipteridaceae ferns had a restricted spatio-temporal distribution in Argentina related to humid conditions: (i) the earliest occurrence was recorded in the Barreal Formation of late Anisian–early Ladinian age, Barreal-Calingasta Depocenter, San Juan Province, Cuyana Basin, were less diverse and their frond size smaller, and (ii) in the late Norian–Rhaetian interval these ferns had their maximum diversification with the development of individuals with large fronds, registered at the north-eastern end of the Neuquén Basin, in the Malargüe Depocenter (Tronquimalal Group), Mendoza Province (Gnaedinger and Zavattieri 2020, and references therein), and in the south-southwestern region of the basin, at Paso Flores depocenter (Gnaedinger and Zavattieri 2017b, and references therein). Such diversity and morphological conditions of the Dipteridaceae family could indicate that at the end of the Triassic, the humidity regimen was more benign than in the Middle Triassic (Bodnar et al. 2018).

Spalletti et al. (2003) and Artabe et al. (2003) analyzed the palaeogeographical reconstructions and the distribution of plants within the phytogeographic Triassic provinces of south-western Gondwana. Towards the end of the Triassic, in Argentina, the third recognized floristic event, the Florian Stage (late Norian–Rhaetian) (Spalletti et al. 1999) was characterized by an important change in plant composition, when the long-lasting endemic Triassic Dicroidium-dominated communities declined and the incoming of taxa that persisted during the Jurassic occurred. At this time, the basins located on the western margin of southern South America had a seasonal temperate-warm and subtropical humid to sub-humid climate with marine influence from the west, allowing for more humid conditions,
characterized by monsoonal climates (Artabe et al. 2001, 2003; Spalletti et al. 2003).

The Dipteridaceae grew in herb-shrub and tree communities developed on floodplains of braided and meandering fluvial systems (Artabe et al. 2001). The record of the new taxon, *Patagoniapteris artabaeae* gen. and sp. nov. in the uppermost section of the Paso Flores Formation in the Cañadón de Pancho area, is congruent with above the mentioned pattern of distribution and climatic and environmental conditions.

**Conclusions**

A new taxon, *Patagoniapteris artabaeae* gen. and sp. nov. is established based on sterile and fertile fronds showing affinities with fossil and extant Dipteridaceae from the Paso Flores Formation, late Norian–Rhaetian, Neuquén Basin, Argentina. *Patagoniapteris artabaeae* gen. and sp. nov. shows a unique combination of morphological and reproductive features that do not belong to any described genera of Dipteridaceae. It comprises an interesting combination because it shares some morphological frond features with the fossil genera *Clathropteris*, *Digitopteris*, *Thaumatopteris*, *Sewardalea*, and with some species of *Dictyophyllum*. It also has some reproductive features in common with the extant species *Dipteris lobbiana* (e.g., sori distribution and number of sporangia).

*Patagoniapteris artabaeae* gen. and sp. nov. represents a new transitional form which shows an increasing complexity in the venation patterns and laminal fusion. In addition, it shows an increase in the number of sporangia per sori, only observed in the Recent genus *Dipteris*. The Cañadón de Pancho plant assemblages are late Norian–Rhaetian in age, being the youngest fossil flora recorded from Argentinean Triassic basins to date. The record of the new taxon, *Patagoniapteris artabaeae* gen. and sp. nov. in the uppermost section of the Paso Flores Formation at Cañadón de Pancho area, is consistent with the higher diversity of Dipteridaceae at the end of the Triassic in Argentina. In the western margin of southern South America, during “Florian Stage” (late Norian–Rhaetian), the climate was temperate-warm seasonal with humid conditions influenced by the penetration of maritime air masses from the west (Spalletti et al. 2003). These humid to sub-humid regimes favoured the development of the Dipteridaceae with large fronds.

**Acknowledgements**

We wish to thank Rodolfo Coria (CONICET-Neuquén, Argentina), who led the Project “Searching for primitive dinosaurs in the Late Triassic of Northern Patagonia, Neuquén, Argentina”, which was financed by The Dinosaur Society (1996–1997) and in the frame of which the plant material herein described was collected. Furthermore, as Managing Director R. Coria allowed us to access to the Paleontological Collection of the Museo Municipal “Carmen Funes”, Plaza Huincul, Argentina, to study the plant assemblages housed therein. We also extend our appreciation and gratitude to Flavio Bellardini (Paleontological Curator of the Museo Municipal “Carmen Funes”), for his constant encouragement and assistance with the materials and repository. Grateful acknowledgement is extended to Eugenia Zavattieri (illustrator and graphic designer, Puerto Madryn, Argentina) for the improvement of the photo-micrographic figures and plates. Thanks are due to an anonymous reviewer and Maria Barbacka (Hungarian Natural History Museum, Budapest, Hungary) for their insightful comments and corrections that improved the manuscript. The manuscript greatly benefited from thorough review by Roslyn Lacey Wallace (Houston, USA) who made extensive English language and stylistic improvements. This research has been partially funded by the Agencia Nacional de Promoción Científica y Tecnológica, Argentina (research grants ANPCYT–FONCyT, Argentina, PICT 2011-2546 to AMZ). Parts of the research that led to the study of this material were financially supported by the Secretaría General de Ciencia y Técnica, SGCTY-UNNE, Corrientes Province (PI 2018-2022, FOI3 to SCG) and CONICET (PIP 2014-2016, 112 201301 00317 to SCG-AMZ).

**References**

Andrae, K.J. 1853. Beiträge zur Kenntnis der fossilen Flora Siebenburgens und des Banates. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt 2: 1–48.

Armour, H.M. 1907. On the sorus of *Dipteris*. New Phytologist 6: 238–244.

Arrondo, O.G. and Pettigrew, B. 1982. Revisión del género *Goeperttiella* Ōishi y Yamashita emend (Goeperttilloideae–Dipteridaceae). *Ameghiniana* 19: 67–78.

Arrondo, O.G., Spalletti, L.A., Morel, E., and Gana uz, D.G. 1991. The sedimentological and paleobotanical characteristics of an Upper Triassic–Lower Liassic basin in southwestern Patagonia (Argentina). In: H. Ulbrich and A.C. Rocha Campos (eds.), *Gondwana Symposium 7*th Proceedings, Instituto de Geociencias, *Universidade de Sao Paulo* 714: 517–532.

Artabe, A.E., Morel, E.M., and Spalletti, L.A. 2001. Paleoecología de las floras triásicas argentinas. In: A.E. Artabe, E.M. Morel, and A.B. Zamuner (eds.), *El Sistema Triásico de Argentina*, 199–225. Fundación Museo de La Plata “Francisco Pasca só Moreno”, La Plata.

Artabe, A.E., Morel, E.M., and Spalletti, L.A. 2003. Caracterización de las provincias fitogeográficas triásicas del Gondwana Extratropical. *Ameghiniana* 40: 387–405.

Artabe, A.E., Morel, E.M., and Zamuner, A.B. 1994. Estudio paleobotánico y taxonómico en la Formación Paso Flores (Triásico Superior), en el Cañadón de Pancho, Neuquén, Argentina. *Ameghiniana* 31: 153–160.

Ash, S.R. 1970. *Ferns From the Chinle Formation (Upper Triassic) in the Fort Wingate Area, New Mexico.* 72 pp. United States Government Printing Office, Washington D.C.

Bodnar, J., Drovandi, J.M., Morel, E.M., and Gana zu, D.G. 2018. Middle Triassic dipterid ferns from west-central Argentina and their relationship to paleoeclimatic changes. *Acta Palaeontologica Polonica* 63: 397–417.

Bonfleur, B. and Kerp, H. 2010. The first record of the dipterid fern leaf *Clathropteris* Bronnhiart from Antarctica and its relation to *Polyphe- cels stromensis* Yao, Taylor and Taylor nov. emend. Review of Palaeo- botany and Palynology 160: 143–153.

Bonetti, M.I.R. and Herbst, R. 1964. Dos especies de *Dictyophyllum* del Triásico de Paso Flores. Provincia del Neuquén, Argentina. *Ameghiniana* 3: 273–279.

Brauns, D. 1862. Der Sandstein bei Seinstedt unwesentlich Falbitsteine und die in ihm vorkommenden Pflanzenreste. *Palaeontographica* 9: 47–62.

Brongniart, A. 1853. Beiträge zur Kenntnis der fossilen Flora Siebenburgens und des Banates. Abhandlungen der Kaiserlich-Königlichen Geologischen Reichsanstalt 2: 1–48.

Bodnar, J., Drovandi, J.M., Morel, E.M., and Gana zu, D.G. 2018. Middle Triassic dipterid ferns from west-central Argentina and their relationship to paleoeclimatic changes. *Acta Palaeontologica Polonica* 63: 397–417.

Bonfleur, B. and Kerp, H. 2010. The first record of the dipterid fern leaf *Clathropteris* Bronnhiart from Antarctica and its relation to *Polyphe- cels stromensis* Yao, Taylor and Taylor nov. emend. Review of Palaeo- botany and Palynology 160: 143–153.

Bonetti, M.I.R. and Herbst, R. 1964. Dos especies de * Dictyophyllum* del Triásico de Paso Flores. Provincia del Neuquén, Argentina. *Ameghiniana* 3: 273–279.

Brauns, D. 1862. Der Sandstein bei Seinstedt unwesentlich Falbitsteine und die in ihm vorkommenden Pflanzenreste. *Palaeontographica* 9: 47–62.

Brongniart, A. 1828. *Prodrome d’une histoire des végétaux fossiles.* vii + 223 pp. Levrault, Paris.
De Jersey, N.J. 1964. Triassic spores and pollen from the Bundamba.

Ganuza, D.G., Spalletti, L.A., Morel, E.M., and Arrondo, O.G. 1995. Paleo-

Ganuza, D., Morel, E.M., Spalletti, L.A., and Arrondo, O.G. 1992. Las plan-

Dictyophyllum

Fabre, J. and Greber, C. 1960. Presence d’un

dans la flo-

Dunker, W. 1846.

González Díaz, E.F. 1982. Sedimentitas del Triásico superior continental

Die Gattungen der fossilen Pflanzen vergli-

Gnaedinger, S. and Zavattieri A.M. 2020. Coniferous woods from the Up-

Gnaedinger, S. and Zavattieri A.M. 2017a. A new name for

Gnaedinger, S. and Zavattieri A.M. 2017b. Propuesta de clasificación de las Dipteridaceae con un

Atlas de las especies argentinas. D’Orbignyana 6: 1–71.

Herbst, R. 1992b. Propuesta de clasificación de las Dipteridaceae. Publi-

Herbst, R. 1993. Dipteridaceae (Filicales) del Triásico del Arroyo Llantenes (provincia de Mendoza) y del Paso Flores (provincia del Neuquén), Argentina. Ameghiniana 30: 155–162.

Hooker, W.J. 1853. Hooker’s Journal of Botany and Kew Garden Miscell-

Kon’no, E. 1968. Some Upper Triassic species of Dipteridaceae from Japan

Kramer, K.U., Green, P.S., and Gotz, E. 1990.

Kustatscher, E. and van Konijnenburg-van Cittert, J.H.A. 2011. The ferns

Herbst, R. 1965. La flora fósil de la formación Roca Blanca, provincia de Santa Cruz, Patagonia. Con consideraciones geológicas y estratigra-

Hooker, W.J. 1853. Hooker’s Journal of Botany and Kew Garden Miscell-

Harris, T.M. 1931. The fossil flora of Scoresby Sound East Greenland.

Henry and Cohen, Bonn.

González Díaz, E.F. 1982. Sedimentitas del Triásico superior continental en el valle del Río Collón Curá, entre los arroyos Quemquemtreu y Calefú, Provincia de Neuquén. Revista de la Asociación Geológica Argentina 37: 214–220.

Guignard, G., Wang, Y., Ni, Q., Tian, N., and Jiang, Z. 2009. A dipteridea-

cern fern with in situ spores from the Lower Jurassic in Hubei, China. Review of Palaeobotany and Palynology 156: 104–115.

Gulison, C.A., Gutiérrez Pleimling, A.R., and Digregorio, R.E. 1984. Esquema estratigráfico de la secuencia jurásica del oeste de la pro-

vinca del Neuquén. 9º Congreso Geológico Argentino, San Carlos de Bariloche. Actas 1: 236–259.

Harris, T.M. 1931. The fossil flora of Scoresby Sound East Greenland. Part 1: Cryptogams (exclusive of Lycopodiales). Mededelser om Grøn-

GNAEDINGER AND ZAVATTIERI—LATE TRIASSIC DIPTERIDACEAN FERN FROM ARGENTINA 899

Geological Survey of Queensland Publication Group.

Gnaedinger, S. and Zavattieri A.M. 2017b. Propuesta de clasificación de las Dipteridaceae. In: V. Ramos (ed.), Geología y recursos naturales de la Provincia de Río Negro. Relatorio 9º Congreso Geológico Argentino 1: 139–162.

Lagarreta, L. and Uliana, M.A. 1996. The Jurassic succession in west-cen-

tral Patagonia, Argentina: implications for taxonomy and palaeo-

Gnaedinger, S. and Zavattieri A.M. 2017a. A new name for

De Jersey, N.J. 1959. Jurassic spores and pollen grains from the Rosewood Coalfield. Queensland Government Mining Journal 60: 346–366.

De Jersey, N.J. 1964. Triassic spores and pollen from the Bundamba Group. Geological Survey of Queensland Publication 321: 1–21.

De Jersey, N.J. and Raine, J.J. 1990. Triassic and earliest Jurassic mios-

De Jersey, N.J. 1959. Jurassic spores and pollen grains from the Rosewood Coalfield. Queensland Government Mining Journal 60: 346–366.

Dilma, O.R., Lizuaín, A., and Núñez, E. 1984. La cobertura sedimentaria desempeño de la Cuenca Neuquina (Triásico superior–Terciario inferior). Serie Corre lación Geológica 6: 37–53.

D’Elia, L., Muravchik, M., Franzese, J.R., and López, L. 2012. Tectono-

Dunko, J. and Greber, C. 1960. Presence d’un Dictyophyllum dans la flo-

Dunker, W. 1846. Monographie der norddeutschen Wea
denbildung. 85 pp. Oehme, Braunschweig.

Fabre, J. and Greber, C. 1960. Presence d’un Dictyophyllum dans la flo-

Fengielli, J. 1937. La flóra Jurásica de Paso Flores en el Neuquén, con referencias a la de Piedra Pintada y otras floras jurásicas argentinas. Revista del Museo de La Plata (Nueva Serie) 1. Paleontología 3: 67–108.

Fengielli, J. 1941. Las Camptopteridias del Lias de Piedra Pintada en el Neuquén (Patagonia). Revista del Museo de La Plata, nueva serie, Paleontología 27: 27–57.

Ganuza, D., Morel, E.M., Spalletti, L.A., and Arrondo, O.G. 1992. Las plan-

Ganuza, D., Morel, E.M., Spalletti, L.A., and Arrondo, O.G. 1992. Las plan-

Ganuza, D., Morel, E.M., Spalletti, L.A., and Arrondo, O.G. 1992. Las plan-

Ganuza, D.G., Spalletti, L.A., Morel, E.M., and Arrondo, O.G. 1995. Paleo-

Ganuza, D.G., Spalletti, L.A., Morel, E.M., and Arrondo, O.G. 1995. Paleo-

Gauzán, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, J. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.

Danzé-Corsin, P. and Laveine, J.P. 1963. Microflora. In: P. Briche, P. Danzé-

Cronquist, A., Takhtajan, A., and Zimmerman, W. 1966. On the higher taxa of Embryobionta. Taxon 15: 129–134.
Moore, T. 1857. *Index Filicum*. 204 pp. William Pamplin, London.

Morel, E.M., Ganuza, D.G., and Zúñiga, A. 1999. Revisión paleoflorística de la Formación Paso Flores, Triásico superior de Río Negro y del Neuquén. *Revista de la Asociación Geológica Argentina* 54: 389–406.

Morel, E.M., Spalletti, L.A., Arrondo, O.G., and Ganuza, D.G. 1992. Los estratos plantíferos de la Formación Paso Flores (Triásico Superior) de las Lomas y Cañadón de Ranquel Huao, provincia del Neuquén, Argentina. *Revista del Museo de La Plata (Nueva Serie) 9* Paleontología 58: 199–221.

Nathorst, A.G. 1878. Floran vid Höganäs och Helsingborg. *Kungliga Svenska Vetenskapsakademiens Handlingar* 16: 1–53.

Nullo, F.E. 1979. Descripción geológica de la Hoja 39c, Paso Flores, Provincia de Río Negro. *Boletín Servicio Geológico Nacional* 167: 1–70.

Ôishi, S. 1940. The Mesozoic floras of Japan. *Journal of the Faculty of Science, Hokkaido Imperial University Series 4, Geology and Mineralogy* 1: 257–379.

Ôishi, S. 1940. Los Mesozoic floras of Japan. *Journal of the Faculty of Science, Hokkaido University. Series 4, Geology and Mineralogy* 5: 123–480.

Ôishi, S. and Yamashita, K. 1936. On the fossil Dipteridaceae. *Journal of the Faculty of Science, Hokkaido Imperial University Series 4, Geology and Mineralogy* 3: 135–184.

Popp, O. 1863. Der Sandstein von Jägersberg bei Forchheim und die in ihm vorkommende Pflanzen. *Neues Jahrbuch für Mineralogie und Geologie* 1863: 399–417.

Pott, C., Bouchal, J.M., Choo, T.Y.S., Yousif, R., and Bonfleur, B. 2018. Ferns and fern allies from the Carnian (Upper Triassic) of Lunz am See, Lower Austria: A melting pot of Mesozoic fern vegetation. *Palaeontographica, Abteilung B* Palaeobotanik – Palaeophytology 297 (1–6): 1–101.

Presl, C.B. 1851. Epimeliales Botanice Abhandlungen der Böhmischen Ge- sellchaft der Wissenschaften 6: 361–624.

Rees, P.M. 1993. Dipterid ferns from the Mesozoic of Antarctica and New Zealand and their stratigraphical significance. *Palaeontology* 36: 637–656.

Rees, P.M. and Cleal, C.J. 2004. Lower Jurassic ferns from Hope Bay and Botany Bay, Antarctica. *Special Papers in Palaeontology* 72: 1–90.

Reinwardt, C.G.C. 1828. Nova Plantarum indicarum genera. *In: C.F. Hornschuch* (ed.), Sylloge Plantarum Novarum itemque minus cognitarum, Tomus secundus, 3. Regensburgische Botanische Gesellschaft, Regensburg.

Reiser, R.F. and Williams, A.J. 1969. Palynology of the lower Jurassic sediments of the northern Surat Basin. *Geological Survey of Queens- land* 399, Palaeontological Papers 15: 1–24.

Schimper, W.P. 1869–1874. Traité de paléontologie végétale, ou, la flore du monde primitif dans ses rapports avec les formations géologiques et la flore du monde actuel. Vol. 1: iv+738 pp.; Vol. 2: 996 pp.; Vol. 3: 896 pp. Baillièr, Paris.

Schweitzer, H.J., Schweitzer, U., Kirchner, M., Van Konijnenburg-van Cittert, J.H.A., Van der Burgh, J., and Ashraf, R.A. 2009. The Rhaeto–Jurassic flora of Iran and Afghanistan. 14. Pteridophyta–Leptosporangiateae. *Acta Palaeontologica Polonica* 54: 257–379.

Shirley, J. 1898. Additions to the flora of Queensland. Mainly from the Ipswich Formation, Trias–Jura System. *Bulletin Geological Survey of Queensland* 7: 1–25.

Smith, A.R., Pryer, K.M., Schuettpeilz, E., Korall, P., Schneider, H., and Wolf, P.G. 2006. A classification for extant ferns. *Taxon* 55: 705–731.

Spalletti, L.A., Arrondo, O.G., Morel, E.M., and Ganuza, D.G. 1988. Estudio sedimentológico y paleoflorístico de la Formación Paso Flores – Triásico superior – en el sector occidental del Macizo Norpatagónico, Argentina. *Actas 5º Congreso Geológico Chileno* 2: 395–413.

Spalletti, L.A., Arrondo, O.G., Morel, E.M., and Ganuza, D.G. 1990. Los depósitos fluviales de la cuenca Triásica superior en el sector noroeste del Macizo Norpatagónico. *Revista de la Asociación Geológica Argentina* 43 (for 1988): 544–557.

Spalletti, L.A., Artabe, A.E. and Morel, E.M. 2003. Geological factors and evolution of southwestern Gondwana Triassic plants. *Gondwana Research* 6: 119–134.

Spalletti, L., Artabe, A.E., Morel, E.M., and Brea, M. 1999. Biocenización paleoflorística y cronoestratigrafía del Triásico Argentino. *Ameghiniana* 36: 419–451.

Stevens, J. 1981. Palynology of the Callide Basin, east-central Queensland. *University of Queensland Papers, Department of Geology* 9: 1–35.

Stipanic, P.N. and Menéndez, C.A. 1949. Contribución al conocimiento de la flora fósil de Barreal (provincia de San Juan). I. Dipteridaceae. *Boletín de Informaciones Petroleras, Buenos Aires* 24: 44–73.

Taylor, E.L., Taylor, T.N., and Krings, M. 2009. *Paleobotany: The Biology and Evolution of Fossil Plants*. 2nd Edition. 1230 pp. Academic Press, Amsterdam.

Tidwell, W.D. and Ash, S.R. 1994. A review of selected Triassic to Early Cretaceous ferns. *Journal Plant Research* 107: 417–442.

Turtunanova-Ketova, A.I. 1962. New genus of ferns from the Mesozoic sediments of Kazakhstan [in Russian]. *Paleontologičeskij žurnal* 1962 (2): 145–148.

Varela, R., Basei, M.A.S., Cingolani, C.A., Siga J.O., and Passarelli, C.R. 2005. El basamento cristalino de los Andes norpatagónicos en Argentina: geocronología e interpretación tectónica. *Revista Geológica de Chile* 52: 167–187.

Vergani, G.D., Tankard, J., Belotti, J., and Welsink, J. 1995.TECTonic evolution and paleogeography of the Neuquén Basin, Argentina. *In: A.J. Tankard, R. Suárez, and H.J. Welsink (eds.), Petroleum Basins of South America. American Association of Petroleum Geologists, Memoir 62: 383–402.

Wallkom, A.B. 1917. Mesozoic floras of Queensland. Part 1: The Flora of the Ipswich and Walloon Series (c), Filiicales. *Geological Survey of Queensland* 257: 1–46.

Webb, J.A. 1982. Triassic species of *Dicytophyllum* from eastern Australia. *Alcheringa* 6: 79–91.

Zamuner, A.B. and Artabe, A.E. 1994. Estudio de un leño fósil, *Proto- circoporoxylon marianaensis* n. sp., de la Formación Paso Flores (Neotriásico), Provincia de Río Negro, Argentina. *Ameghiniana* 31: 203–207.

Zamuner, A.B., Zavattieri, A.M., Artabe, A.E., and Morel, E.M. 2001. Palaeobotánica. *In: A.E. Artabe, E.M. Morel, and A.B. Zamuner (eds.), El Sistema Triásico en Argentina*, 143–184. Fundación Museo de La Plata “Francisco Pascasio Moreno”, La Plata.

Zavattieri, A.M. 2002. Anexo 4: Microfloras. *In: P.N. Stipanicic and C.A. Marsicano (eds.), Léxico Estratigráfico de la Argentina. Vols. 2–3* Nequén. *Paleontología* 6: 119–134.

Zavattieri, A.M. and Volkheimer, W. 2008. Palynofacies analysis and sedimentary environment of Early Jurassic coastal sediments at the southern border of the Neuquén Basin, Argentina. *Journal of South American Earth Sciences* 25: 227–245.

Zeiller, R. 1903. *Flore fossile des gîtes de charbon du Tonkin*. 316 pp. Ministère des Travaux Publications, Paris.

Zhang, X.C., Kato, M., and Nooteboom, H.P. 2013. *Dipteridaceae*. *In: Z.Y. Wu, P.H. Raven, and D.Y. Hong (eds.), Flora of China, Vols. 2–3* (Pteridophytes), 116–117. Science Press, Beijing.

Zhou, N., Wang, Y., Li, L., and Zhang, X. 2015. Diversity variation and temporal-spatial distributions of the Dipteridaceae ferns in the Mesozoic of China. *Palaeoworld* 25: 263–286.