

1 **Plant diversity turnovers in the Triassic–Jurassic transition: evidence from the**
2 **paleobotanical record of Argentina**

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15 **ABSTRACT.** The Triassic–Jurassic transition was an important interval in Earth’s history,
16 which encompassed two great biotic crises: the End-Triassic mass extinction and the early
17 Toarcian Oceanic Anoxic Event. The response of the terrestrial vegetation to these crises has
18 been the subject of debate. In Argentina, a floristic turnover occurring in the Triassic-Jurassic
19 boundary has been suggested, which led to the disappearance of corystoperms and
20 peltasperms, and to a greater diversity of ferns, bennettitaleans, and conifers. Recent findings
21 have thrown into question the previous evolutionary models. To understand the terrestrial plant
22 turnovers during the Triassic-Jurassic transition in Argentina, we calculated diversity indexes,
23 survivorship proportions, and origination and extinction rates, based on the megafossil record.
24 The results indicated that, despite only one lineage disappeared in the Late Triassic
25 (Pleuromeiaceae), there was an important but gradual vegetation change in the Triassic-
26 Jurassic transition. Late Triassic diagnostic lineages extended into the Jurassic but most of
27 them with few records and became extinct in the Early Jurassic. Floristic turnovers during the
28 Jurassic, can be correlated with paleoclimatic changes. The Osmundaceae, Dipteridaceae,

29 and Bennettiales diversified in the Sinemurian. A taxa richness increase in the Toarcian is
30 recognized, marked by a conifer diversification. In the Bathonian-Kimmeridgian interval, the
31 conifers and bennettitaleans acquired greater importance. The dataset can be improved in the
32 future by intensifying fossil sampling in order to obtain more robust and precise results.

33 **Keywords.** Mesozoic, South America, Paleofloras, Extinction.

34 **RESUMEN. Cambios en la diversidad de plantas en la transición Triásico-Jurásico:**

35 **evidencia del registro paleobotánico de Argentina.** La transición Triásico-Jurásico fue un

36 intervalo importante en la historia de la Tierra, que abarcó dos grandes crisis bióticas: la

37 extinción masiva de fines del Triásico y el Evento Anóxico Oceánico del Toarciano Temprano.

38 La respuesta de la vegetación a estas crisis ha sido objeto de debate. En la Argentina se ha

39 sugerido un recambio en el límite Triásico-Jurásico, que llevó a la desaparición de

40 corystopermas y peltaspermas, y a una mayor influencia de helechos, bennettitales y

41 coníferas. Hallazgos recientes han puesto en duda los modelos evolutivos previos. Con el

42 objetivo de comprender los cambios florísticos durante la transición Triásico-Jurásico en la

43 Argentina, calculamos índices de diversidad, proporciones de supervivencia y tasas de origen

44 y extinción, a partir del registro megafosilífero. Los resultados indicaron que, aunque sólo un

45 linaje desaparece en el Triásico Tardío (Pleuromeiaceae), se produjo un cambio de vegetación

46 importante pero gradual en la transición Triásico-Jurásico. Los linajes diagnósticos del Triásico

47 Tardío pasan al Jurásico, pero la mayoría con pocos registros y se extinguen en el Jurásico

48 Temprano. Los cambios florísticos durante el Jurásico, pueden ser correlacionados con las

49 fluctuaciones paleoclimáticas. Las Osmundaceae, Dipteridaceae y Bennettitales se

50 diversifican en el Sinemuriano. Se reconoce un aumento en la riqueza taxonómica en el

51 Toarciano, marcado por la diversificación de las coníferas. En el Bathoniano-Kimmeridgiano

52 adquirieron una mayor importancia las coníferas y bennettitales. El conjunto de datos podría

53 ser mejorado en el futuro intensificando el muestreo de fósiles para obtener resultados más

54 sólidos y precisos.

55 **Palabras clave.** Mesozoico, Sudamérica, Paleofloras, Extinción.

56 **INTRODUCTION**

57 The Triassic–Jurassic transition, which extends from the Late Triassic through the
58 Early Jurassic (~237 Ma to 174 Ma), was an important interval in Earth’s history (Schoepfer
59 et al. 2022). The End-Triassic mass extinction (ETME), at ~201.5 Ma, ranks among the ‘Big
60 Five’ Phanerozoic mass extinctions (Raup and Sepkoski 1982; Bambach 2006). Significant
61 losses have been recorded in all marine habitats and some terrestrial communities (Wignall
62 and Atkinson 2020; and references cited therein). In addition, the Triassic–Jurassic transition
63 encompasses a second-order mass extinction during the early Toarcian Oceanic Anoxic
64 Event (T-OAE), at ~181 Ma (Harries and Little, 1999; Schoepfer et al., 2022).

65 The ETME was triggered by Central Atlantic Magmatic Province (CAMP) magmatism
66 (Marzoli et al. 1999), and the T-OAE by Karoo-Ferrar Large Igneous Province (KFLIP)
67 magmatism (Pálffy and Smith 2000), both associated with the stepwise disintegration of the
68 Pangean supercontinent (Schoepfer et al. 2022). Large outgassing of CO₂, CH₄, and SO₂,
69 driven by CAMP, is believed to have triggered ETME through a series of cascading
70 environmental perturbations, including changes in the global carbon cycle, rapid warming
71 leading to "hyperthermic" climate events and increased continental weathering (Fox et al.
72 2022; and references cited therein).

73 Because there are few adequate fossiliferous sections globally, the ETME and the
74 subsequent biotic recovery are not as well-known as other mass extinctions (Damborenea et
75 al. 2017). However, in recent years it gained greater interest. Numerous publications
76 increased our understanding of the event's causes, consequences, and extent (e.g. Bonis
77 and Kürschner 2012; Haworth et al. 2012; Petersen and Lindström 2012; Blackburn et al.
78 2013; Bond and Wignall 2014; Pálffy and Kocsis 2014; Van de Schootbrugge and Wignall
79 2016; Barbacka et al. 2017; Lindström 2021).

80 The impact of the ETME on terrestrial plants diversity is less understood than on
81 terrestrial animals. Estimating the severity of the crisis in land plants is complicated by
82 provinciality and conflicting records between extinctions based on fossil leaf taxa and those
83 based on spores and pollen (Bond and Wignall 2014; Lindström 2016; Barbacka et al. 2017).
84 Local extinctions in spore and pollen taxa vary globally between 17 and 73% and for most of

85 the taxa that went extinct, the parent plant affinity is not fully resolved or even unknown
86 (Lindström 2016). Likely, many of the spore-producing plants that disappeared at the end of
87 the Rhaetian or in the earliest Hettangian were plants with ecological preferences that
88 resulted in low preservation potential for their macroscopic remains, e.g. epiphytic plants or
89 plants growing in sites with little potential for fast burial (Lindström 2021).

90 The response of the terrestrial vegetation to this crisis is debated with some authors
91 suggesting that no abrupt floral extinction but gradual shifts took place across the Triassic–
92 Jurassic transition (Lucas and Tanner 2015; Barbacka et al. 2017; Cascales-Miñana, et al.
93 2016, 2018), while others present major turnovers (McElwain et al. 2007; Bonis and
94 Kürschner 2012; de Jersey and McKellar 2013; Lindström et al. 2017; Gravendyck et al.
95 2020; Zhang et al. 2020). Macroflora records from East Greenland and southern Sweden
96 show that the Late Triassic *Lepidopteris* flora was replaced by the Early Jurassic
97 *Thaumatopteris* flora (Lundblad 1959; McElwain et al. 1999, 2007; Kustatscher et al. 2018).
98 On the Southern Hemisphere record, the Triassic seed-fern dominated flora was replaced by
99 a more complex flora with conifers (cheirolepids), bennettitaleans, and new seed-ferns during
100 the Early Jurassic (Turner et al. 2009). In China, the Late Triassic *Dictyophyllum*–
101 *Clathropteris* macroflora was substituted by the Early Jurassic *Ptilophyllum*–*Coniopteris* flora
102 (Ye et al. 1986). In European successions, Rhaetian palynological assemblages are
103 characterized by the abundance of the gymnosperm pollen *Ricciisporites tuberculatus*
104 Lundblad —recently interpreted as produced by the peltasperm *Lepidopteris ottonis* (Goepf.)
105 Schimp. by Vajda et al., 2023—, followed by a fern spore spike across the Triassic–Jurassic
106 transition, and high abundances of *Classopollis* (Hirmerellaceae= Cheirolepidiaceae) in the
107 Lower Jurassic successions (Götz et al. 2009; Larsson 2009; van de Schootbrugge et al.
108 2009; Bonis et al. 2009, 2010; Pieńkowski et al. 2012; Vajda et al. 2013). A fern spike was
109 also identified within the Triassic–Jurassic sedimentary succession in North America,
110 followed by the dominance of *Classopollis meyeriana* (Klaus) de Jersey in the Lower
111 Jurassic successions (Olsen et al. 2002; Whiteside et al. 2007). In China, the Upper Triassic
112 *Dictyophyllidites*–*Kyrtomisoris*–*Ovalipollis*–*Ricciisporites* palynological assemblage is

113 replaced by the Early Jurassic *Dictyophyllidites–Classopollis–Cycadopites* assemblage (Lu
114 and Wang 1987; Wang et al. 2010). Rhaetian palynofloras from the Southern Hemisphere
115 (New Zealand) are dominated by lycophyte spores and corystosperm pollen, followed by a
116 high abundance of bryophyte spores in the uppermost Rhaetian, elevated osmundaceous
117 fern spore quantity in the Hettangian, and profuse *Classopollis* occurrence in the Sinemurian
118 (Akikuni et al. 2010; de Jersey and McKellar 2013). Similar stratigraphical abundance
119 patterns were observed in eastern Australian records across the Triassic–Jurassic transition,
120 with a large number of fern and bryophyte spores in the uppermost Rhaetian, and common
121 cheirolepid pollen occurrences in the Hettangian and Sinemurian (de Jersey and McKellar
122 2013).

123 It was suggested that very few plant families became extinct during the End-Triassic
124 crisis, as the seed ferns of the family Peltaspermeaceae (McElwain and Punyasena 2007);
125 and Umkomasiaceae (=Corystospermaceae) (Iglesias et al. 2011), being the overall balance
126 of plant groups essentially similar in Hettangian and Rhaetian floras (Cascales-Miñana et al.
127 2018). For example, cheirolepid conifers originated before the Triassic-Jurassic boundary,
128 and were affected by the extinction, but became markedly dominant in the Early Jurassic, up
129 to >90% in many assemblages (Whiteside et al. 2007; Lindström 2016). However, in lower
130 taxonomic levels extinctions are estimated to be higher, with up to 95% of species locally, in
131 Greenland and Sweden (McElwain et al. 1999). In other parts of central Europe (e.g.,
132 Austria, Poland), plant diversity increased across the Triassic-Jurassic boundary, probably
133 due to climatic warming resulting from redirected oceanic currents (Cascales-Miñana et al.
134 2018). The macroflora of Poland show no extinction event as the statistical analyses did not
135 signal any significant differences in plant composition between the Rhaetian and Hettangian
136 stages (Barbacka et al. 2017).

137 Other authors argued that the crisis amongst land plants at the end of the Triassic
138 was profound from an ecological perspective (McGhee et al. 2013; Lindström 2016). In the
139 United Kingdom and Austria, a replacement of Triassic broad-leaved forms with Jurassic
140 narrow-leaved forms was seen, which may have been favored by the warming associated

141 with increased atmospheric carbon dioxide levels (Belcher et al. 2010). In Greenland and
142 Sweden, the sequences through the Triassic/Jurassic boundary reveal clear evidence of
143 disruption to the vegetation, changing stomatal densities and carbon isotopes indicate a
144 marked increase in atmospheric CO₂, and abundant fusain suggests extensive wildfires
145 (McElwain et al. 1999). Combined mercury and teratology records demonstrate that
146 terrestrial plants in Northwestern Europe were subject to mutagenesis correlative with CAMP
147 volcanism. Thus, in addition to environmental stress from global warming and sulfuric acid
148 deposition due to emissions from CAMP (Steinthorsdottir et al. 2011, 2018), mercury-induced
149 phytotoxicity may have played a role in the demise of Late Triassic land plants (Lindström et
150 al. 2019).

151 The rapidity of the event has also been discussed. Wignall and Atkinson (2020)
152 showed that it can be resolved into two distinct, short-lived extinction pulses separated by a
153 several hundred thousand-year interlude phase; both consistent precisely with turnover and
154 extinction losses in the palynological record in Europe. The first one coincides with the
155 decline of conifer pollen and diversity reduction followed by the interlude phase, with
156 assemblages dominated by fern spores. The second extinction corresponds to another
157 abrupt, palynological change, which is the loss of several pollen taxa, and the brief
158 proliferation of *Classopollis* (Wignall and Atkinson 2020).

159 The biotic crisis related to the rapid global warming and oceanic oxygen deficiency
160 during the early Toarcian Oceanic Anoxic Event (T-OAE) has long been considered a major
161 event in the marine realm (Harris and Little 1999; Ullmann et al. 2014), and records of the
162 effects on terrestrial ecosystems have remained scant. However, palynological studies of
163 United Kingdom sequences have shown that during the T-OAE, terrestrial floras shifted from
164 a high-diversity mixture of conifers, seed ferns, wet-adapted ferns, and lycophytes to a low-
165 diversity assemblage dominated by cheirolepid conifers, cycads, and *Cerebropollenites*-
166 producers, which were able to survive in warm, drought-like conditions; and despite a rapid
167 recovery of vegetation after the T-OAE, the overall community composition remained notably
168 different than that previous to the event (Slater et al. 2019).

169 In Argentina it has been suggested that, towards the Triassic-Jurassic boundary, a
170 floristic turnover occurred, which led to the disappearance of Umkomasiales
171 (=Corystospermales) and Peltaspermales; an impoverishment in Cycadales, Ginkgoales, and
172 Gnetales, and to a greater abundance of ferns (e.g. Gleicheniaceae, Dipteridaceae,
173 Osmundaceae), bennettitaleans, and conifers (e.g. Araucariaceae, Cupressaceae,
174 Podocarpaceae) (cf. Artabe et al. 2007a; Iglesias et al. 2011). Concerning megafloras, about
175 50 taxa of ferns, bennettitaleans, and conifers originate in the Early Jurassic, different from
176 those recorded in the Triassic (cf. Artabe et al. 2007a; Iglesias et al. 2011).

177 However, in recent decades, new findings of plant megafossils and more accurate
178 radiometric datings cast doubts on previously proposed evolutionary models. Based on their
179 similarities in both macro- and micromorphology, a recent study considered *Alicurana* Herbst
180 & Gnaedinger (an Early Jurassic cycad genus from Argentina) as a junior synonym of
181 *Komlopteris* Barbacka *emend.* Slodownik et al., a genus assigned to a persistent lineage of
182 post-Triassic corystosperms (Slodownik et al., 2023). *Lepidopteris* and *Dicroidium* lineages,
183 dominant in Southern Hemisphere Triassic ecosystems, show a similar overall pattern of
184 origination (late Permian), diversification (late Early-Middle Triassic), and decline (Late
185 Triassic), with relict occurrences during the Early Jurassic (Elgorriaga et al., 2019; Sagasti et
186 al., 2019).

187 Few quantitative studies have been conducted on the Triassic-Jurassic megafloras of
188 Argentina. Among them are the similarity analysis by Damborenea et al. (1975) on
189 taphofloras from the Upper Triassic to the Lower Jurassic, and the contribution of Artabe et
190 al. (2007a), which assesses diversity changes, extinction rates, and origination rates during
191 the Middle Triassic to Early Jurassic interval.

192 The general purpose of this paper is to understand the terrestrial plant turnovers
193 during the Triassic-Jurassic Transition (End-Triassic Mass extinction event, the subsequent
194 recovery, and the early Toarcian Oceanic Anoxic Event) in Argentina, based on megafossil
195 records using a quantitative approach. Our results are compared with previous information
196 and correlated with environmental and climatic changes. Though this analysis covers the

197 Carnian-Berriasian time interval, it is mostly focused on the events occurring between the
198 Rhaetian and the Oxfordian. Given that the information available about floristic changes in
199 the Triassic-Jurassic transition is scarcer in the Southern Hemisphere than in the Northern
200 Hemisphere, this work aims to provide data for the comprehension of plant evolution
201 worldwide.

202

203 **MATERIALS AND METHODS**

204 *Database assembly*

205 In order to analyze the dynamics of the diversity of fossil plants in the Triassic-Jurassic
206 interval in Argentina, a database was developed on the records of plant megafossils for the
207 following lithostratigraphic units (formations), covering the time interval from the Carnian (Late
208 Triassic) to the Berriasian (Early Cretaceous): Los Rastros and Ischigualasto (Ischigualasto-
209 Villa Unión Basin); Carrizal (Marayes-El Carrizal Basin); Potrerillos, Cacheuta, and Río Blanco
210 (Cuyo Basin); Chihuido, Llantenes, Paso Flores, Arroyo Malo, Lapa, El Freno, Nestares,
211 Piedra del Águila, and Piedra Pintada (Neuquén Basin); Rancho de Lata (Frontal Cordillera);
212 Santo Domingo (also known as Quebrada de Santo Domingo Formation, Northern
213 Precordillera of La Rioja); Cerro Piche (North Patagonian Massif); unnamed formation at Cerro
214 Bayo, Lonco Trapial, Cañadón Asfalto, and Cañadón Calcáreo (Cañadón Asfalto Basin); Roca
215 Blanca, La Matilde, and Chon Aike (Deseado Massif); and Sprinhgill (Austral Basin) (Fig. 1).
216 We considered the ages of each formation as indicated by the geochronological dating when
217 available, or by biostratigraphic criteria when that information is controversial or not feasible.
218 Based on radioisotopic information, the Los Rastros Formation is considered Carnian (234.47
219 ± 0.44 Ma; Mancuso et al. 2020), while the Ischigualasto Formation is Carnian to Norian (231.4
220 to 225.9 Ma; Rogers et al. 1993; Martínez et al. 2011). The Carrizal Formation is correlated
221 with the Ischigualasto Formation through paleofloristic and palynological associations (Morel
222 et al. 2015, Césari et al. 2021; Colombi et al. 2021), assigning it to the Late Carnian. The
223 available radiometric information from the Potrerillos Formation includes U-Pb SHRIMP ages,
224 which date the central part of the unit as Carnian (230.3 ± 2.3 Ma) (Spalletti et al. 2008). The

225 biostratigraphic analysis of Bodnar et al. (2020) suggested that the Cacheuta Formation was
226 deposited during the Carnian-Norian interval, and the Río Blanco Formation, during the Norian.
227 Volkheimer and Papú (1993) analyzed the microfloristic assemblages recorded from the upper
228 levels of the Chihuido Formation, and they assigned the flora to the Norian stage. The
229 Llantenes Formation, which underlies the Chihuido Formation, was assigned to the Norian-
230 Rhaetian *Dictyophyllum tenuiserratum-Linguifolium arctum-Protocircoporoxydon marianaensis*
231 (DLM) Biozone (Spalletti et al. 1999; Morel et al. 2003), thus we referred this lithostratigraphic
232 unit to that interval. The Paso Flores Formation was also ascribed to the DLM Biozone
233 (Spalletti et al. 1999; Morel et al. 2003) and recently Gnaedinger and Zavattieri (2021)
234 proposed that its paleobotanical contents represents the youngest Triassic flora known in
235 Argentina, as a consequence, here we inferred a Rhaetian age for this formation. The Rancho
236 de Lata Formation was suggested to have been deposited since the Rhaetian to the
237 Sinemurian, but it can be differentiated into two sections according to its palynoflora: a
238 Rhaetian lower section and a Hettangian-Sinemurian upper section (Alvarez et al. 1994). The
239 Lapa Formation was considered as Rhaetian-Hettangian according to its fossiliferous content
240 and stratigraphic relationships (Spalletti et al. 1991; Gulisano and Gutiérrez Pleimling 1995).
241 The Santo Domingo Formation was regarded as uppermost Triassic–lowermost Jurassic in
242 age based on radiometric, paleomagnetic, and paleontological evidence (Caminos et al. 1995;
243 Coughlin 2000; Vizán et al. 2013), corresponding with doubts to the Rhaetian-Hettangian
244 interval. The El Freno Formation was suggested Hettangian-Sinemurian on the basis of its
245 paleofloristic assemblages and stratigraphical relationships (Spalletti et al. 2007; Lanés et al.
246 2013; Gnaedinger et al. 2015). The most recent sedimentological, environmental, and
247 paleobotanical data support a Sinemurian age for the Nestares Formation (Sagasti et al. 2019).
248 The calculated U-Pb SHRIMP age for the time of deposition of the tuffs of the Piedra del Águila
249 Formation is 191.7 ± 2.8 Ma (Sinemurian) (Spalletti et al. 2010). The Cerro Piche formation
250 was restricted to the Sinemurian- Pliensbachian interval by a combined U–Pb/Hf isotope study
251 (Falco et al 2021). For the Piedra Pintada Formation, we followed the stance of Damborenea
252 and Manceñido (1993) who assigned it to the Early Jurassic (Pliensbachian), latter followed by

253 Martínez and Olivera (2016). The ages of the Cañadón Asfalto Basin units were considered
254 according to U-Pb geochronology carried out by Cúneo et al. (2013), which indicates a
255 Sinemurian to Pliensbachian age for the Las Leoneras Formation and its equivalents (e.g. the
256 unnamed formation at Cerro Bayo), a Pliensbachian to Toarcian age for the Lonco Trapial
257 Formation, a Toarcian to Aalenian age for the Cañadón Asfalto Formation (extended to
258 Bajocian by Ruiz Gonzalez et al. 2024), and an Oxfordian to Kimmeridgian age for the
259 Cañadón Calcáreo Formation. The Roca Blanca Formation lacks radiometric ages, but an
260 early Pliensbachian to Toarcian age was suggested based on the fossil flora contents (Herbst,
261 1965; Gnaedinger and Herbst, 2009), which can be delimited by the $180,1 \pm 1,5$ Ma age of the
262 overlying Cerro Leon Formation (Guido et al. 2004). The Bahía Laura Group comprises the
263 interdigitated deposits of the Chon Aike and La Matilde formations. According to the ages
264 obtained by De Barrio (1993) and Pankhurst et al. (1993), 162 ± 11 Ma and 168 ± 1.9 Ma
265 respectively, the Bahía Laura Group can be dated as Bathonian to Oxfordian-Kimmeridgian.
266 Based on U-Pb radiometric dating, Lovecchio and Naipauer (2022) determined Tithonian to
267 Hauterivian maximum ages for the deposition of the Springhill Formation. This unit can be
268 differentiated into two sectors with fossil plant assemblages: Lago Argentino (Tithonian) and
269 western Austral Basin (Berriasian-Valanginian) (Del Fueyo et al. 2021; Tomas and Acuña
270 2022).

271 Some Upper Triassic and Jurassic formations were not included due to uncertainties
272 in the age of the fossil-bearing levels. The Laguna Colorada Formation (El Tranquilo Basin,
273 Santa Cruz Province), which has long been regarded as Norian in age mainly based on
274 occurrences of the *Dicroidium* paleoflora, was recently assigned to the Sinemurian due to
275 results of U-Pb geochronology from rocks intercalated with the vertebrate-bearing levels (Pol
276 et al. 2021). Until now, a new interpretation of the age of the plant-fossil-bearing levels of the
277 Laguna Colorada Formation has not been published according to the latest dating. The
278 Cañadón Largo (El Tranquilo Basin, Santa Cruz Province), was inferred to be Ladinian-early
279 Carnian in age because it underlies the Laguna Colorada Formation, previously assumed as
280 Norian, and there is an important hiatus between them (Jalfin and Herbst 1995). Thus, the new

281 age of the Laguna Colorada Formation will also change the inferred age of the Cañadón Largo
282 Formation. The Cepeda Formation (Cuyo Basin, San Juan Province) is supposed to belong to
283 the Late Triassic, probably Carnian, but with doubts (Bodnar et al. 2019), and for that reason
284 it was not considered for this analysis.

285 As many plant fossil assemblages were not precisely located stratigraphically along
286 each formation, the age of each one is not known in detail. Accordingly, we had to consider
287 that the age of each plant fossil assemblage was the same as that of the entire formation in
288 which they occur. This may cause some imprecision in the calculated each stage diversity,
289 since several formations span long intervals, extending over more than one stage.

290 In the database, we incorporated the number of specimens of each species recorded
291 in each formation. All fossilization types and organs were included, but when there is a whole-
292 plant concept, only the species which gives its name to the reconstruction was counted. Data
293 were taken from published data in the reference list available in Supplementary Material 1.

294 The database was prepared in various steps using MS Excel and Google Sheets
295 (Supplementary Material 2). First, data were compiled in MS Excel, resulting in: a 1/0
296 (presence/absence) matrix for species per formation; a table with the ages to which the
297 formations are assigned; and a species dataset with taxa by formation and stage, including
298 the taxonomic assignment from class to species and the age of the record. Then, an
299 occurrence dataset was gathered for each stage, with the collection number and the
300 references where samples were originally published or reported. Early and Late interval, and
301 the lower and upper boundaries, were restricted by each stage. In those cases where the
302 formation appeared in two stages, the record was multiplied. This was indicated by adding
303 a suffix to the collection number. The full table was compiled in one sheet and exported as a
304 .csv file.

305 *Repositories of specimens.*

306 The sample number indicated in the database corresponds to their repository number
307 according to the bibliography (see Supplementary Material 1). These repositories are:

308 -BA Pb (Colección Nacional de Paleobotánica del Museo Argentino del Ciencias Naturales
309 “Bernardino Rivadavia”, Buenos Aires, Argentina)
310 -BAFC-Pb (Colección de Paleobotánica de la Facultad de Ciencias Exactas y Naturales de
311 la Universidad de Buenos Aires, Buenos Aires, Argentina.
312 -CIRGEO-PB (Centro de Investigaciones en Recursos Geológicos, Buenos Aires, Argentina)
313 -CORD-PB (Museo de Paleontología de la Facultad De Ciencias Exactas, Físicas y
314 Naturales, Universidad Nacional de Córdoba, Córdoba, Argentina)
315 -CTES-PB (Colección Paleontológica de la Universidad Nacional del Nordeste, sección
316 Paleobotánica, Corrientes, Argentina)
317 -DNGyM (Dirección Nacional de Geología y Minería, Buenos Aires, Argentina)
318 -FMNH-P (Field Museum of Natural History, Paleobotany Collection, Chicago, USA)
319 -IANIGLA-PB (Colección de Paleobotánica del IANIGLA-CCT CONICET Mendoza,
320 Mendoza, Argentina);
321 -LIL-PB (Colección Paleobotánica de la Fundación Miguel Lillo, Tucumán, Argentina)
322 -LPPB (División Paleobotánica, Museo de La Plata, La Plata, Argentina)
323 -MAPBAR (Museo Paleontológico Bariloche, Bariloche, Argentina)
324 -MCF-PBPH (Colección Paleontológica del Museo Municipal “Carmen Funes”, Plaza
325 Huinca, Argentina)
326 -MCNAM-PB (Colección Paleobotánica del Museo de Ciencias Naturales y Antropológicas
327 “Juan Cornelio Moyano”; Mendoza, Argentina)
328 -MHNSR-PB (Museo de Historia Natural de San Rafael, San Rafael, Argentina)
329 -MLP (División Paleontología Invertebrados, Museo de La Plata, La Plata, Argentina)
330 -MNBP (Colección del Monumento Natural Bosques Petrificados, Jaramillo, Argentina)
331 -MPEF-PB (Museo Paleontológico “Egidio Feruglio”, Trelew, Argentina)
332 -MPM-PB (Museo Padre Jesús Molina, Río Gallegos, Argentina)
333 -Pb (Hunterian Museum -Palaeobotany-, Glasgow, United Kingdom)
334 -PBSJ (Colección Paleobotánica del Museo de Ciencias Naturales de San Juan, San
335 Juan, Argentina)

336 When the repository numbers were not provided, the specimens were mentioned
337 without them. Some of the fossil specimens are illustrated in Figures 2-4.

338 *Taxonomic assignments and systematic affinities*

339 The supra-generic taxonomic assignments were made according to those made in
340 the references and updated in the corresponding cases (see References available as
341 Supplementary Material 1). To organize the results concisely, some informal names of taxa
342 were used, such as: pteridophytes (comprising free-sporing vascular plants, as lycophytes,
343 sphenophytes, and ferns), pteridosperms or seed ferns (containing peltasperms,
344 corystosperms, petriellaleans, caytonialeans), cycadophytes (including cycadaleans,
345 bennetitaleans, and nilssonialeans), and coniferophytes (grouping ginkgoaleans,
346 czekanowskialeans, conifers, and gnetaleans).

347 We followed the most recently published and supported criteria for the affinity of each
348 taxon, but there are still discrepancies or ambiguities with the systematic alliance of some
349 genera that can lead to certain results to be carefully considered. One example is the genus
350 *Cladophlebis*, which was usually assigned to the family Osmundaceae (see Van
351 Konijnenburg-van Cittert 1996; Artabe et al. 2007b) but its morphology also resembles that of
352 Cyatheaceae or Schizaeaceae, among others (Skog and Dilcher 1994; Tidwell and Ash 1994).
353 *Kurtziana* is a genus related both to the cycads (Artabe et al. 1991) or the seed ferns
354 (Bomfleur et al. 2011), however, we followed the first criterion because it is the most widely
355 used in the Argentine Mesozoic literature. As both *Cladophlebis* and *Kurtziana* are taxa with
356 several records in the Triassic-Jurassic of Argentina, they have relevance when limiting the
357 temporal ranges of the lineages.

358 Another ambiguity relates to the Triassic megafossil record of Coniferales (=derived
359 conifers), which consists mainly of permineralized woods and leafy twigs, which are more
360 equivocal than reproductive structures (see Bodnar et al. 2023). Consequently, the temporal
361 ranges of Araucariaceae, Hirmerellaceae, and Cupressaceae could be narrowed if we adopt
362 a more restricted proposal.

363 *Calculated indices*

364 The table was then converted to csv and imported in RStudio, where it was analyzed
365 using DivDyn v.0.8.2 Package (Kocsis et al. 2019) (available as Supplementary Material 2).
366 Since this is an initial exploratory work, the equations provided in this package were used,
367 following the premise of Kocsis et al. (2019) that studies are repeatable and can be
368 continued by adding and improving information.

369 The value of species richness per stage was estimated based on the range-through
370 assumption and including singletons, which are taxa described from only one collection and
371 one stage. Species richness per stage of each plant lineage was estimated without the
372 range-through assumption.

373 Diversity was estimated in each case by applying the DivDyn v.0.8.2 Package of the
374 R software (Kocsis et al. 2019). The divDyn function calculates various metrics from
375 occurrence data sets in the form of time series. Through this function, we calculated the
376 following indexes of species and genera richness: Range-through diversity (divRT) (Newell,
377 1952), Sampled-in-bin diversity (divSIB) (Miller and Foote 1996), and Corrected sampled-in-
378 bin diversity (divCSIB) (Alroy et al. 2008). In this study, Range-through diversity considered
379 all species or genera in the interval, that is, if a genus appeared in one geological interval
380 and then disappeared later, one assumes that the species or genus existed for the entire
381 time between its first and last known fossil occurrences, regardless of gaps that there
382 may be on that geologic record. Sampled-in-bin diversity is the number of species or
383 genera actually sampled in the focal bin or interval. It is a metric that avoids the range-
384 interpolation bias, but it is more affected by changes in sampling intensity (Kocsis et al.
385 2022). Corrected sampled-in-bin diversity is a variant that corrects for residual error by
386 assessing the proportion of species or genera found immediately before and after a sampling
387 bin but not inside it (Alroy et al. 2008). Although this can be a convenient correction, it also
388 increases the estimation error. Nevertheless, this is the least biased estimator for diversity
389 (Kocsis et al. 2022). The diversity curves were compared with the number of occurrences in
390 each time bin. For species diversity, we regarded the number of specimens of each species
391 as occurrences, as provided by the references. For genera diversity, occurrences are the

392 number of genera per formation, coming from the same collection. In this work collections
393 correspond to the references, as it is explained below.

394 We applied the function “ranges” to plot ranges and occurrence distributions through
395 the Late Triassic-earliest Cretaceous interval and evaluated the First appearance data (FAD)
396 and Last appearance data (LAD).

397 Extinction and origination rates at generic level were calculated with time-averaging
398 by two methods: 1) per capita rates of Foote (1999) use the range-through assumption to
399 establish ranges for the taxa in the dataset; the rate value expresses what proportion of the
400 taxa decayed until the end of the interval or appeared at the beginning of the time-bin; and 2)
401 ‘gap-filler’ rates (Alroy 2014), which is a different estimator of the per capita rates but will
402 converge on them when sampling tends to completeness. We also estimated the proportions
403 of survivorship at genus and species level using the “survivors” function, which calculates
404 both forward and backward survivorship proportions from a given occurrence dataset. These
405 are tools to visualize changes in the composition of a group over time (Raup, 1978). The
406 curves show how a once coexisting set of taxa, called a cohort, loses its participants (forward
407 survivorship) as time progresses, or gains its elements as time is analyzed backward. Each
408 value corresponds to a cohort in bin (a) and one other bin (b). The value expresses what
409 proportion of the analyzed cohort (present together in bin a) is present in bin b.

410 Spearman Rank correlation coefficient was obtained to assess the correlations
411 between the richness values and the number of collections where they occur. The collections
412 refer to the fossil assemblages gathered by the same research team on a field trip (which
413 coincides with the published reference). To assess the sampling biases across the different
414 stages, sampling completeness was calculated using the three-timer sampling completeness
415 function (Alroy 2008) at generic and specific levels.

416

417 **RESULTS**

418 **Species richness**

419 Figure 5a illustrates the Range-through diversity (divRT), Sampled-in-bin diversity
420 (divSIB), and Corrected sampled-in-bin diversity (divCSIB) of species. During the Late
421 Triassic, a severe reduction of the species richness is seen from the Carnian to the Norian,
422 but in the Rhaetian, this decline is less pronounced (divRT and divCSIB) or even the values
423 slightly increase (divSIB). The results indicate that the species richness in the Hettangian is
424 reduced to half of the values recorded in the Rhaetian (divRT and divSIB), although
425 according to (divCSIB) this reduction is much more moderate. The divRT and divSIB curves
426 show that in the Sinemurian the number of species grows, but to less than observed in the
427 latest Triassic. This pattern is not observed in divCSIB curve, which drops from the
428 Hettangian to the Toarcian without clear peaks in the interval. If we consider the
429 occurrences, the Sinemurian has a significantly higher value than the Hettangian, which
430 could be influencing the observed increase in divRT and divSIB curves. However, this effect
431 is not reflected in the divCSIB curve, as the latter is the least affected by sampling intensity.
432 A decrease in species richness is documented from the Pliensbachian to the Bajocian in all
433 the curves, and then an increase is recorded in the Bathonian and another in the Oxfordian.
434 In the Tithonian, the species richness is strikingly reduced. During the Jurassic, species
435 richness (per stage) does not reach the Late Triassic values.

436 When the diversity curve is compared with the occurrences (Fig. 5b), the stages in
437 which the species richness is greater agree with those with larger number of records and
438 collections. The Pliensbachian has the highest number of occurrences; however, the
439 diversity curve slightly drops in this stage. Spearman's rank correlation coefficient is
440 0.6433087, which indicates a moderate to high positive correlation between species richness
441 and occurrences. The p-value equals 0.009672 (less than 0.05) which means that the
442 correlation is statistically significant. The sampling completeness shows scores higher than
443 0.9 in all the Jurassic stages except for the Hettangian, in which the lowest sampling
444 completeness is recorded. That the reason why the diversity decreases in this stage are less
445 marked in CSIB metrics (Tab. 1; Fig. 5).

446 Regarding the species richness of each plant lineage (Tab. 2), the corystosperms
447 (Umkomasiaceae) are the most diverse group in the Carnian, Norian, and Rhaetian, with 31,
448 16, and 13 species, respectively (see also Fig. 2). This family passes just as relicts to the
449 Early Jurassic with four species (Tab. 2; Fig. 3). Other gymnosperm orders important during
450 the Late Triassic are the Peltaspermales (Peltaspermaceae and Mattatiellaceae),
451 Petriellales, Cycadales, Ginkgoales, and Gnetales. All these orders occur in the Jurassic but
452 only with very few species. During the Hettangian, the species richness of each recorded
453 order or family does not exceed four species. The bennettitaleans, with 14 species, and the
454 dipteridacean and osmundacean ferns, with 11, are the lineages with greatest specific
455 richness in the Sinemurian (Tab. 2; Fig.3). The dipteridacean species richness decreases
456 from the Toarcian onwards. On the contrary, osmundacean ferns diversify again in the
457 Bathonian. A diminution of the bennettitalean richness is observed from the Pliensbachian
458 and they became the most diverse plant lineage during the Bathonian-Kimmeridgian. The
459 modern conifer lineages show a progressive diversification from the Sinemurian, and in the
460 Oxfordian and Kimmeridgian, the Araucariaceae and Podocarpaceae are the most diverse
461 conifer families (Tab. 2; Fig. 4). In the studied interval, the Marchantiopsida, Bryopsida, and
462 the lycopodiacean lycophytes are recorded only in the Carnian, with one species each.
463 Pleuromeiacean and isoetacean lycophytes are present in the Carnian-Norian and
464 Bathonian-Kimmeridgian, respectively, also with one species each. Within sphenophytes, the
465 Equisetaceae family shows species richness values quite stable from the Late Triassic to the
466 Late Jurassic, with one to five species, while Neocalamitaceae (= Apocalamitaceae) reach
467 four species in the Carnian, three in the Sinemurian, and disappears afterward.

468 **Genera richness**

469 The calculated diversity curves (divRT, divSIB, and divCSIB; Fig. 6a) show a
470 decrease between the Rhaetian and Hettangian, and a slight rise in the Sinemurian (divRT
471 and divSIB) or in the Toarcian in the corrected-sample diversity (divCSIB). As in species
472 richness, the Sinemurian has the greatest number of genera occurrences, which could
473 influence the observed increase in divRT and divSIB curves. During the Jurassic, the genera

474 richness does not reach the same levels as in the Rhaetian. Diversity increases in the
475 Toarcian (divRT, divSIB, and divCSIB) and the Bathonian (divRT, and divCSIB) or Callovian
476 (divSIB). A general diminution in genera richness is recorded after the Oxfordian (divRT,
477 divSIB, and divCSIB). The stages with the greatest plant diversity in the Jurassic are different
478 depending on the estimator: the Toarcian for the divRT, the Toarcian, Bathonian, Callovian,
479 Oxfordian, and Kimmeridgian for divSIB; and the Oxfordian and Kimmeridgian for divCSIB
480 diversity.

481 When the diversity curve is compared with the occurrences (Fig. 6b), the stages in
482 which the genera richness is greater coincide with those with the largest number of records.
483 Spearman's rank correlation coefficient gives a value of 0.4555421, which indicates a
484 moderate positive correlation between the genera richness and occurrences. The p-value
485 equals 0.08793 (greater than 0.05) which means that the correlation is not statistically
486 significant. In summary, although there appears to be a moderate positive correlation
487 between the two variables, it is not significant at the common threshold of 5% (p-value
488 <0.05).

489 Comparing the diversity curves of different plant groups (Fig. 7-8), seed plants (Fig.
490 7) undergo a marked reduction of the genera richness between Rhaetian and Hettangian,
491 and a general decrease from the Upper Triassic to the Upper Jurassic, while these trends
492 are less significant for free-sporing vascular plants (pteridophytes). This last plant group goes
493 through generic-level diversification in the Sinemurian and Pliensbachian, impoverishment in
494 the Aalenian-Bajocian, and a recovery from the Bathonian (Fig. 7, 8a). Within seed plants,
495 the pteridosperms (Umkomasiales, Peltaspermales, Petriellales, Caytoniales, and incertae
496 sedis pteridosperms) show a gradual reduction of the diversity from the Upper Triassic to the
497 Upper Jurassic, with a mild recuperation during the Sinemurian and Toarcian (Fig. 8b). The
498 divRT curve of cycadophytes (Cycadales, Nilssoniales, and Bennettitales) exhibits minor
499 changes from the Upper Triassic to the Upper Jurassic, but with the major values in the
500 Sinemurian and Bathonian to Kimmeridgian (Fig. 8c). The divSIB curve of this group also
501 shows the peaks in the Sinemurian and Bathonian- Kimmeridgian, being, in this case, more

502 noticeable. In both curves, a reduction of the cycadophyte genera richness is not recorded
503 between the Rhaetian and Hettangian. The coniferophytes (Voltziales, Coniferales,
504 Ginkgoales, Czekanowskiales, and Gnetales) generic-level diversity diminishes in the
505 Hettangian, progressively increases up to the Toarcian, and experiments a peak in the
506 Bathonian-Kimmeridgian (Fig. 8d). The sampling completeness shows the highest values
507 between the Sinemurian to the Kimmeridgian, and the lowest scores in the Hettangian and
508 Tithonian (Tab. 1).

509 **Occurrences**

510 The number of occurrences of free-sporing vascular plants is much greater in the
511 Pliensbachian than in other analyzed stages (Fig. 9a), with Dipteridaceae and Osmundaceae
512 being the most representative families (Fig. 9a, 9b). The occurrences for Neocalamitaceae
513 abruptly drop in the Rhaetian and Hettangian and persist up to the Sinemurian as relicts. The
514 families Dipteridaceae, Osmundaceae, and Equisetaceae are the most frequent in the Lower
515 Jurassic (Fig. 4, 9a, 9b), whereas in the Middle Jurassic, the most common are the
516 Osmundaceae. The number of pteridophyte records abruptly declines in the Upper Jurassic.
517 Considering the proportion of occurrences (Fig. 9b), Osmundaceae is a dominant family
518 within ferns during almost all the studied interval, Dipteridaceae is important in the Rhaetian
519 and Lower Jurassic, the Equisetaceae increases its representation throughout the Jurassic,
520 and Gleicheniaceae is more relevant in the uppermost Lower and lowermost Middle Jurassic.
521 As the few fern fossil records during the Tithonian correspond to specimens assigned to
522 *Coniopteris*, which is a putative Dicksoniaceae genus, the importance of this family in the
523 Upper Jurassic is overrated. Lycophytes are very infrequent in the studied bin.

524 The occurrences of pteridosperms severely decline from the Carnian up to the
525 Hettangian, increase from the Sinemurian up to the Toarcian, and drop again from the
526 Aalenian onwards (Fig. 9c). There is a reversal in the importance of the Umkomasiaceae in
527 the Upper Triassic, to the Caytoniaceae, Peltaspermeaceae, and Matattiellaceae in the
528 Jurassic (Fig. 9d).

529 The number of cycadophyte occurrences is greatest in the Lower Jurassic (Fig. 9e).
530 The proportion of records displays that the Cycadales are dominant in the Upper Triassic and
531 from the Lower Jurassic upwards, the Bennettitales represents almost all the cycadophyte
532 occurrences (Fig. 9f).

533 Within coniferophytes, the voltzian conifers, gnetaleans, and ginkgoaleans are
534 common and dominant during the Carnian, but their importance considerably decreases from
535 the Hettangian onwards (Fig. 9g, 9h). The proportion of occurrences of modern conifers
536 (Coniferales) markedly increases from the Lower Jurassic onwards. Cupressaceae has
537 significance in the Lower-Middle Jurassic, Araucariaceae is important during the Middle and
538 Upper Jurassic, and Hirmerellaceae (=Cheirolepidiaceae) acquires significance in the Upper
539 Jurassic (Fig. 9g, 9h).

540 **Ranges and FAD-LAD**

541 At species level, a floral replacement is seen in the Early Jurassic (Fig. 10), with the
542 appearance of several species that last for a short time and are successively replaced. A few
543 taxa persist from the Late Triassic or Early Jurassic, but there is no long-term stability in the
544 composition of plant communities.

545 Regarding the genera ranges (Supplementary Material 3), 43 genera disappear
546 through all the Late Triassic in Argentina. Although the genera *Hepaticites*, *Muscites*,
547 *Protophylocladoxylon*, and *Androstrobus*, are not present in the Jurassic or the first stage of
548 the Cretaceous, they were described in younger Cretaceous or Cenozoic sequences from
549 Argentina (Archangelsky 1965; Archangelsky and Villar de Seoane 2004; Puebla et al. 2012;
550 Pujana 2022). Ten genera are singletons, present only in one stage. It is important to point
551 out that *Lycopodites*, *Gontriglossa*, *Scytophyllum*, *Rissikianthus*, and *Protocircoporoxylon* are
552 not considered singletons since these taxa were described in Early-Middle Triassic
553 formations from Argentina (Coturel et al., 2016; Bodnar et al., 2019; Drovandi et al., 2020),
554 despite that they are recorded in one stage of the studied interval. Excluding singletons and
555 those that occur in the Cretaceous or Cenozoic, 29 genera have their last appearance in the
556 Rhaetian.

557 Thirty-two genera appear for the first time in the Early Jurassic in Argentina according
558 to this study, from these, six genera are recorded in the Middle Triassic or in Triassic units
559 not included in this work because their age is under discussion: *Rienitsia*, *Elatocladus*,
560 *Desmiophyllum*, *Pelourdea*, *Pterophyllum*, and *Todites* (Troncoso et al. 2000; Zamuner et al.
561 2001). On the other hand, *Komlopteris* is a singleton genus as is only present in the
562 Sinemurian. From the 25 remaining taxa, seven appear for the first time in the Hettangian,
563 six in the Sinemurian, seven in the Pliensbachian, and four in the Toarcian.

564 Finally, forty-six genera present in the Argentine Triassic pass to the Jurassic; from
565 these, twenty-one disappear in the Early Jurassic.

566 Two lineages disappear in the Late Triassic: the pleuromeiacean lycophytes, and the
567 gymnosperm family Hamshawviaceae (Tab. 3). The families that first occurred in the
568 Jurassic are the Gleicheniaceae, Caytoniaceae, and Williamsoniaceae in the Early Jurassic,
569 the Isoetaceae in the Middle Jurassic, and the Matoniaceae in the Late Jurassic (Tab. 3).

570 **Survivorship curves, and origination and extinction rates**

571 In the 'forward' survivorship curves of plant genera, two different groups of cohorts
572 can be recognized in the studied bin (Fig. 11a), one extending from the Late Triassic up to
573 the Jurassic and the other spanning from the Jurassic up to the Early Cretaceous. The
574 slopes of these curves for the Late Triassic cohorts are similar to those of the Early Jurassic
575 and Middle Jurassic cohorts. The more pronounced slopes of the curves are observed
576 between the Rhaetian and Hettangian; the Sinemurian and Pliensbachian; and the
577 Kimmeridgian and Tithonian. The 'backward' survivorships show a more marked cohort
578 emergence between Rhaetian and Hettangian, and between Bajocian and Bathonian (Fig.
579 11b).

580 The slopes of survivorship curves of species (Fig. 11c) tend to be steeper than those
581 of genus curves, which is inherent to the method. All the species curves show abrupt
582 inflections in the intervals in which the genera curves exhibit sharper slopes. It is noticeable a
583 cohort with very marked slopes, extending from the Late Triassic and only to the
584 Pliensbachian.

585 The extinction rates at the genus level show the greatest values in the Rhaetian, the
586 Toarcian, and between the Oxfordian and Kimmeridgian (Fig. 12a). The origination rates at
587 genera-level show the greatest values in the Rhaetian, and Bathonian, and a less
588 pronounced peak in the Pliensbachian, (Fig. 12b). The peaks of extinction and origination in
589 the Rhaetian show a floral replacement. The peaks of origination of the passages
590 Pliensbachian-Toarcian and Bathonian-Calloviaian coincide with the lower values of extinction.
591 In comparison with the diversity curves, the first is correlated with a slight decrease in
592 diversity and the second with a clearer increase. The extinction and origination rates exhibit
593 no values after the Kimmeridgian because of the absence of temporal definition of the
594 paleofloras (see Discussion), by the same reason, there are no changes in the diversity
595 between the Oxfordian and Kimmeridgian.

596

597

598 **DISCUSSION**

599 **Sampling and taphonomical biases**

600 This study comprised all fossil types without discrimination according to organs. While
601 this inclusion approach may be subject to debate, it is essential to note that the fossilization
602 type varies from one formation to another, while in some only permineralized logs are
603 preserved, others only carry impressions-compressions of leaves. Being the plants modular
604 organisms, each individual can produce a great number of leaves, branches, etc., that are
605 preserved in the fossil record as disarticulated plant remains. For example, in a fossiliferous
606 locality, a large number of fossil leaves belonging to deciduous trees may have been
607 preserved by the nature of the original organ but not as a consequence of the number of
608 individuals which lived in the plant community (see Serrano-Brañas and Reyes-Luna, 2014).
609 For this reason, if we restrict ourselves to one type of fossil or organ, we would not have
610 enough information on some of the analyzed stages. From the Carnian to the Toarcian, the
611 predominance of leafy branches and leaves (mostly preserved as impressions and
612 compressions) over other plant organs is very clear, representing about 67-85%

613 (Supplementary Material 4). Reproductive structures (preserved as impressions-
614 compressions or permineralizations) are more abundant from the Pliensbachian to the
615 Kimmeridgian, however, from the Aalenian onwards, they represent a significantly larger
616 proportion of the total occurrences. On the other hand, the permineralized woods are more
617 abundant in the Bathonian-Kimmeridgian interval, where they account for a quarter of the
618 occurrences. In general, it can be observed that in stages with a lower proportion of fossilized
619 leaves, the total number of occurrences also decreases, suggesting that leaves contribute
620 more records than other plant organs. This is due to the intrinsic nature of plants, as they
621 produce a large number of leaves throughout their lifetime, while they form only one stem. In
622 comparison with the diversity curves, variations in the proportion of organ types and
623 fossilization modes could be attributed to both taphonomic and taxonomic causes. For
624 instance, fossiliferous sites with permineralized forests become very common from the
625 Middle Jurassic onwards in Argentina, and in these locations, due to paleoenvironmental
626 factors, impressions and compressions are uncommon. The greater abundance of fossil
627 woods coincides with an increase in coniferophyte diversity (Fig. 8d), which are tree woody
628 plants with large trunks.

629 Sampling biases in the paleobotanical record have been scarcely evaluated (Cleal et
630 al. 2021; Pardoe et al. 2021). Our results show a high positive correlation between species
631 diversity and collections but this correlation is lower at genera level. This means that
632 increased sampling effort in the same locations may result in more species or morphological
633 types. In paleobotany, the morphological species concept is applied and variability within a
634 species is only occasionally considered, and thus morphological combinations that minimally
635 differ from already known ones are classified as new species.

636 The methodological scope varies in each research included in this analysis. Some
637 references (here considered as collections) are studies focused on one to three species with
638 a very large number of specimens of each taxon (see for example Escapa et al. 2008, 2015;
639 Elgorriaga et al. 2015, 2019; Choo et al. 2016; Contreras et al. 2019) while other papers
640 aims are floristic analyses, where there are comparatively fewer specimens for many more

641 taxa (see for example, Petriella and Arrondo 1984; Spalletti et al. 1991, 2010; Morel et al.
642 1999, 2013). This could affect sampling completeness.

643 Another issue that may affect the index is the bias produced by the wide time ranges
644 of the Jurassic formations. While the Triassic formations are better constrained in age, and
645 even an intra-formational analysis could be performed, the Jurassic formations with their
646 respective floras artificially span several stages.

647 Another analyzed bias in the estimation of past diversity has been the type of
648 environments in which plant fossils are preserved. Plant preservation potential in surface
649 environments increases dramatically in the presence of surface water bodies, high water
650 tables, or with rapid burial to below the vadose zone, which are circumstances more
651 frequently present in humid climatic environments than in arid or semi-arid environments and
652 where sedimentation rates are high and, as a consequence, wetland floras preserved in
653 basinal environments during humid climatic intervals dominate much of the plant fossil record
654 (Channing and Edwards 2013). On the other hand, specific richness tends to be greater in
655 the wetter biomes (McNeely 2003), hence a larger number of plant remains in the humid
656 intervals would only overstate a diversity increase which would have actually occurred.

657 In our results, there is a correlation between the diversity and the number of
658 formations per stage, but in some stages, this relation is not so evident. For instance, there
659 are only two units (i.e. Chon Aike and La Matilde Formations) from the Bathonian-Callovian,
660 which are among the stages with greatest diversity; nevertheless, La Matilde Formation is
661 one of those with the largest number of outcrop sites of the Argentinean Jurassic and its
662 paleoflora was studied unceasingly since 1924 for several authors (resulting in 31
663 collections).

664 Despite the estimated plant diversity could appear as essentially the result of sampling
665 bias, Capel et al. (2022) note that the number of localities may co-vary with abundance and
666 diversity, and so diversity may not be just a response to heterogeneous sampling or lack of
667 fossiliferous localities. For example, the records of plant macrofossils after an extinction event
668 can be scarce as plants were not as widespread as they were later in the recovery interval. In

669 this study, the RT and SIB diversity Hettangian decrease and the increase in the Pliensbachian
670 (and their corresponding drop and climb of occurrences) can be an example. In fact, the low
671 number of localities probably reflects actual low plant abundance and diversity of the
672 vegetation, implying a redundancy effect. The concomitant increase in diversity and number
673 of localities probably also reflects the progressive colonization of the plants after the extinction
674 event (Capel et al. 2022).

675 **Comparison with previous studies**

676 A major floristic change in Argentina towards the Triassic-Jurassic boundary was
677 suggested, leading to the disappearance of the Umkomasiales and Peltaspermales, the
678 impoverishment of the Ginkgoales, and an increase in the diversity of ferns, Bennettiales
679 and conifers (Artabe et al. 2007a; Iglesias et al. 2011). According to the most recent findings,
680 included in this contribution, Umkomasiales and Peltaspermales (Peltaspermaceae and
681 Mattatielaceae) reached the Early and Late Jurassic, respectively. During the Early Jurassic,
682 Ginkgoales declined, while derived conifers, Bennettiales, and ferns of the families
683 Osmundaceae and Dipteridaceae diversified.

684 In their reviews of Mesozoic floras, Artabe et al. (2007a) and Iglesias et al. (2011)
685 stated that the Argentine Triassic flora comprises 86 genera and 238 species, from which
686 only 28 species survive in the Jurassic. Although it is not explicit in their contribution, Artabe
687 et al. (2007a) elaborated diversity curves of species richness under the range-through
688 assumption, based on an absence/presence matrix of species using Past software (Artabe,
689 pers. comm.). Our study shows that from 166 species present in the Late Triassic, only 26
690 Triassic species survive into the Jurassic, but at the genera level the survival is greater (from
691 87 genera present in the Late Triassic, 48 pass to the Jurassic). Artabe et al. (2007a)
692 recorded 131 species in the Carnian (BNP Biozone) and 54 in the Norian (OL Biozone),
693 while in our analysis we recorded 122 species in the Carnian, and 77 species in the Norian.
694 These discrepancies may be due to the new interpretation about the age of some units and
695 the new paleobotanical data published since those authors carried out their investigation in
696 2007 (see Introduction and Material and Methods). Artabe et al. (2007a) and Iglesias et al.

697 (2011) also asserted that 27 genera appeared for the first time in the Early Jurassic and that
698 Jurassic floras are characterized by a low diversity. Our results show that in the Early
699 Jurassic 24 genera appeared for the first time and that, although the Jurassic diversity is
700 lower than the Carnian one, the species and genera richness values of some Jurassic stages
701 are comparable to those of the Norian and Rhaetian.

702 Quattrochio et al. (2007) suggested that the megafloras of the Middle and Late
703 Jurassic of Argentina show an impoverishment in the number of species compared with the
704 Early Jurassic. Our estimations exhibit high diversity in Middle Jurassic megafloras and a
705 decrease in diversity afterward.

706 Regarding the controversy about the abrupt or gradual extinction of the terrestrial
707 vegetation across the Triassic–Jurassic transition, at the lineage level only the
708 Pleuromeiaceae became extinct, but there is an important change in the dominance of the
709 groups. Despite the most abundant groups in the Late Triassic (Neocalamitaceae,
710 Dipteridaceae, Osmundaceae, Umkomasiaceae, Cycadales, Ginkgoales, Voltziaceae, and
711 Gnetales, Fig. 2) pass through the Jurassic, most of them pass as relicts (Neocalamitaceae,
712 Umkomasiaceae, Ginkgoales, Voltziaceae, and Gnetales) or with less dominance
713 (Cycadales). Only Dipteridaceae and Osmundaceae diversified during the Jurassic.
714 According to our database, several genera considered diagnostic of the Argentine Triassic
715 survive as relict forms to the Early Jurassic: *Neocalamites* (Neocalamitaceae), *Rietnisia*
716 (Asterothecaceae), *Dicroidium* (see Fig. 3e), and *Rhexoxylon* (Umkomasiaceae), *Rochipteris*
717 (Petriellaceae), *Dejerseya* (Matatiellaceae), *Heidiphyllum* (Voltziaceae); *Kurtziana*
718 (Cycadales) or even to the Late Jurassic *Lepidopteris* (Peltaspermaceae). Most lineages and
719 genera that go through the Triassic–Jurassic boundary become extinct in the Early or Middle
720 Jurassic. The extinction rates indicate a peak at the Rhaetian; however, the origination rates
721 also have high values indicating a turnover. The slopes of the survivor curves between
722 Rhaetian and Hettangian are pronounced. Taking our results with caution due to the
723 sampling biases, we can see that there is an important floristic change in the Triassic–
724 Jurassic transition, but it is not so abrupt, and several taxa became extinct gradually during

725 the Late Triassic, between 230 Ma and 202 Ma, before the T-J boundary and afterward in the
726 Jurassic.

727 After the end-Triassic, the terrestrial ecosystems appear to have recovered by late
728 Hettangian time in several areas of the world (Austria, UK, Denmark, Greenland, Arctic
729 Canada, Australia, and New Zealand), with the return of common to abundant woody plants
730 (Lindström 2016). In Argentina, the richness values and occurrences are low for all plant
731 groups in the Hettangian as seen in our analysis. However, it is important to recognize that
732 sampling at this stage remains the least comprehensive. Recovery can be inferred in the
733 Sinemurian since an important increase in diversity of pteridophytes, seed ferns, and
734 cycadophytes is recorded. Ferns, which are considered colonizer taxa commonly found in
735 disturbed environments (Zhou et al. 2021), are more abundant and diverse (at species and
736 family levels) in the Sinemurian of Argentina, which could be an indication that they
737 accompanied the ecosystem recovery. The coniferophytes show an important diversification
738 from the Pliensbachian.

739 Regarding the early Toarcian Oceanic Anoxic Event, we observed an increase of
740 overall diversity at species and genus level in the Toarcian, in particular of the seed plants,
741 with a major contribution of the derived conifers. However, given the nature of our data, the
742 information from the Toarcian is insufficient, as the materials may come from layers both
743 below and above the event. A peak of the extinction rate curves is observed in the Toarcian
744 and the diversity falls in the Aalenian, and these changes could be linked to the Toarcian
745 crisis.

746 **Paleoclimatic discussion**

747 The Rhaetian was characterized by a warm wet climate with a definite seasonality
748 (Parrish 1993). A peak of species and genera richness, both in pteridophytes and
749 gymnosperms, is observed in Argentina.

750 The Early Jurassic climate was characterized by global warming, more intense than
751 the present; average summer temperatures exceeded 35 °C in low-latitude regions of
752 western Pangea, where eolian sandstones testify to the presence of vast deserts (Chandler

753 et al. 1992). High rainfall rates were associated primarily with monsoons that originated over
754 the warm Tethys Ocean, characterizing an overall paleoclimate for the Jurassic (Chandler et
755 al. 1992). The climatic conditions at the beginning of the early Jurassic (Hettangian) were
756 similar to those of the Late Triassic, and the Pliensbachian to Toarcian was a time of global
757 warming, probably driven by the Karoo magmatism (Holz 2015). For Argentina, warm (-
758 temperate) and humid conditions during the Sinemurian and increasing warmth and aridity
759 from the late Pliensbachian to the Toarcian were suggested (Volkheimer et al. 2008a). Our
760 results indicate a diversity reduction in the Hettangian, and during the Sinemurian a
761 diversification of pteridophytes (especially Osmundaceae and Dipteridaceae) and
762 cycadophytes (bennettitaleans) in the second case (Fig. 3). The family Osmundaceae and
763 the genus *Otozamites* are indicative of wet and warm climates (Van Konijnenburg-Van Cittert
764 2002; Wang et al. 2008). The Pliensbachian appears as an interval with an overall diversity
765 decline and a decline of seed ferns, but a slight increase in fern and conifer diversity.

766 The Early Toarcian global warming trend is marked by an oceanic anoxic event (Holz
767 2015). A change from wet (fern-dominated) to warmer and seasonally dry landscapes during
768 the Toarcian was interpreted from palynological data from Denmark (Wade-Murphy et al. 2006)
769 and plant fossils have also been interpreted to reflect short-term warming in the Toarcian of
770 Siberia (Vakhrameev 1991). The Toarcian is one of the moments of the Jurassic with highest
771 overall diversity at the genus level for Argentina as can be seen from our analysis. A
772 diversification in seed plants (mainly the coniferophytes), but a reduction in the genera richness
773 of ferns, agrees with the floristic change in other parts of the world and drier conditions. At the
774 Toarcian, there is an extinction rate peak in Argentine megafloras. Although this peak does not
775 coincide with the early Toarcian crisis, the its age needs to be adjusted to conclude in this
776 regard, considering that there is no precise dating of each plant fossil assemblage.

777 For the Early Bajocian, Hesselbo et al. (2003) characterized strong seasonally arid and
778 weak seasonally arid climates. Our data show a decrease in plant diversity and occurrences
779 in the Aalenian and Bajocian.

780 From Late Callovian to Early Oxfordian (i.e., the transition from Mid-to Late Jurassic),
781 migration of marine fauna and isotopic thermometry pointed out drastic cooling during the early
782 Late Callovian, suggesting continental ice formation at this time (Holz 2015). Dromart et al.
783 (2003) designated this episode the “Callovian Ice Age”, hence, the Middle Jurassic seems to
784 display one of the coldest climates of the entire Mesozoic. For Argentina, Volkheimer et al.
785 (2008b) suggested warm and relatively humid climatic conditions during the Middle Jurassic
786 due to the mega- and microfloras with high percentages of the thermophilic Hirmerellaceae
787 (=Cheirolepidiaceae), associated with Araucariaceae, which need relatively humid conditions.
788 Consistent with the present analysis, in the Bathonian-Callovian, the overall diversity is high in
789 every plant group except for the seed ferns, and as in the previous studies, with a major
790 representation of Osmundaceae, Bennettiales, Araucariaceae and a minor importance of the
791 Hirmerellaceae (Fig. 4). The Araucariaceae maintained a preference for subtropical or
792 mesothermal conditions (Kershaw and Wagstaff, 2001); therefore, it cannot be ruled out that
793 the climate was temperate in this interval.

794 More recent papers corroborate the Oxfordian-Kimmeridgian warming and the more
795 humid paleoclimatic conditions towards the Tithonian (e.g. Wierzbowski et al., 2013). In the
796 Oxfordian-Kimmeridgian interval, a mild increase in species diversity is seen, but at genus
797 level this is imperceptible. In the Oxfordian, the extinction rates reach the highest value, and
798 the origination rates, the lowest value. This could be exaggerated by the lack of a precise age
799 definition of the plant-fossil strata from the formations included in that bin (i.e. Chon Aike and
800 La Matilde). This lack of precision results in origination rates of zero and extinction rates greater
801 than one, making it impossible to graph in Figure 12. A decrease in floral diversity towards the
802 Tithonian is observed, but as the number of total occurrences is also very few for that stage in
803 Argentina (25 occurrences, one formation with two collections), these results should be taken
804 with caution.

805 .

806 **CONCLUSIONS**

807 We carried out a detailed characterization of the megafloras of the Triassic-Jurassic
808 transition, based on recent advances in the knowledge of the Upper Triassic and Jurassic of
809 Argentina. This analysis offers guidance for future collection and analytical efforts to address
810 knowledge gaps in plant fossil assemblages from intervals and localities with scarce or
811 imprecise records. The dataset generated in this work can be improved by updating,
812 modifying, or adding new information in the future and in this way the results obtained will be
813 more robust.

814 The obtained curves of diversity and origination and extinction rates, display
815 fluctuations along the Triassic-Jurassic transition, but there is no absolute certainty that these
816 patterns precisely align with globally dated events. In Argentina megafloras, the End Triassic
817 and early Toarcian mass extinctions are reflected more as gradual turnovers than as abrupt
818 crises. Several lineages that were dominant during the Triassic survive up to Sinemurian, but
819 taxa that will become more important in Jurassic and Early Cretaceous floras emerge after the
820 Triassic–Jurassic boundary.

821 The Sinemurian was an interval in which ferns and bennettitaleans diversified, which
822 can be related to the previous interpretation of a wet climate during the Sinemurian for
823 Argentina. In the Toarcian, the increase of conifer diversity and a decrease in fern richness,
824 could be correlated with the reduction in humidity proposed for this stage. The greater
825 importance of the Osmundaceae, Araucariaceae, and Bennettitales, from the Bathonian
826 onwards coincides with the warmer and wetter climates suggested for the Bathonian-
827 Kimmeridgian interval.

828

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834

835 **References**

- 836 Akikuni, K., Hori, R., Vajda, V., Grant-Mackie, J.A. and Ikehara, M. 2010. Stratigraphy of
837 Triassic–Jurassic boundary sequences from the Kawhia coast and Awakino gorge, Murihiku
838 Terrane, New Zealand. *Stratigraphy* 7(1): 7-24.
- 839 Alroy, J. 2008. Dynamics of origination and extinction in the marine fossil record. *Proceedings*
840 *of the National Academy of Science* 105: 11536–11542.
- 841 Alroy, J. 2014. Accurate and precise estimates of origination and extinction rates. *Paleobiology*
842 40(3): 374-397.
- 843 Alroy, J., Aberhan, M., Bottjer, D.J., Foote, M., Fürsich, F.T., Harries, P.J., Hendy, A.J.,
844 Holland, S.M., Ivany, L.C., Kiessling, W., Kosnik, M.A., Marshall, C.R., McGowan, A.J.,
845 Miller, A.I., Olszewski, T.D., Patzkowsky, M.E., Peters, S.E., Villier, L., Wagner, P.J.,
846 Bonuso, N., Borkow, P.S., Brenneis, B., Clapham, M.E., Fall, L.M., Ferguson, C.A., Hanson,
847 V.L., Krug, A.Z., Layou, K.M., Leckey, E.H., Nürnberg, S., Powers, C.M., Sessa, J.A.,
848 Simpson, C., Tomasovych, A. and Visaggi, C.C. 2008. Phanerozoic trends in the global
849 diversity of marine invertebrates. *Science* 321(5885): 97-100.
- 850 Álvarez, P.P., Benoit, S.V. and Ottone, E.G. 1994. Las formaciones Rancho de Lata, Los
851 Patillos y otras unidades mesozoicas de la Cordillera Principal de San Juan. *Revista de la*
852 *Asociación Geológica Argentina* 49(1-2): 123-142.
- 853 Archangelsky, S. 1965. Fossil Ginkgoales from the Ticó Flora Santa Cruz Province, Argentina.
854 *Bulletin of the British Museum (Natural History) Geology* 10(5): 119-138.
- 855 Archangelsky, S. and Villar de Seoane, L. 2004. Cycadean diversity in the Cretaceous of
856 Patagonia, Argentina. Three new *Androstrombus* species from the Baqueró Group. *Review*
857 *of Palaeobotany and Palynology* 131(1-2): 1-28.
- 858 Artabe, A.E., Zamuner, A.B. and Archangelsky, S. 1991. Estudios cuticulares en
859 Cycadópsidas fósiles. El género *Kurtziana* Frenguelli 1942. *Ameghiniana* 28(3-4): 365-374.
- 860 Artabe, A.E., Morel, E.M and Ganuza D.G. 2007a. Las floras triásicas de la Argentina.
861 *Publicación Electrónica de la Asociación Paleontológica Argentina* 11: 75-86.

862 Artabe, A.E., Morel, E.M., Ganuza, D.G., Zavattieri, A.M. and Spalletti, L.A. 2007b. La
863 paleoflora triásica de Potrerillos, provincia de Mendoza, Argentina. *Ameghiniana* 44(2):
864 279-301.

865 Bambach, R.K. 2006. Phanerozoic biodiversity mass extinctions. *Annual Review of Earth and*
866 *Planetary Sciences* 34: 127-155.

867 Barbacka, M., Pacyna, G., Kocsis, Á.T., Jarzynka, A., Ziaja, J. and Bodor, E. 2017. Changes
868 in terrestrial floras at the Triassic-Jurassic Boundary in Europe. *Palaeogeography,*
869 *Palaeoclimatology, Palaeoecology* 480: 80-93.

870 Belcher, C.M., Mander, L., Rein, G., Jervis, F.X., Haworth, M., Hesselbo, S.P., Glasspool, I.J.
871 and McElwain, J.C. 2010. Increased fire activity at the Triassic/Jurassic boundary in
872 Greenland due to climate-driven floral change. *Nature Geoscience* 3(6): 426-429.

873 Blackburn, T.J., Olsen, P.E., Bowring, S.A., McLean, N.M., Kent, D.V., Puffer, J., Mchone, G.,
874 Rasbury, E.T. and Et-Touhami, M. 2013. Zircon U-Pb geochronology links the End-Triassic
875 Extinction with the Central Atlantic Magmatic Province. *Science* 340(6135): 941-945.

876 Bodnar, J., Iglesias, A., Colombi, C.E. and Drovandi, J.M. 2019. Stratigraphical,
877 sedimentological and palaeofloristic characterization of the Sorocayense Group (Triassic)
878 in Barreal depocenter, San Juan Province, Argentina. *Andean Geology* 46(3): 567-603.

879 Bodnar, J., Morel, E.M., Coturel, E.P. and Ganuza, D.G. 2020. New plant fossil records and
880 biostratigraphic analysis from the Uspallata Group (Triassic) at Cacheuta Hill, Cuyo Basin,
881 west-central Argentina. *Geobios* 60: 3-27.

882 Bodnar, J., Cuesta, V., Escapa, I., and Nunes, C. 2023. Exploring the first appearance of the
883 main derived conifer families of Gondwana: evidence provided by the Triassic Woods from
884 Argentina. *Ameghiniana* 60(1): 18-47.

885 Bomfleur, B., Taylor, E.L., Taylor, T.N., Serbet, R., Krings, M. and Kerp, H. 2011. Systematics
886 and paleoecology of a new peltaspermalean seed fern from the Triassic polar vegetation of
887 Gondwana. *International Journal of Plant Sciences* 172(6): 807-835.

888 Bond, D.P.G. and Wignall, P.B. 2014. Large igneous provinces and mass extinctions: An
889 update. In: Keller, G. and Kerr, A.C. (eds.), *Volcanism, Impacts, and Mass Extinctions:*
890 *Causes and Effects*. Geological Society of America, Special Paper 505: 29-55.

891 Bonis, N.R. and Kürschner, W.M. 2012. Vegetation history, diversity patterns, and climate
892 change across the Triassic/Jurassic boundary. *Paleobiology* 38(2): 240-264.

893 Bonis, N.R., Kürschner, W.M. and Krystyn, L. 2009. A detailed palynological study of the
894 Triassic-Jurassic transition from key sections in the Eiberg Basin (Northern Calcareous
895 Alps, Austria). *Review of Palaeobotany and Palynology* 156(3-4): 376-400.

896 Bonis, N.R., Kürschner, W.M. and Van Konijnenburg-van Cittert, J.H.A. 2010. Changing CO₂
897 conditions during the end-Triassic inferred from stomatal frequency analysis on *Lepidopteris*
898 *ottonis* (Goeppert) Schimper and *Ginkgoites taeniatus* (Braun) Harris. *Palaeogeography,*
899 *Palaeoclimatology, Palaeoecology* 295(1-2): 146-161.

900 Caminos, R., Zamuner, A., Limarino, C. and Fauqué, L. 1995. Hallazgo de Triásico superior
901 en la Precordillera riojana. *Revista Asociación Geológica Argentina* 50(1-4): 262-265.

902 Capel, E., Cleal, C.J., Xue, J., Monnet, C., Servais, T. and Cascales-Miñana, B. 2022. The
903 Silurian–Devonian terrestrial revolution: Diversity patterns and sampling bias of the vascular
904 plant macrofossil record. *Earth-Science Reviews* 231: 104085.

905 Cascales-Miñana, B., Diez, J.B., Gerrienne, P. and Cleal, C.J. 2016. A palaeobotanical
906 perspective on the great end-Permian biotic crisis. *Historical Biology* 28(8): 1066-1074.

907 Cascales-Miñana, B., Servais, T., Cleal, C.J., Gerrienne, P. and Anderson, J. 2018. Plants—
908 the great survivors! *Geology Today* 34(6): 224-229.

909 Césari S.N., Drovandi J.M., Colombi C.E., Correa G. A. and Spalletti L.A. 2021. A new Late
910 Triassic palynological assemblage from western Gondwana (Carrizal Formation, Marayes
911 Basin, Argentina). *Comptes Rendus Palevol* 33: 677-700.

912 Chandler, M.A., Rind, D. and Ruedy, R. 1992. Pangaeon climate during the Early Jurassic:
913 GCM simulations and the sedimentary record of paleoclimate. *Geological Society of*
914 *America Bulletin* 104(5): 543-559.

- 915 Channing, A. and Edwards, D. 2013. Wetland megabias: ecological and ecophysiological
916 filtering dominates the fossil record of hot spring floras. *Palaeontology* 56(3): 523-556.
- 917 Choo, T., Escapa, I. and Benjamin, B. 2016. Monotypic colonies of *Clathropteris meniscioides*
918 (Dipteridaceae) from the Early Jurassic of central Patagonia, Argentina: implications for
919 taxonomy and palaeoecology. *Palaeontographica. Abteilung B, Palaophytologie* 294: 85-
920 109.
- 921 Cleal, C., Pardoe, H.S., Berry, C.M., Cascales-Miñana, B., Davis, B.A.S., Diez, J.B., Filipova-
922 Marinova, M.V., Giesecke, T., Hilton, J., Ivanov, D., Kustatscher, E., Lery, S.A.G.,
923 McElwain, J.C., Opluštil, S., Popa, M.E., Seyfullah, L.J., Stolle, E., Thomas, B.A. and Uhl,
924 D. 2021. Palaeobotanical experiences of plant diversity in deep time. 1: How well can we
925 identify past plant diversity in the fossil record? *Palaeogeography, Palaeoclimatology,*
926 *Palaeoecology* 576: 110481.
- 927 Colombi, C., Martínez, R.N., Césari, S. N., Alcober, O., Limarino, C.O. and Montañez, I. 2021.
928 A high-precision U–Pb zircon age constrains the timing of the faunistic and palynofloristic
929 events of the Carnian Ischigualasto Formation, San Juan, Argentina. *Journal of South*
930 *American Earth Sciences* 111: 103433.
- 931 Contreras, D.L., Escapa, I.H., Iribarren, R.C. and Cúneo, N.R. 2019. Reconstructing the early
932 evolution of the Cupressaceae: a whole-plant description of a new *Austrohamia* species
933 from the Cañadón Asfalto Formation (Early Jurassic), Argentina. *International Journal of*
934 *Plant Sciences* 180(8): 834-868.
- 935 Coturel, E.P., Morel, E.M. and Ganuza, D. 2016. Lycopodiopsids and equisetopsids from the
936 Triassic of Quebrada de los Fósiles formation, San Rafael Basin, Argentina. *Geobios* 49(3):
937 167-176.
- 938 Coughlin, T.J. 2000. Linked origin-oblique fault zones in the central Argentine Andes: the
939 basis for a new model for Andean orogenesis and metallogenesis. Ph D Thesis
940 (Unpublished), Universidad de Queensland, 207 pp.
- 941 Cúneo, R., Ramezani, J., Scasso, R., Pol, D., Escapa, I., Zavattieri, A.M. and Bowring, S.A.
942 2013. High-precision U–Pb geochronology and a new chronostratigraphy for the Cañadón

943 Asfalto Basin, Chubut, central Patagonia: Implications for terrestrial faunal and floral
944 evolution in Jurassic. *Gondwana Research* 24(3-4): 1267-1275.

945 Damborenea, S.E. and Manceñido, M.O. 1993. Piedra Pintada. In: Riccardi, A.C. and
946 Damborenea, S.E. (eds.), *Léxico estratigráfico de la Argentina IX: Jurásico*. Asociación
947 Geológica Argentina, Serie B: Didáctica y Complementaria 21: 313-316.

948 Damborenea, S.E., Manceñido, M.O. and Riccardi, A.C. 1975. Biofacies y estratigrafía del
949 Liásico de Piedra Pintada, Neuquén, Argentina. 1° Congreso Argentino de Paleontología y
950 Bioestratigrafía, Actas 2: 173-228. Tucumán.

951 Damborenea, S.E., Echevarría, J. and Ros-Franch, S. 2017. Biotic recovery after the end-
952 Triassic extinction event: Evidence from marine bivalves of the Neuquén Basin, Argentina.
953 *Palaeogeography, Palaeoclimatology, Palaeoecology* 487: 93-104.

954 De Barrio, R.E. 1993. El vulcanismo ácido jurásico en el noroeste de Santa Cruz, Argentina.
955 12° Congreso Geológico Argentino, Actas 4: 189-198. Mendoza.

956 de Jersey, N.J. and McKellar, J.L. 2013. The palynology of the Triassic–Jurassic transition in
957 southeastern Queensland, Australia, and correlation with New Zealand. *Palynology* 37(1):
958 77–114.

959 Del Fueyo, G.M., Carrizo, M.A., Poire, D.G. and Lafuente Díaz, M. 2021. Recurrent volcanic
960 activity recorded in araucarian wood from the Lower Cretaceous Springhill Formation,
961 Patagonia, Argentina: Palaeoenvironmental interpretations. *Acta Palaeontologica Polonica*,
962 66(1): 231-253.

963 Dromart, G., Garcia, J.P., Picard, S., Atrops, F., Lécuyer, C. and Sheppard, S.M.F. 2003. Ice
964 age at the Middle–Late Jurassic transition? *Earth and Planetary Science Letters* 213(3-4):
965 205-220.

966 Drovandi, J.M., Correa, G.A., Bodnar, J., Colombi, C.E., Coturel, E.P. and Morel, E.M. 2020.
967 A new paleofloristic assemblage from the Cuyana Basin (Agua de los Pajaritos depocenter),
968 Argentina and its paleobiogeographic and paleoenvironmental implications. *Journal of*
969 *South American Earth Sciences* 104: 102819.

970 Elgorriaga, A., Escapa, I.H., Bomfleur, B., Cúneo, R. and Ottone, E.G. 2015. Reconstruction
971 and phylogenetic significance of a new *Equisetum* Linnaeus species from the Lower
972 Jurassic of Cerro Bayo (Chubut Province, Argentina). *Ameghiniana* 52(1): 135-152.

973 Elgorriaga, A., Escapa, I.H. and Cúneo, N.R. 2019. Relictual *Lepidopteris* (Peltaspermales)
974 from the Early Jurassic Cañadón Asfalto Formation, Patagonia, Argentina. *International*
975 *Journal of Plant Sciences* 180(6): 578-596.

976 Escapa, I., Cúneo, R. and Axsmith, B. 2008. A new genus of the Cupressaceae (sensu lato)
977 from the Jurassic of Patagonia: implications for conifer megasporangiate cone homologies.
978 *Review of Palaeobotany and Palynology* 151(3-4): 110-122.

979 Escapa, I.H., Bomfleur, B., Cuneo, N.R. and Scasso, R. 2015. A new marattiaceous fern from
980 the Lower Jurassic of Patagonia (Argentina): the renaissance of *Marattiopsis*. *Journal of*
981 *Systematic Palaeontology* 13(8): 677-689.

982 Falco, J.I., Hauser, N., Olivera, D., Bodnar, J. and Reimold, W.U. 2021. A multi-proxy study of
983 the Cerro Piche Graben – a Lower Jurassic basin in the central North Patagonian Massif,
984 Argentina. *Journal of South American Earth Sciences* 109: 103287.

985 Foote, M. 1999. Morphological diversity in the evolutionary radiation of Paleozoic and Post-
986 Paleozoic Crinoids. *Paleobiology* 25(S2): 1–115.

987 Fox, C.P., Whiteside, J.H. Olsen, P.E., Cui, X., Summons, R.E., Idiz, E. and Grice, K. 2022.
988 Two-pronged kill mechanism at the end-Triassic mass extinction. *Geology* 50 (4): 448-453.

989 Gnaedinger, S. and Herbst, R. 2009. Primer registro de maderas gimnospérmicas de la
990 Formación Roca Blanca (Jurásico Inferior), provincia de Santa Cruz, Argentina.
991 *Ameghiniana* 46(1): 59-71.

992 Gnaedinger, S. and Zavattieri, A.M. 2021. A new Late Triassic dipteridacean fern from the
993 Paso Flores Formation, Neuquen Basin, Argentina. *Acta Palaeontologica Polonica* 66(4):
994 885-900.

995 Gnaedinger, S., Massini, J.L.G., Bechis, F. and Zavattieri, A.M. 2015. Coniferous woods and
996 wood-decaying fungi from the El Freno Formation (Lower Jurassic), Neuquén Basin,
997 Mendoza Province, Argentina. *Ameghiniana* 52(4): 447-467.

998 Götz, A.E., Ruckwield, K., Pálffy, J. and Haas, J. 2009. Palynological evidence of synchronous
999 changes within the terrestrial and marine realm at the Triassic/Jurassic boundary (Csovár
1000 section, Hungary). *Review of Palaeobotany and Palynology* 156(3-4): 401-409.

1001 Gravendyck, J., Schobben, M., Bachelier, J.B. and Kürschner, W.M. 2020. Macroecological
1002 patterns of the terrestrial vegetation history during the end-Triassic biotic crisis in the central
1003 European Basin: A palynological study of the Bonenburg section (NW-Germany) and its
1004 supra-regional implications. *Global and Planetary Change* 194: 103286.

1005 Guido, D., Escayola, M., de Barrio, R., Schalamuk, I. and Takashi Onoe, A. 2004. Edad y
1006 rasgos petrográficos y geoquímicos de cuerpos subvolcánicos asignables a la Formación
1007 Cerro León, este del Macizo del Deseado, Santa Cruz. *Revista de la Asociación Geológica
1008 Argentina* 59(4): 707-714.

1009 Gulisano, C.A. and Gutiérrez Pleimling, A.R. 1995. Guía de Campo: El Jurásico de la Cuenca
1010 Neuquina. Provincia de Mendoza. Asociación Geológica Argentina, Serie E, Guías de
1011 Campo 2: 1-103. Buenos Aires.

1012 Harris, P.J. and Little, C.T.S. 1999. The early Toarcian (Early Jurassic) and the Cenomanian-
1013 Turonian (Late Cretaceous) mass extinctions: similarities and contrasts. *Palaeogeography,
1014 Palaeoclimatology, Palaeoecology* 154(1-2): 39-66.

1015 Haworth, M., Elliott-Kingston, C., Gallagher, A., Fitzgerald, A. and McElwain, J.C. 2012.
1016 Sulphur dioxide fumigation effects on stomatal density and index of non-resistant plants:
1017 Implications for the stomatal palaeo-[CO₂] proxy method. *Review of Palaeobotany and
1018 Palynology* 182: 44-54.

1019 Herbst, R. 1965. La flora fósil de la Formación Roca Blanca, provincia de Santa Cruz,
1020 Patagonia, con algunas consideraciones geológicas y estratigráficas. *Opera Lilloana* 12: 1-
1021 101.

1022 Hesselbo, S.P., Morgans-Bell, H.S., McElwain, J.C., Rees, P.M., Robinson, S.A. and Ross,
1023 C.E. 2003. Carbon-cycle perturbation in the Middle Jurassic and accompanying changes in
1024 the terrestrial paleoenvironment. *The Journal of Geology* 111(3): 259-276.

- 1025 Holz, M. 2015. Mesozoic paleogeography and paleoclimates – a discussion of the diverse
1026 greenhouse and hothouse conditions of an alien world. *Journal of South American Earth*
1027 *Sciences* 61: 91-107.
- 1028 Iglesias, A., Artabe, A.E. and Morel, E.M. 2011. The evolution of Patagonian climate and
1029 vegetation from the Mesozoic to the present. *Biological Journal of the Linnean Society* 103
1030 (2): 409-422.
- 1031 Jalfin, G.A. and Herbst, R. 1995. La flora triásica del Grupo El Tranquilo, provincia de Santa
1032 Cruz (Patagonia). *Estratigrafía. Ameghiniana* 32(3): 211-229.
- 1033 Kershaw, P. and Wagstaff, B. 2001. The southern conifer family Araucariaceae: history,
1034 status, and value for paleoenvironmental reconstruction. *Annual Review of Ecology and*
1035 *Systematics* 32(1); 397-414.
- 1036 Kocsis, A.T., Reddin, C.J., Alroy, J. and Kiessling, W. 2019. The R package divDyn for
1037 quantifying diversity dynamics using fossil sampling data. *Methods in Ecology and*
1038 *Evolution* 10 (5): 735-743.
- 1039 Kocsis, A.T., Reddin, C.J. and Kiessling, W. 2022. Handout to the R package divDyn v0.8.2
1040 for diversity dynamics using fossil sampling data. Available from [https://CRAN.R-](https://CRAN.R-project.org/package=divDyn)
1041 [project.org/package=divDyn](https://CRAN.R-project.org/package=divDyn)
- 1042 Kustatscher, E., Ash, S.R., Karasev, E., Pott, C., Vajda, V., Yu, J. and McLoughlin, S. 2018.
1043 Flora of the Late Triassic. In: Tanner, L.H. (ed.), *The Late Triassic World. Topics in*
1044 *Geobiology* 46: 545-622, Springer, Cham.
- 1045 Lanés, S., Gnaedinger, S.C., Zavattieri, A.M., and Lezama, L. 2013. Sedimentary
1046 paleoenvironment and fossil plants of the El Freno Formation (early Jurassic) in Las Leñas
1047 valley, Neuquén basin. *Revista de la Asociación Geológica Argentina* 70(4): 465-476.
- 1048 Larsson, L.M. 2009. Palynostratigraphy of the Triassic–Jurassic transition in southern Sweden.
1049 *GFF* 131(1-2): 147-163.
- 1050 Lindström, S. 2016. Palynofloral patterns of terrestrial ecosystem change during the end-
1051 Triassic event – a review. *Geological Magazine* 153(2): 223-251.

- 1052 Lindström, S. 2021. Two-phased mass rarity and extinction in land plants during the End-
1053 Triassic climate crisis. *Frontiers in Earth Sciences* 9: 780343.
- 1054 Lindström, S., van de Schootbrugge, B., Hansen, K.H., Pedersen, G.K., Alsen, P., Thibault,
1055 N., Dybkjaer, K., Bjerrum, C.J. and Nielsen, L.H. 2017. A new correlation of Triassic-
1056 Jurassic boundary successions in NW Europe, Nevada and Peru, and the Central Atlantic
1057 Magmatic Province: A time-line for the end-Triassic mass extinction. *Palaeogeography,*
1058 *Palaeoclimatology, Palaeoecology* 478: 80-102.
- 1059 Lindström, S., Sanei, H., Van De Schootbrugge, B., Pedersen, G.K., Leshner, C.E., Tegner, C.,
1060 Heunisch, C., Dybkjaer, K. and Outridge, P.M. 2019. Volcanic Mercury and Mutagenesis in
1061 Land Plants during the End-Triassic Mass Extinction. *Science Advances* 5(10): eaaw4018.
- 1062 Lovecchio, J.P. and Naipauer; M. 2022. Proveniencia de la Formación Springhill en las
1063 cuencas Austral y Malvinas a partir de edades U-Pb en circones detríticos. 11º Congreso
1064 de Exploración y Desarrollo de Hidrocarburos, Actas: 291-313. Instituto Argentino del
1065 Petróleo y del Gas, Mendoza
- 1066 Lu, M.N. and Wang, R.S. 1987. Pollen and spore assemblages and distribution characteristics
1067 from Late Triassic to Early Jurassic epoch in Sichuan Basin. In: *Collections of Petroleum*
1068 *Stratum Paleontology Conferences*. Geological Publishing House, Beijing: 207-212 (in
1069 Chinese).
- 1070 Lucas, S.G. and Tanner, L.H. 2015. End-Triassic nonmarine biotic events. *Journal of*
1071 *Palaeogeography* 4(4): 331-348.
- 1072 Lundblad, A.B. 1959. Rhaeto–Liassic floras and their bearing on the stratigraphy of Triassic–
1073 Jurassic rocks. Stockholm. *Contributions to Geology* 3: 83–102.
- 1074 Mancuso, A.C., Benavente, C.A., Irmis, R.B. and Mundil, R. 2020. Evidence for the Carnian
1075 Pluvial Episode in Gondwana: New multiproxy climate records and their bearing on early
1076 dinosaur diversification. *Gondwana Research* 86: 104-125.
- 1077 Martínez, M.A. and Olivera, D.E. 2016. Jurassic organic-walled marine microplankton from the
1078 Neuquén Basin. Distribution, biostratigraphy and paleobiogeography. A review. In:
1079 Martínez, M. and Olivera, D. (eds.), *Palinología del Meso-Cenozoico de Argentina -*

1080 Volumen en homenaje a Mirta Elena Quattrocchio. Publicación Electrónica de la Asociación
1081 Paleontológica Argentina 16 (2): 106–128.

1082 Martínez, R.N., Sereno, P.C., Alcober, O.A., Colombi, C.E., Renne, P.R., Montañez, I.P. and
1083 Currie, B.S. 2011. A basal dinosaur from the dawn of the dinosaur era in southwestern
1084 Pangaea. *Science* 331(6014): 206-210.

1085 Marzoli, A., Renne, P.R., Piccirillo, E.M., Ernesto, M., Bellieni, G. and De Min, A. 1999.
1086 Extensive 200-million-year-old continental flood basalts of the Central Atlantic Magmatic
1087 Province. *Science* 284 (5414): 616-618.

1088 McElwain, J.C. and Punyasena, S.W. 2007. Mass extinction events and the plant fossil record.
1089 *Trends in Ecology and Evolution* 22(10): 548-557.

1090 McElwain J.C., Beerling D.J. and Woodward F.I. 1999. Fossil plants and global warming at the
1091 Triassic-Jurassic boundary. *Science* 285(5432): 1386-1390.

1092 McElwain, J.C., Popa, M.E., Hesselbo, S.P., Haworth, M. and Surlyk, F. 2007. Macroecological
1093 responses of terrestrial vegetation to climatic and atmospheric change across the
1094 Triassic/Jurassic boundary in East Greenland. *Paleobiology* 33 (4): 547-573.

1095 McGhee, G.R., Clapham, M.E., Sheehan, P.M., Bottjer, D.J. and Droser, M.L. 2013. A New
1096 Ecological-Severity Ranking of Major Phanerozoic Biodiversity Crises. *Palaeogeography,*
1097 *Palaeoclimatology, Palaeoecology* 370: 260-270.

1098 McNeely, J.A. 2003. Biodiversity in arid regions: values and perceptions. *Journal of Arid*
1099 *Environments* 54(1): 61-70.

1100 Miller, A.I. & Foote, M. 1996. Calibrating the Ordovician radiation of marine life: Implications
1101 for Phanerozoic diversity trends. *Paleobiology* 22(2): 304-309

1102 Morel, E.M., Ganuza, D.G. and Zúñiga, A. 1999. Revisión paleoflorística de la Formación Paso
1103 Flores, Triásico superior de Río Negro y del Neuquén. *Revista de la Asociación Geológica*
1104 *Argentina* 54(4): 389-406.

1105 Morel, E.M., Artabe, A.E. and Spalletti, L.A. 2003. The Triassic floras of Argentina:
1106 Biostratigraphy, Floristic events and comparison with other areas of Gondwana and
1107 Laurasia. *Alcheringa* 27(3): 231-243.

1108 Morel, E.M., Ganuza, D.G., Artabe, A.E. and Spalletti, L.A. 2013. Revisión de la paleoflora de
1109 la Formación Nestares (Jurásico Temprano), provincias del Neuquén y Río Negro,
1110 Argentina. *Ameghiniana* 50(5): 493-508.

1111 Morel, E.M., Artabe, A.E., Ganuza, D.G., Bodnar, J., Correa, G. and Spalletti, L.A., 2015. El
1112 Triásico de la Formación Carrizal en el depocentro de Marayes (San Juan, Argentina):
1113 paleobotánica, tafonomía y bioestratigrafía. *Revista de la Asociación Geológica Argentina*
1114 72(4): 456-469.

1115 Newell, N.D. 1952. Periodicity in invertebrate evolution. *Journal of Paleontology* 26 (3): 371-
1116 385.

1117 Olsen, P.E., Kent, D.V., Sues, H.D., Koeberl, C., Huber, H., Montanari, A., Rainforth, E.C.,
1118 Fowell, S.J., Szajna, M.J. and Hartline, B.W. 2002. Ascent of dinosaurs linked to an iridium
1119 anomaly at the Triassic-Jurassic boundary. *Science* 296(5571): 1305-1307.

1120 Pálffy, J. and Smith, P.L., 2000. Synchrony between Early Jurassic extinction, oceanic anoxic
1121 event, and the Karoo-Ferrar flood basalt volcanism. *Geology* 28(8): 747-750.

1122 Pálffy, J. and Kocsis, A.T. 2014. Volcanism of the Central Atlantic magmatic province as the
1123 trigger of environmental and biotic changes around the Triassic-Jurassic boundary. In:
1124 Keller, G. and Kerr, A.C. (eds.) *Volcanism, Impacts, and Mass Extinctions: Causes and*
1125 *Effects*. Geological Society of America, Special Paper 505: 245-261, Boulder.

1126 Pankhurst, R.J., Sruoga, P. and Rapela, C.W. 1993. Estudio geocronológico Rb–Sr de los
1127 complejos Chon Aike y El Quemado a los 47°30' L.S. 12° Congreso Geológico Argentino,
1128 Actas 4: 171–178, Mendoza.

1129 Pardoe, H.S., Cleal, C.J., Berry, C., Cascales-Miñana, B., Davis, B.A.S., Diez, J.B., Filipova-
1130 Marinova, M.V., Giesecke, T., Hilton, J., Ivanov, D., Kustatscher, E., Leroy, S.A.G.,
1131 McElwain, J.C., Opluštil, S., Popa, M.E., Seyfullah, L.J., Stolle, E., Thomas, B.A. and Uhl,
1132 D. 2021. Palaeobotanical experiences of plant diversity in deep time. 2: How to measure
1133 and analyse past plant biodiversity. *Palaeogeography, Palaeoclimatology, Palaeoecology*
1134 580: 110618.

- 1135 Parrish, J.T. 1993. Climate of the supercontinent Pangea. *The Journal of Geology* 101(2): 215-
1136 233.
- 1137 Petersen, H.I. and Lindström, S. 2012. Synchronous Wildfire Activity Rise and Mire
1138 Deforestation at the Triassic–Jurassic Boundary. *PLoS ONE* 7(10): e47236.
- 1139 Petriella, B. and Arrondo, O.G. 1984. La tafolórula liásica de la Estancia La Juanita, provincia
1140 de Santa Cruz, Argentina. *Ameghiniana* 21(1): 35–41.
- 1141 Pieńkowski, G., Niedźwiedzki, G. and Waksmundzka, M. 2012. Sedimentological,
1142 palynological and geochemical studies of the terrestrial Triassic–Jurassic boundary in
1143 northwestern Poland. *Geological Magazine* 149(2): 308-332.
- 1144 Pol, D., Mancuso, A.C., Smith, R.M., Marsicano, C.A., Ramezani, J., Cerda, I.A., Otero, A. and
1145 Fernandez, V. 2021. Earliest evidence of herd-living and age segregation amongst
1146 dinosaurs. *Scientific Reports* 11(1): 20023.
- 1147 Puebla, G.G., Mego, N. and Prámparo, M.B. 2012. Asociación de briofitas de la Formación La
1148 Cantera, Aptiano tardío, cuenca de San Luis, Argentina. *Ameghiniana* 49(2): 217-229.
- 1149 Pujana, R. 2022. Fossil woods from Argentina (1884–2021). *Revista Del Museo Argentino de*
1150 *Ciencias Naturales, Nueva Serie* 24(2): 217-240.
- 1151 Quattrocchio, M.E., Martínez, M.A. and Volkheimer, W. 2007. Las floras jurásicas de la
1152 Argentina. *Publicación Electrónica de la Asociación Paleontológica Argentina* 11: 87-100.
- 1153 Raup, D.M. 1978. Approaches to the Extinction Problem: Presidential Address to the Society
1154 November 8, 1977. *Journal of Paleontology* 52(3): 517-523.
- 1155 Raup, D.M. and Sepkoski, J.J. 1982. Mass extinctions in the marine fossil record. *Science* 215
1156 (4539): 1501–1503.
- 1157 Rogers, R.R., Swisher III, C.C., Sereno, P.C., Monetta, A.M., Forster, C.A. and Martinez, R.N.
1158 1993. The Ischigualasto tetrapod assemblage (Late Triassic, Argentina) and $^{40}\text{Ar}/^{39}\text{Ar}$
1159 dating of dinosaur origins. *Science* 260(5109): 794-797.
- 1160 Ruiz González, V., Renda, E. M., Vizán, H., Martín-Hernández, F., Palencia-Ortas, A. and
1161 Osete, M. L. 2024. Intraplate deformation during Gondwana breakup: a study of the Jurassic

1162 units of the Cañadón Asfalto Basin (extra-Andean Patagonia, Argentina). *Geophysical*
1163 *Journal International* 238(3): 1723-1741.

1164 Sagasti, A.J., Morel, E.M., Ganuza, D. and Knight, P.A. 2019. New paleofloristic elements and
1165 stratigraphic considerations for the Nestares Formation (Lower Jurassic, Argentina).
1166 *Journal of South American Earth Sciences* 94: 102245.

1167 Schoepfer, S.D., Algeo, T.J., van de Schootbrugge, B. and Whiteside, J.H. 2022. The Triassic-
1168 Jurassic transition – A review of environmental change at the dawn of modern life. *Earth-*
1169 *Science Reviews* 232: 104099.

1170 Serrano-Brañas, C.I. and Reyes-Luna, P.C. 2014. Paleobotánica forense: Una aproximación
1171 a la tafonomía de plantas. *Boletín de la Sociedad Geológica Mexicana* 66(1): 25-39.

1172 Skog, J.E. and Dilcher, D.L. 1994. Lower vascular plants of the Dakota Formation in Kansas
1173 and Nebraska, USA. *Review of Palaeobotany and Palynology* 80(1-2): 1-18.

1174 Slater, S.M., Twitchett, R.J., Danise, S. and Vajda, V. 2019. Substantial vegetation response
1175 to Early Jurassic global warming with impacts on oceanic anoxia. *Nature Geoscience* 12(6):
1176 462–467.

1177 Slodownik, M., Hill, R.S. and McLoughlin, S. 2023. *Komlopteris*: A persistent lineage of post-
1178 Triassic corystosperms in Gondwana. *Review of Palaeobotany and Palynology* 317:
1179 104950.

1180 Spalletti, L.A., Arrondo, O.G., Morel, E.M. and Ganuza, D.G. 1991. Evidencias sobre la edad
1181 Triásica de la Formación Lapa en la región de Chacaico, Provincia del Neuquén. *Revista*
1182 *de la Asociación Geológica Argentina* 46(3-4): 167-172.

1183 Spalletti, L.A., Artabe, A.E., Morel, E.M. and Brea, M. 1999. Biozonación paleoflorística y
1184 cronoestratigrafía del Triásico Argentino. *Ameghiniana* 36(4): 419-451.

1185 Spalletti, L.A., Fanning, M. and Rapela, C. 2008. Dating the Triassic continental rift in the
1186 southern Andes: the Potrerillos Formation, Cuyo basin, Argentina. *Geologica Acta* 6(3):
1187 267-283.

1188 Spalletti, L.A., Morel, E.M., Franzese, J.R., Artabe, A.E., Ganuza, D.G. and Zúñiga, A. 2007.
1189 Contribución al conocimiento sedimentológico y paleobotánico de la Formación El Freno

- 1190 (Jurásico Temprano) en el valle superior del río Atuel, Mendoza, Argentina. *Ameghiniana*
1191 44(2): 367-386.
- 1192 Spalletti, L., Franzese, J., Morel, E., D'elia, L., Zúñiga, A. and Fanning, C.M. 2010.
1193 Consideraciones acerca de la sedimentología, paleobotánica y geocronología de la
1194 Formación Piedra del Águila (Jurásico Inferior, Neuquén). *Revista de la Asociación*
1195 *Geológica Argentina* 66(3): 305-313.
- 1196 Steinthorsdottir, M., Jeram, M. and McElwain, J.C. 2011. Extremely elevated CO₂
1197 concentrations at the Triassic/Jurassic boundary. *Palaeogeography, Palaeoclimatology,*
1198 *Palaeoecology* 308: 418-432.
- 1199 Steinthorsdottir, M., Vajda, V. and Pole, M. 2018. Significant transient pCO₂ perturbation at the
1200 New Zealand Oligocene-Miocene transition recorded by fossil plant stomata.
1201 *Palaeogeography, Palaeoclimatology, Palaeoecology* 515: 152-161.
- 1202 Tidwell, W.D. and Ash, S.R. 1994. A review of selected Triassic to Early Cretaceous ferns.
1203 *Journal of Plant Research* 107: 417-442.
- 1204 Tomas, G.J. and Acuña, A.J. 2022. Comparaciones basadas en biomarcadores de crudos
1205 extraídos de la Formación Springhill (Cuenca Austral) y de la Formación Bajo Barreal
1206 (Cuenca Golfo San Jorge), Argentina. *Estudios Geológicos* 78(1): e143.
- 1207 Troncoso, A., Gnaedinger, S. And Herbst, R. 2000. *Heidiphyllum*, *Rissikia* y *Desmiophyllum*
1208 (Pinophyta, Coniferales) en el Triásico del norte chico de Chile y sur de Argentina.
1209 *Ameghiniana* 37(1), 119-125.
- 1210 Turner, S., Bean, L.B., Dettmann, M., McKellar, J., McLoughlin, S. and Thulborn, T. 2009.
1211 Australian Jurassic sedimentary and fossil successions: current work and future prospects
1212 for marine and non-marine correlation. *GFF* 131(1-2): 49-70.
- 1213 Ullmann, C.V., Tibault, N., Ruhl, M., Hesselbo, S.P. and Korte, C. 2014. Effect of a Jurassic
1214 oceanic anoxic event on belemnite ecology and evolution. *PNAS* 111(28): 10073–10076.
- 1215 Vakhrameev, V.A. 1991. Jurassic and Cretaceous floras and climates of the Earth. Cambridge
1216 University Press, 318 pp., Cambridge.

- 1217 Vajda, V., Calner, M. and Ahlberg, A. 2013. Palynostratigraphy of dinosaur footprint-bearing
1218 deposits from the Triassic-Jurassic boundary interval of Sweden. *GFF* 135(1): 120-130.
- 1219 Vajda, V., McLoughlin, S., Slater, S.M., Gustafsson, O. and Rasmusson, A.G. 2023. The
1220 “seed-fern” *Lepidopteris* mass-produced the abnormal pollen *Ricciisporites* during the end-
1221 Triassic biotic crisis. *Palaeogeography, Palaeoclimatology, Palaeoecology* 627: 111723.
- 1222 van de Schootbrugge B., Quan T.M., Lindström S., Püttmann W., Heunisch C., Pross J.,
1223 Fiebig, J., Petschick, R., Röhling, H.-G., Richoz, S., Rosenthal, Y. and Falkowski, P.G.
1224 2009. Floral changes across the Triassic/Jurassic boundary linked to flood basalt
1225 volcanism. *Nature Geosciences* 2(8): 589-594.
- 1226 van de Schootbrugge, B. and Wignall, P.B. 2016. A tale of two extinctions: converging end-
1227 Permian and end-Triassic scenarios. *Geological Magazine* 153(2): 332-354.
- 1228 van Konijnenburg-van Cittert, J.H.A. 1996. Two *Osmundopsis* species from the Mid Jurassic
1229 of Yorkshire and their sterile foliage. *Palaeontology* 39(3): 719-731.
- 1230 Van Konijnenburg-Van Cittert, J.H.A. 2002. Ecology of some late Triassic to early Cretaceous
1231 ferns in Eurasia. *Review of Palaeobotany and Palynology* 119(1-2): 113-124.
- 1232 Vizan, H., Geuna, S., Melchor, R., Bellosi, E.S., Lagorio, S.L., Vásquez, C., Japas, M.S., Ré,
1233 G. and Do Campo, M. 2013. Geological setting and paleomagnetism of the Eocene red
1234 beds of Laguna Brava Formation (Quebrada Santo Domingo, northwestern Argentina).
1235 *Tectonophysics* 583: 105-123.
- 1236 Volkheimer, W. and Papu, O.H. 1993. Una microflora del Triásico Superior de la Cuenca
1237 Malargüe, localidad Llantenes, provincia de Mendoza, Argentina. *Ameghiniana* 30(1): 93-
1238 100.
- 1239 Volkheimer, W., Rauhut, O.W., Quattrocchio, M.E. and Martínez, M.A. 2008a. Jurassic
1240 paleoclimates in Argentina, a review. *Revista de la Asociación Geológica Argentina* 63(4):
1241 549-556.
- 1242 Volkheimer, W., Quattrocchio, M.E., Cabaleri, N.G. and García, V. 2008b. Palynology and
1243 paleoenvironment of the Jurassic lacustrine Cañadón Asfalto Formation at Cañadón

- 1244 Lahuincó locality, Chubut Province, Central Patagonia, Argentina. *Revista Española de*
1245 *Micropaleontología* 40(1): 77-96.
- 1246 Wade-Murphy, J., Kuerschner, W.M. and Hesselbo, S.P. 2006. Early Toarcian vegetation
1247 History from the Korsodde Section of Bornholm (Denmark) and its Possible Impact on
1248 Terrestrial Carbon Isotope Records. 7th European Palaeobotany Palynology Conference,
1249 Program and Abstracts: 153-164, Prague.
- 1250 Wang, Y.D., Ni, Q., Jiang, Z.K. and Tian, N. 2008. Diversity variation and tempo-spatial
1251 distribution of *Otozamites* (Bennettitales) in the Mesozoic of China. *Palaeoworld* 17(3-4):
1252 222-234.
- 1253 Wang, Y.D., Fu, B.H., Xie, X.P., Huang, Q.S., Li, K., Li, G., Zs, L., Yu, J.X., Pan, Y., Tian, N.,
1254 and Jiang, Z. 2010. The Terrestrial Triassic and Jurassic Systems in the Sichuan Basin,
1255 China. Hefei: University of Science and Technology of China Press: 1-216. (in Chinese with
1256 English abstract).
- 1257 Whiteside, J.H., Olse, P.E., Kent, D.V., Fowell, S.J. and Et-Touhami, M. 2007. Synchrony
1258 between the Central Atlantic magmatic province and the Triassic-Jurassic mass-extinction
1259 event? *Palaeogeography, Palaeoclimatology, Palaeoecology* 244: 345–367.
- 1260 Wierzbowski, H., Rogov, M.A., Matyja, B. A., Kiselev, D. and Ippolitov, A. 2013. Middle–Upper
1261 Jurassic (Upper Callovian–Lower Kimmeridgian) stable isotope and elemental records of
1262 the Russian Platform: indices of oceanographic and climatic changes. *Global and Planetary*
1263 *Change* 107: 196-212.
- 1264 Wignall, P.B. and Atkinson, J.W. 2020. A two-phase end-Triassic mass extinction. *Earth-*
1265 *Science Reviews* 208: 103282.
- 1266 Ye, M.N., Liu, X.Y., Huang, G.Q., Chen, L.X., Peng, S.J., Xu, A.F. and Zhang, B. X. 1986. Late
1267 Triassic and Early–Middle Jurassic fossil plants from northeastern Sichuan. Anhui Science
1268 and Technology Publishing House, 141 pp., Hefei. (in Chinese with English summary).
- 1269 Zamuner, A.B., Zavattieri, A.M., Artabe, A.E. and Morel, E.M., 2001. Paleobotánica. In: Artabe,
1270 A.E., Morel, E.M. and Zamuner, A.B. (eds.), *El Sistema Triásico En La Argentina*.
1271 *Fundación Museo de La Plata “Francisco P. Moreno”*: 143-184, La Plata.

1272 Zhang, J., Lenz, O.K., Hornung, J., Wang, P., Ebert, M. and Hinderer, M. 2020. Palynology
1273 and the Eco-Plant model of peat-forming wetlands of the Upper Triassic Haojiagou
1274 Formation in the Junggar Basin, Xinjiang, NW China. *Palaeogeography, Palaeoclimatology,*
1275 *Palaeoecology* 556: 109888.

1276 Zhou, N., Xu, Y., Li, L., Lu, N., An, P., Popa, M. E., Kürschner; W.M., Zhang,X. and Wang, Y.
1277 2021. Pattern of vegetation turnover during the end-Triassic mass extinction: Trends of fern
1278 communities from South China with global context. *Global and Planetary Change* 205:
1279 103585.

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1282 **FIGURE AND TABLES LEGENDS**

	Three timer sampling completeness	
Stage	Species	Genera
Carnian		
Norian	0,72727273	0,8
Rhaetian	0,8	0,85714286
Hettangian	0,45454545	0,64705882
Sinemurian	1	1
Pliensbachian	1	1
Toarcian	1	1
Aalenian	0,95238095	1
Bajocian	1	1
Bathonian	1	1
Callovian	1	1
Oxfordian	1	1
Kimmeridgian	1	1
Tithonian	1	0,625
Berriasian		

1283

1284 **Table 1.** Values of three-timer sampling completeness of each stage for species and genera.

1285

Plant group / Stage	Ca	No	Rh	He	Si	Pl	To	Aa	Baj	Ba t	Cl	Ox	Ki	Ti	Be
Marchantiopsida i.s.	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Bryopsida i.s.	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Lycopodiaceae	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Pleuromeiaceae	1	1	0	0	0	0	0	0	0	0	0	0	0	0	0
Isoetaceae	0	0	0	0	0	0	0	0	0	1	1	1	1	0	0
Neocalamitaceae	4	1	0	1	3	0	0	0	0	0	0	0	0	0	0
Equisetaceae	1	1	1	1	4	5	3	1	1	2	2	3	3	0	0
Equisetales i.s.	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Asterothecaceae	1	1	1	1	1	0	0	0	0	0	0	0	0	0	0
Marattiaceae	0	0	1	1	2	1	0	0	0	0	0	0	0	0	0
Osmundaceae	7	6	5	6	11	8	5	2	2	7	7	7	7	2	2
Gleicheniaceae	0	0	0	0	2	1	1	1	1	0	0	0	0	0	1
Dipteridaceae	0	2	6	4	11	13	7	0	0	2	2	2	2	0	0
Matoniaceae	0	0	0	0	0	0	0	0	0	0	0	0	0	1	0
Dicksoniaceae?	1	0	1	0	0	2	2	0	0	1	1	2	2	1	0
Ferns i.s.	0	0	0	0	0	2	2	0	0	1	1	1	1	0	0
Umkomasiaceae	31	16	13	1	3	0	0	0	0	0	0	0	0	0	0
Peltaspermeae	7	0	0	0	0	0	2	2	2	0	0	1	1	0	0
Mattatiellaceae	4	2	2	1	1	0	0	0	0	0	0	0	0	0	0
Petriellaceae	5	1	1	0	1	0	0	0	0	0	0	0	0	0	0
Caytoniaceae	0	0	0	1	3	6	5	0	0	0	0	0	0	1	1
Pteridosperms i.s.	1	0	0	2	3	2	1	1	1	1	1	2	2	0	0

Zamiaceae	2	1	1	1	1	0	0	0	0	0	0	0	0	0	0	
Cycadales i.s.	5	3	4	0	0	0	0	0	0	2	2	2	2	0	1	
Nilssoniaceae	1	3	3	0	1	0	0	0	0	0	0	0	0	0	0	
Williamsoniaceae	0	0	0	1	1	1	1	0	0	0	0	0	0	0	0	
Benettitales i.s.	0	0	0	7	14	12	9	0	0	18	18	18	18	4	12	
Cycadophyta i.s.	6	4	5	2	3	0	0	0	0	0	0	0	0	0	0	
Ginkgoaceae	0	0	1	0	0	1	1	0	0	0	0	0	0	0	0	
Hamshawviaceae	2	1	0	0	0	0	0	0	0	0	0	0	0	0	0	
Ginkgoales i.s.	11	6	8	0	0	1	1	0	0	1	1	1	1	0	2	
Czekanowskiaceae	1	0	2	0	0	0	0	0	0	0	0	0	0	0	0	
Czekanowskiales i.s.	0	0	0	0	0	0	1	1	1	0	0	0	0	0	0	
Voltziaceae	5	3	4	1	1	0	0	0	0	0	0	0	0	0	0	
Hirmeriaceae	1	0	0	0	0	1	2	1	1	2	2	3	3	0	0	
Araucariaceae	1	0	0	1	2	3	3	1	1	13	13	14	14	2	2	
Cupressaceae	1	2	2	0	2	2	2	2	2	3	3	3	3	0	0	
Podocarpaceae	3	4	4	1	1	0	0	0	0	4	4	4	4	0	0	
Coniferales i.s.	0	6	7	4	4	6	9	5	5	14	14	15	15	3	4	
Gnetales i.s.	10	5	3	0	0	1	1	0	0	0	0	0	0	0	0	
Coniferophytes i.s.	0	0	1	0	0	1	1	0	0	0	0	0	0	0	0	
Gymnosperms i.s.	6	8	5	0	0	2	3	1	1	0	0	0	0	0	0	
Tracheophytes i.s.	0	0	1	1	4	3	5	3	3	5	5	5	5	0	0	
Total	12	2	77	82	38	79	74	67	21	21	77	77	84	84	14	25

1286

1287

1288 **Table 2.** Species richness per plant lineage (without range through assumption) in each stage

1289 of the Late Triassic-earliest Cretaceous interval based on occurrences from Argentina.

1290 Abbreviations: Ca = Carnian, No = Norian, Rh = Rhaetian, He = Hettangian, Si = Sinemurian,

1291 Pl = Pliensbachian, To =Toarcian, Aa = Aalenian, Baj = Bajocian, Bat = Bathonian, Ca =

1292 Callovian, Ox = Oxfordian, Ki = Kimmeridgian, Ti = Tithonian, Be = Berriasian, i.s.= incertae

1293 sedis.

1294

1295

	Ca	No	Rh	He	Si	Pl	To	Aa	Baj	Bat	Ca	Ox	Ki	Ti	Be
Marchantiopsida i.s.	█	---	---	---	---	---	---	---	---	---	---	---	---	---	---
Bryopsida i.s.	█	---	---	---	---	---	---	---	---	---	---	---	---	---	---
Lycopodiaceae	█	---	---	---	---	---	---	---	---	---	---	---	---	---	---
Pleuromeiaceae	█	█	---	---	---	---	---	---	---	---	---	---	---	---	---
Isoetaceae	█	---	---	---	---	---	---	---	---	█	█	---	---	---	---
Neocalamitaceae	█	█	█	█	█	---	---	---	---	---	---	---	---	---	---
Equisetaceae	█	█	█	█	█	█	█	█	█	█	█	█	█	---	---

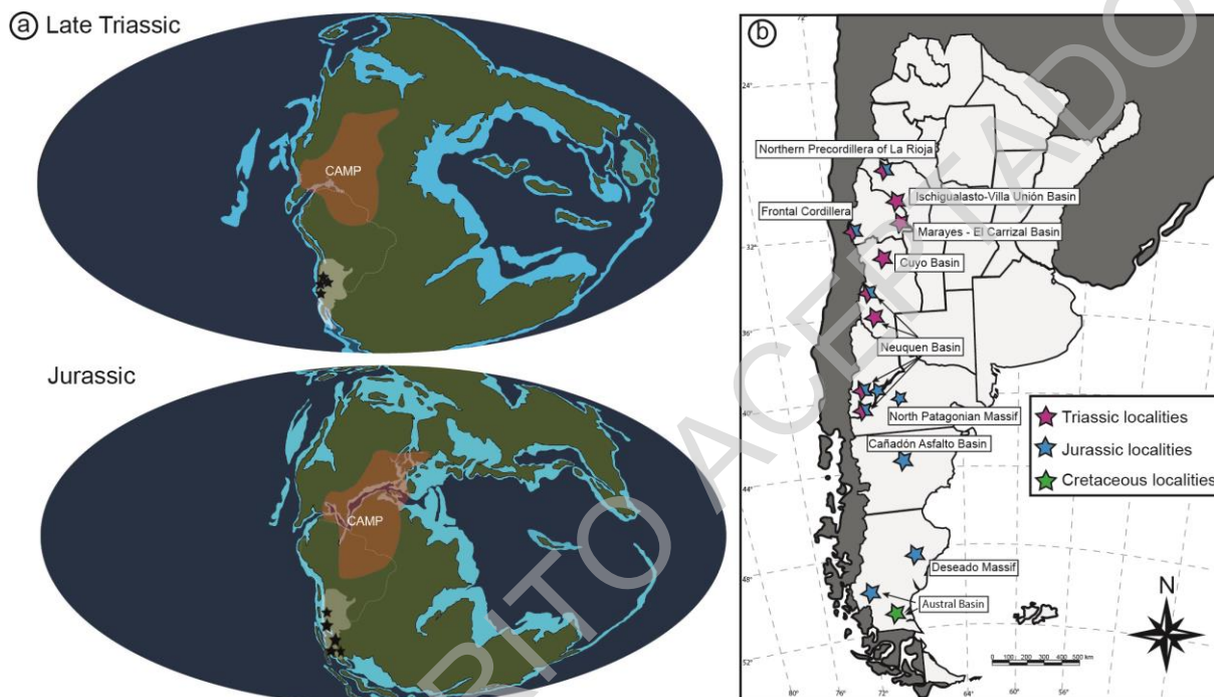
1302 Baj = Bajocian, Bat = Bathonian, Ca = Callovian, Ox = Oxfordian, Ki = Kimmeridgian, Ti =
1303 Tithonian, Be = Berriasian, i.s.= incertae sedis.

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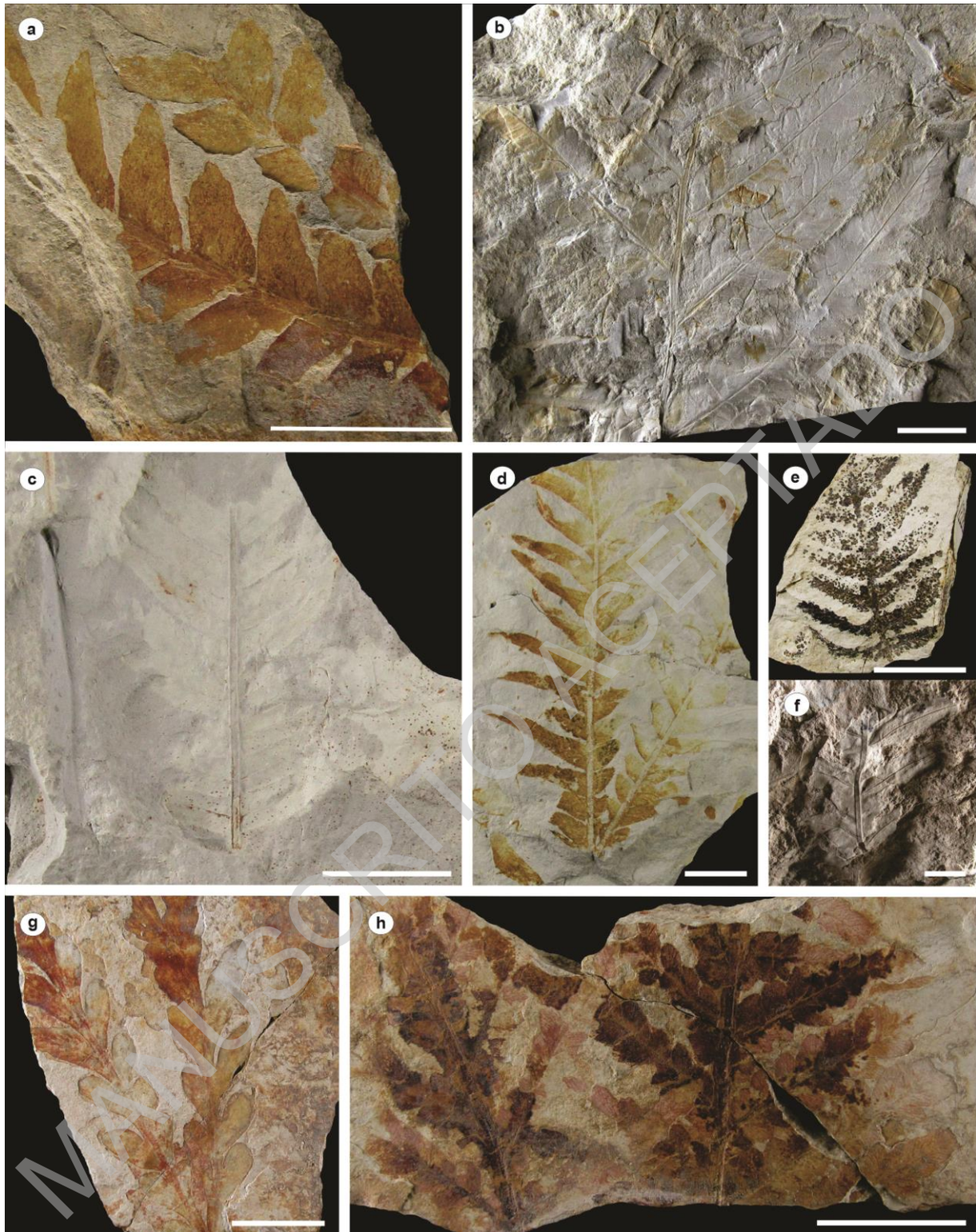
1307



1308

1309 **Figure 1. a**, Paleogeographic maps of the Late Triassic and Jurassic, showing the locations
1310 of the Argentine basins included in the work (maps modified from <http://www.scotese.com>); **b**,
1311 Fossiliferous localities with Late Triassic, Jurassic, and Early Cretaceous flora analyzed in this
1312 work.

1313

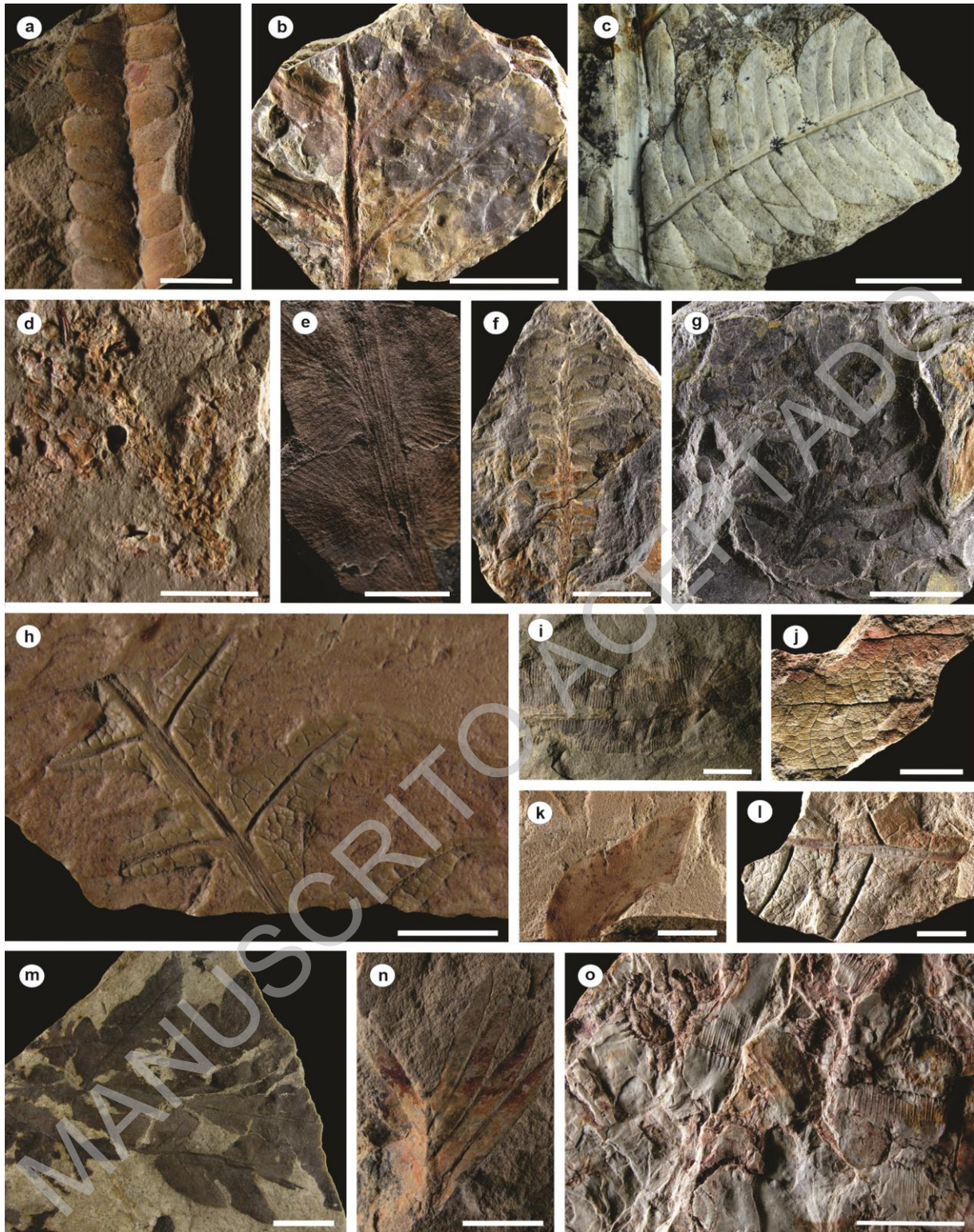


1314

1315 **Figure 2.** Representative flora of the Argentine Rhaetian; **a**, *Dicroidium odontopteroides*
 1316 (Umkomasiaceae; Paso Flores Formation; LPPB 12519 (scale bar= 2 cm); **b**, *Cladophlebis*
 1317 *kurtzi* (Osmundaceae; Paso Flores Formation; LPPB 12218 (scale bar= 2 cm); **c**,
 1318 *Cladophlebis mendozaensis* (Osmundaceae, Paso Flores Formation; LPPB 12219 (scale

1319 bar= 2 cm); **d**, *Dicroidium lancifolium* (Umkomasiaceae; Paso Flores Formation; LPPB 12232
1320 (scale bar= 2 cm); **e**, *Coniopteris harringtoni* (Dicksoniaceae?; Paso Flores Formation; LPPB
1321 12222 (scale bar= 1 cm); **f**, *Dictyophyllum rothi* (Dipteridaceae; Paso Flores Formation; LPPB
1322 12227 (scale bar= 2 cm); **g**, *Dicroidium crassum* (Umkomasiaceae; Paso Flores Formation;
1323 LPPB 12514 (scale bar= 1cm); **h**, *Zuberia sahnii* (Umkomasiaceae; Paso Flores Formation;
1324 LPPB 12520 (scale bar= 2 cm).
1325

MANUSCRITO ACEPTADO



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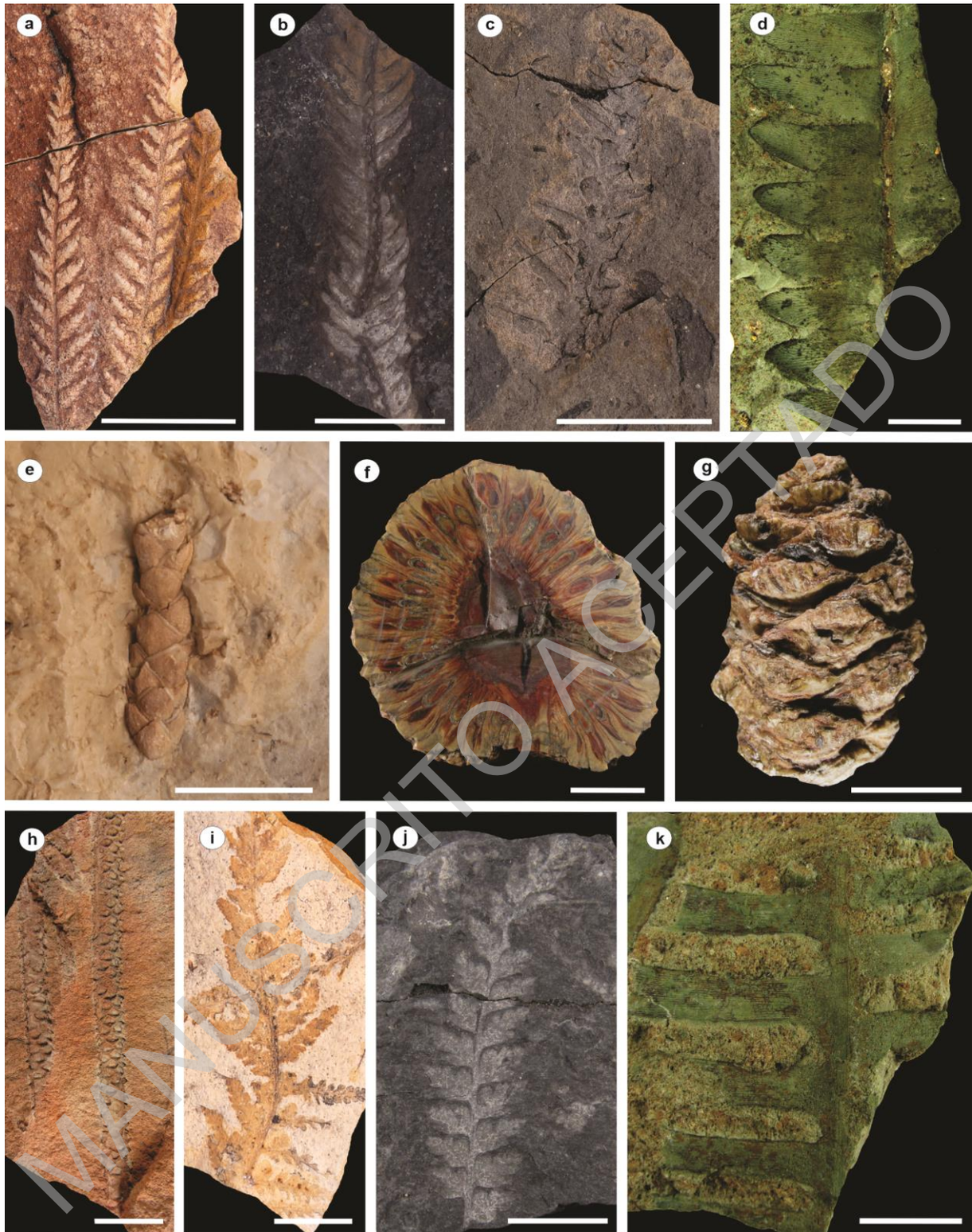
1327 **Figure 3.** Representative flora of the Argentine Early Jurassic; **a**, *Otozamites albosaxatilis*

1328 (*Bennettitales incertae sedis*; Roca Blanca Formation; LPPB 6020 (scale bar= 1 cm); **b**,

1329 *Archangelskya proto-loxoma* (*Pteridosperm incertae sedis*; El Freno Formation; LPPB 13200

1330 (scale bar= 2 cm); **c**, *Cladophlebis antartica* (*Osmundaceae*; Nestares Formation; MAPBAR

1331 5833 (scale bar= 2 cm); **d**, *Pagiophyllum* sp. (Coniferales *incertae sedis*; Lapa Formation;
1332 LPPB 12291 (scale bar= 2 cm); **e**, *Dicroidium odontopteroides* (Umkomasiaceae; Nestares
1333 Formation; MAPBAR 2317 (scale bar= 1 cm); **f**, *Elatocladus conferta* (Coniferales *incertae*
1334 *sedis*; El Freno Formation; LPPB 13210 (scale bar= 1 cm); **g**, *Komlopteris artabeae*
1335 (Umkomasiaceae; Nestares Formation; LPPB 12263 (scale bar= 2 cm); **h**, *Dictyophyllum*
1336 *apertum* (Dipteridaceae; Piedra Pintada Formation; LPPB 3959 (scale bar= 1 cm); **i**,
1337 *Marattiopsis muensteri* (Marattiaceae; El Freno Formation; LPPB 13174 (scale bar= 1 cm); **j**,
1338 *Clathropteris obovata* (Dipteridaceae; Piedra Pintada Formation; LPPB 3983 (scale bar= 2
1339 cm); **k**, *Sagenopteris nilssoniana* (Caytoniaceae; Piedra Pintada Formation; LPPB 5001 (scale
1340 bar= 2,cm); **l**, *Dictyophyllum rothi* (Dipteridaceae; Piedra Pintada Formation; LPPB 3953 (scale
1341 bar= 2 cm); **m**, *Goeppertella stipanicicii* (Dipteridaceae; Lonco Trapial Formation; LPPB 438
1342 (scale bar= 2 cm); **n**, *Ptilophyllum aculifolium* (Bennettitales *incertae sedis*; El Freno
1343 Formation; LPPB 13203 (scale bar= 1 cm); **o**, *Equisetites patagonica* (Equisetaceae; Roca
1344 Blanca Formation; LPPB 5987 (scale bar= 2 cm).
1345

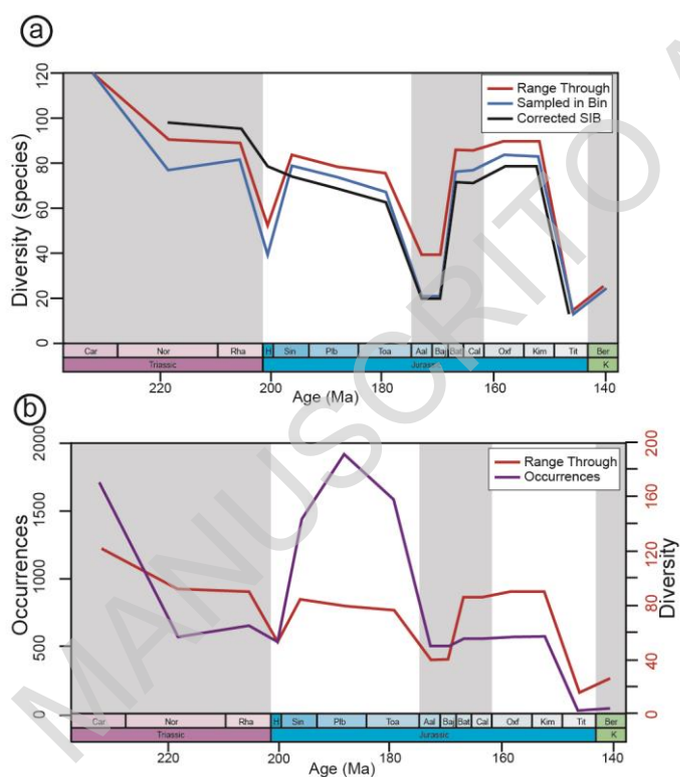


1346

1347 **Figure 4.** Representative flora of the Argentine Middle-Late Jurassic; **a**, *Korallipteris*
 1348 *potrerillensis* (Fern *incertae sedis*; Chon Aike Formation; MPM PB 15965 (scale bar= 2 cm);
 1349 **b**, *Otozamites sanctae-crucis* (Bennettitales *incertae sedis*; Chon Aike Formation; MPM PB
 1350 15840 (scale bar= 2 cm); **c**, *Elatocladus conferta* (Coniferales *incertae sedis*; Chon Aike

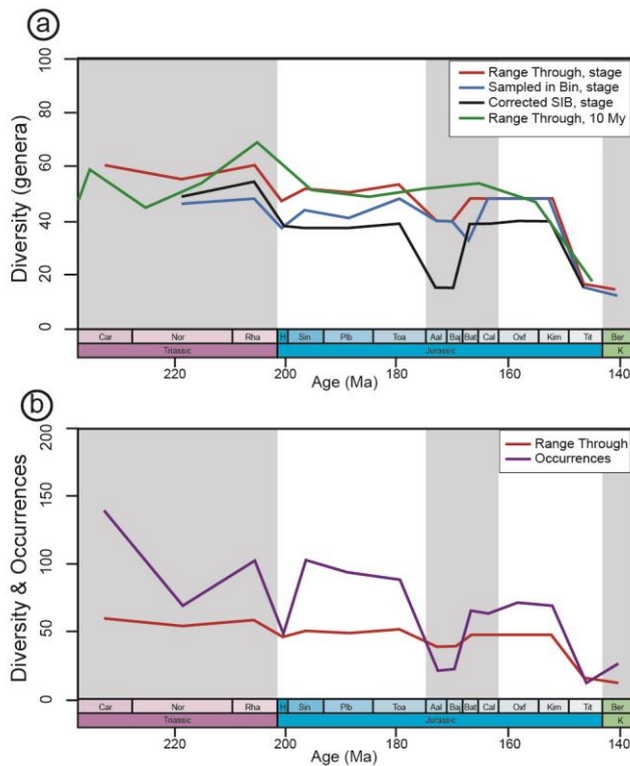
1351 Formation; MPM PB 16000 (scale bar= 2cm); **d**, *Dictyozamites latifolius* (Bennettitales *incertae*
1352 *sedis*; Chon Aike Formation; LPPB 11555 (scale bar= 2 cm), **e**, *Brachyphyllum lotenaense*
1353 (Coniferales *incertae sedis*; Chon Aike Formation; MPM PB 15982 (scale bar= 1cm), **f**, ovulate
1354 cone of *Araucaria mirabilis* (Araucariaceae; La Matilde Formation; LPPB 8079 (scale bar= 2
1355 cm), **g**, *Pararaucaria patagonica* (Hirmerellaceae; La Matilde Formation; LPPB 13748 (scale
1356 bar= 1 cm); **h**, leafy shoot of *Araucaria mirabilis* (Araucariaceae; La Matilde Formation; LPPB
1357 13755 (scale bar= 2 cm); **i**, *Scleropteris vincei* (Tracheophyte *incertae sedis*; Chon Aike
1358 Formation; MPM PB 16006 (scale bar= 1 cm); **j**, *Coniopteris hymenophylloides*
1359 (Dicksoniaceae?; Springhill Formation; LPPB 11161 (scale bar= 1cm); **k**, *Pseudoctenis*
1360 *eathiensis* (Cycadales *incertae sedis*; Chon Aike Formation; LPPB 11543 (scale bar= 2 cm)

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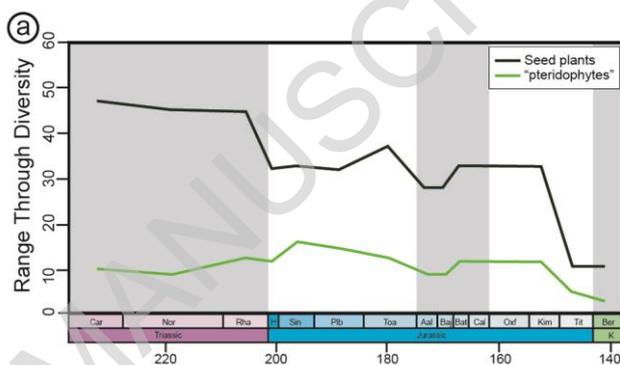
1363 **Figure 5. a**, Diversity (richness) curves at species level in the Late Triassic- Late Jurassic
1364 interval of Argentina; **b**, Range-through diversity compared with the occurrences of each
1365 species per stage in Argentina. In black numbers, scale used for the occurrences; in red
1366 number, scale used for the diversity curve.



1367

1368 **Figure 6. a**, Diversity (richness) curves at genera level in the Late Triassic- Late Jurassic
 1369 interval of Argentina; in green, range-through diversity with a 10-million-year bin resolution; **b**,
 1370 Range-through diversity compared with the occurrences of each genus per stage in Argentina.

1371

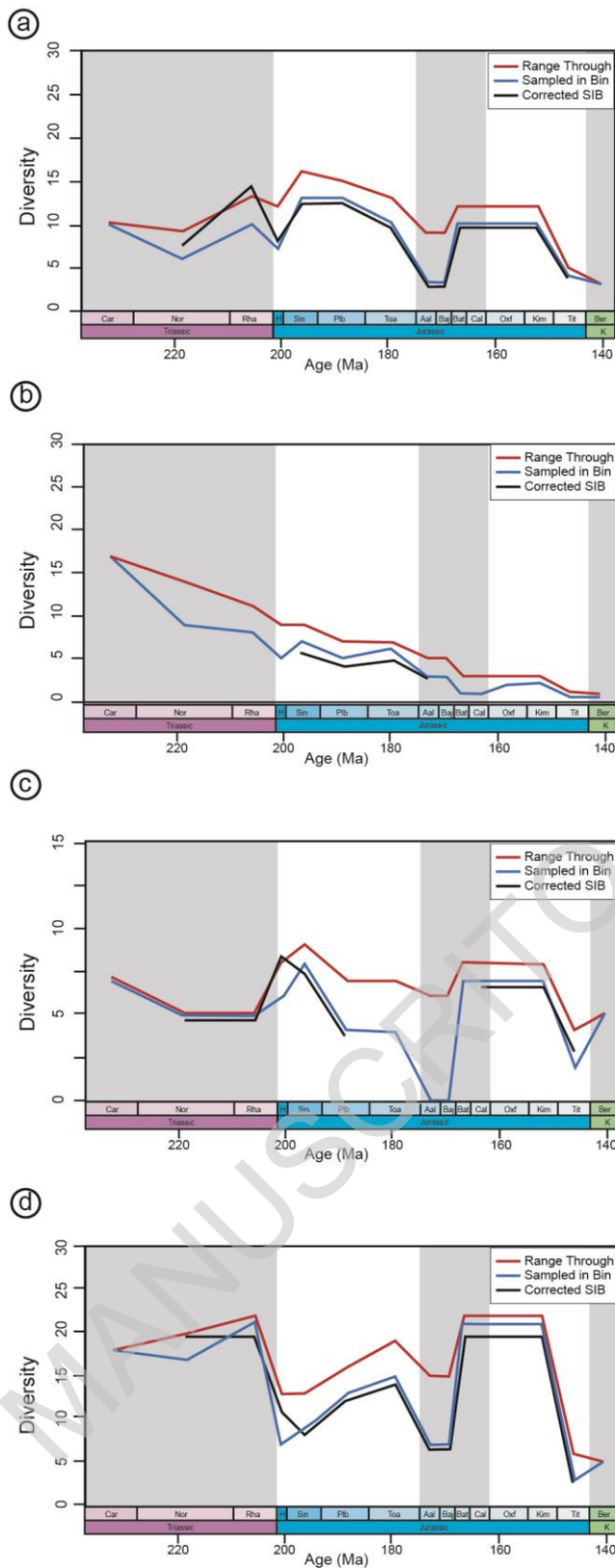


1372

1373 **Figure 7.** Comparison of the range-through diversity at generic level of the seed plants
 1374 (gymnosperms) and pteridophytes or free-spring vascular plants (lycophytes, sphenophytes,
 1375 and ferns) in the Late Triassic- Late Jurassic interval of Argentina.

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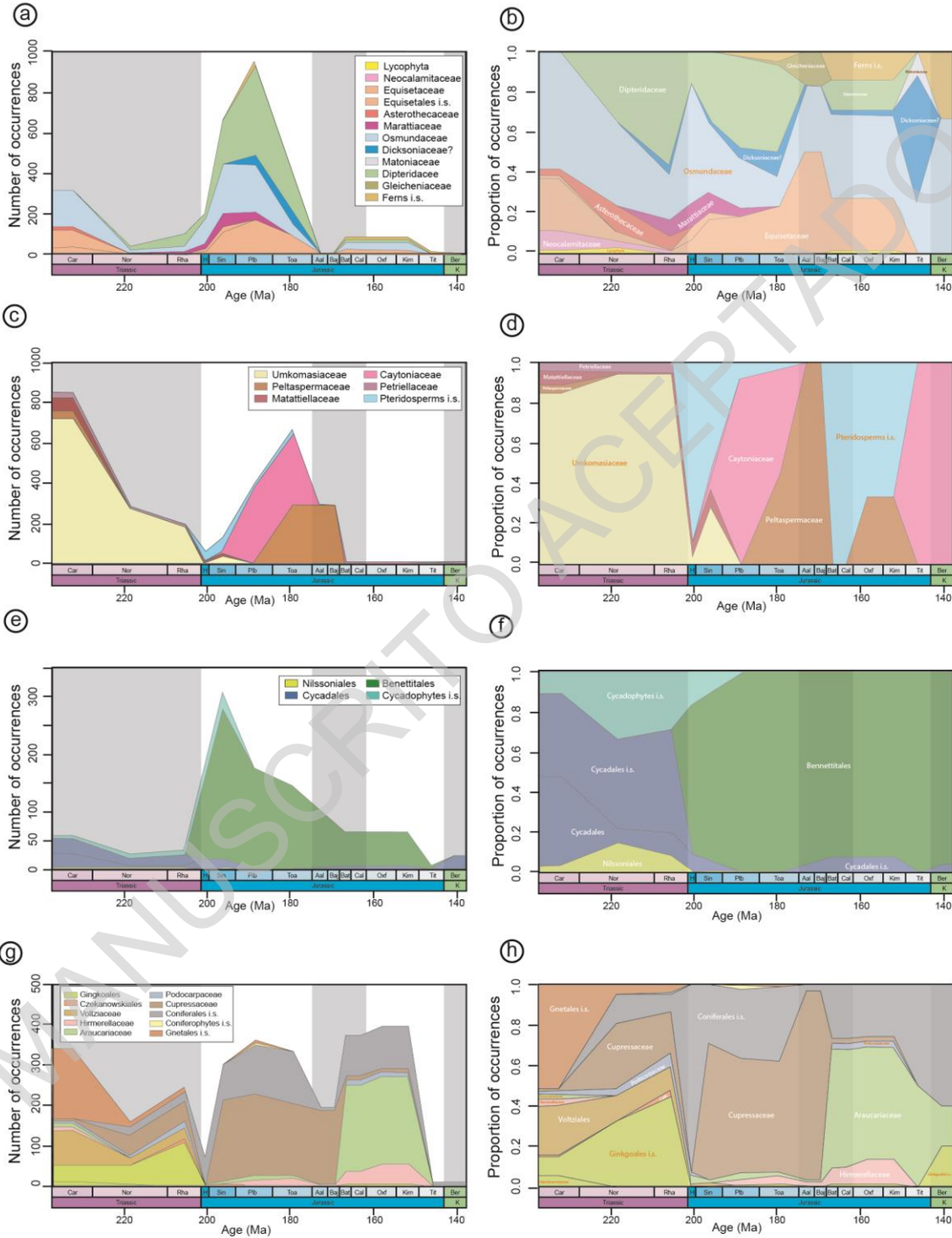
1378

1379 **Figure 8.** Diversity (richness) curves at genera level of different plant groups in the Late

1380 Triassic-Late Jurassic interval of Argentina; **a**, pteridophytes; **b**, pteridosperms or seed ferns;

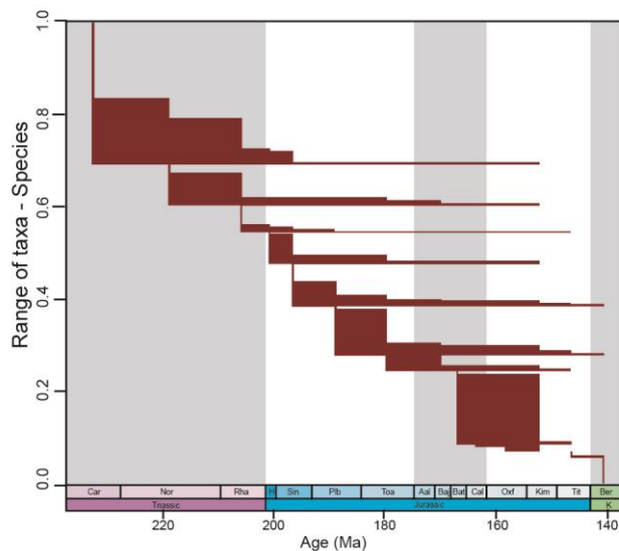
1381 **c**, cydadophytes; **d**, coniferophytes. CSIB is cut when the values three timer sampling
 1382 completeness are 1.

1383

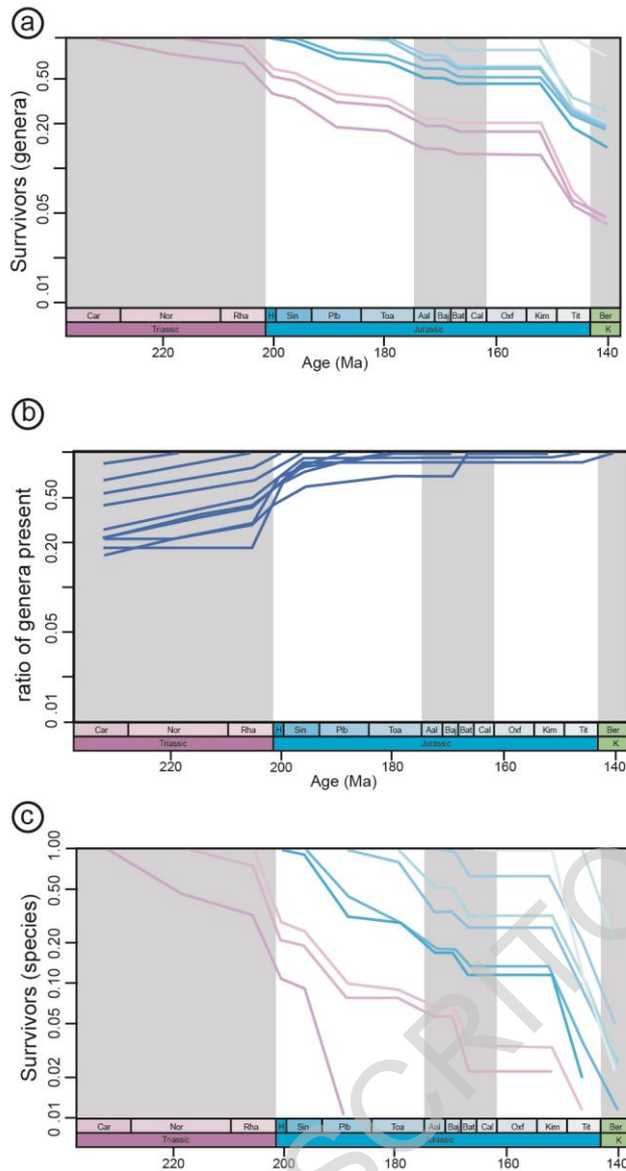


1384

1385 **Figure 9.** Number of occurrences and proportion of occurrences of each plant group in the
1386 Late Triassic-Late Jurassic interval of Argentina; **a-b**, pteridophytes; **c-d**, pteridosperms; **e-f**,
1387 cycadophytes; **g-h**, coniferophytes.
1388



1389 **Figure 10.** Patterns of FAD-LAD of fossil plants species in the Late Triassic-Late Jurassic
1390 interval of Argentina.
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1392



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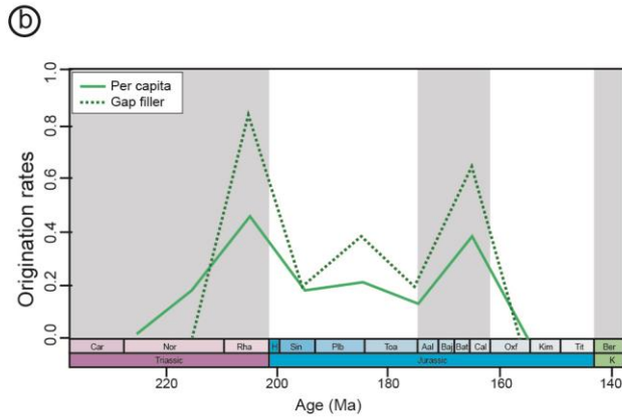
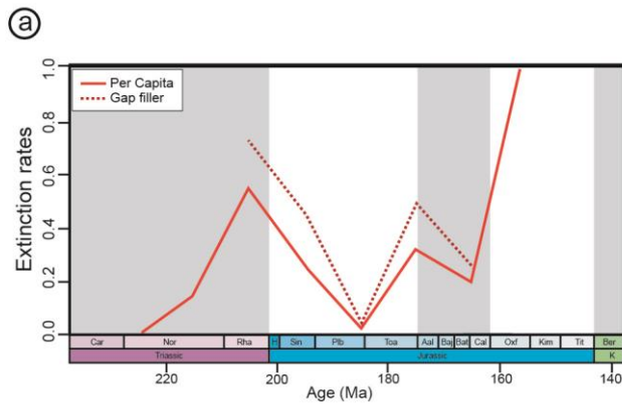
1394 **Figure 11. a**, “Forward” survivorship proportion of plant genera in the Late Triassic-Late

1395 Jurassic interval of Argentina; **b**, “Backward” survivorship proportion of plant genera in the

1396 Late Triassic-Late Jurassic interval of Argentina; **c**, “Forward” survivorship proportion of plant

1397 species in the Late Triassic-Late Jurassic interval of Argentina;

1398



1399

1400 **Figure 12. a**, Extinction rates at genera-level in the Late Triassic-Late Jurassic interval of
 1401 Argentina; **b**, Origination rates at genera-level in the Late Triassic-Late Jurassic interval of
 1402 Argentina.