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End Permian to Middle Triassic plant species richness and abundance patterns in South China

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- End Permian to Middle Triassic plant species richness and
- 2 abundance patterns in South China: coevolution of plants and
- 3 the environment through the Permian–Triassic transition
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24	ABSTRACT This study reviews plant species richness and abundance change from
25	the End Permian to Middle Triassic in South China and examines the co-evolutionary
26	relationship between the flora and the environment through this critical interval in the
27	history of terrestrial biotas. A normalized macro-fossil plant record, that considers
28	only one taxon per whole plant, is produced. This identifies four broad phases of plant
29	evolution. Phase 1 is marked by pre-extinction floras that demonstrate a long-term
30	decline of species richness beginning in the Late Permian (lower Changhsingian) that
31	culminates in the distinct End Permian Plant Crisis (EPPC) at the end of the
32	Changhsingian. Other evidence for the health of the flora, including palynology,
33	biomarkers, wildfire proxies, soil erosion and weathering proxies show a drastic loss
34	of plant abundance (biomass) and increase of wildfire frequency (suggestive of
35	increasing seasonal aridity) during the EPPC. A Phase 2 survival interval, during the
36	Changhsingian-Griesbachian transition, has a severely impoverished plant
37	assemblage consisting of opportunistic lycopods and a short-lived holdover flora.
38	Phase 3 (Late Griesbachian-Smithian) saw the modest recovery of species richness as
39	several groups began to radiate, notably conifers and ferns. Diversity increases
40	substantially and persistently during the succeeding Phase 4 and sees the dominant
41	lycopods/herbaceous bryophytes of Phase 3 replaced by conifer-dominated floras.
42	Plant abundance recovery began earlier than the resumption of coal formation which
43	only initiated in the Anisian following its disappearance during the EPPC. Only in the
44	Late Triassic did the flora recover to a level comparable to that seen in the Permian.
45	The flora of South China thus took ~15 million years to completely recover from the
46	profound environmental and climatic effects of the Permo-Triassic mass extinction.
47	
48	Keywords: plant-environment coevolution, mass extinction, coal gap, Permo-Triassic
49	and end-Permian, gigantopterids, lycopod
50	
51	1. Introduction
52	The 60 million years from the Middle Permian to the end-Triassic was one of the
53	most stressful in life's history. It witnessed the Permo-Triassic Mass Extinction

- 54 (PTME), the most severe crisis of the Phanerozoic, together with the end-Triassic mass extinction and several lesser crises (e.g., Wignall, 2015). The result was a 55 56 fundamental change of incumbents in both marine and terrestrial realms. Terrestrial tetrapod dynasties changed several times. The dinocephalians of the Middle Permian 57 were a distant memory by the time dinosaurs rose to dominance in the Jurassic. The 58 composition of plant communities also underwent fundamental overhaul during the 59 Permian and Triassic but the nature of this transition and its relationship with other 60 61 biosphere changes has long remained enigmatic (e.g., Knoll, 1984; Rees, 2002). In 62 particular, it is unclear if plant communities underwent a series of abrupt mass extinctions, such as seen amongst the marine biota and tetrapods, or instead showed 63 more gradual long-term changes (McElwain & Punyasena, 2007). 64 In an influential study, Knoll (1984) suggested that there was a protracted 65 66 changeover from a Paleophytic to a Mesophytic flora separated by a transitional or mixed flora, but with no abrupt extinction event at the Permo-Triassic boundary. 67 Subsequent work suggests that this distinction is between floras from different biomes 68 69 and has no chronostratigraphic significance (DiMichele et al., 2008). Nonetheless, the idea that plants did not suffer mass extinction, even during the Permo-Triassic 70 71 transition survives. Many studies based on literature compilations favour a noncatastrophic floral history at the end of the Permian (e.g., Rees, 2002; Ouyang and 72 Zhu, 2007; Nowak et al., 2019), and some field studies also support the notion that 73 there were relatively few losses at this time (e.g., Krassilov & Karasev, 2009; Hochuli 74 75 et al., 2010; Xiong & Wang, 2011; Yang et al., 2021). In contrast, there is clear evidence for a major upheaval in plant communities at the end of the Permian, 76 77 including the short-term proliferation of fungal spores (Reduviasporonites) at the expense of plant palynomorphs (Visscher et al., 1996), the abrupt loss of 78 palynomorphs from woody species suggesting forest die-off (Looy et al., 1999) and 79 the abrupt and prolonged disappearance of coals from the geological record: a 20 myr 80 "coal gap" (Retallack, 1995, 1996). Clearly, something happened to plants at the end 81 82 of the Permian (Yu et al., 2015).
 - Prior to the PTME, plants are divided into four main paleogeographic regions

- 84 (e.g., Hilton and Cleal, 2007) whilst low diversity survivors were similar after the
- crisis (Grauvogel-Stamm and Ash., 2005; Yu et al., 2015; Feng et al., 2020).
- 86 Compositionally, the survivors consist of holdover elements of the late Permian floras,
- 87 notably in South China, together with a few, new Triassic forms (Yao et al., 1980;
- 88 Chen et al., 2011; Yu et al., 2015). Some have argued, based on the palynological
- 89 record from South China, Xinjiang and Greenland, that the main plant crisis occurred
- at the end of the Griesbachian, the first substage of the Triassic (Qu et al., 1986;
- 2004; Yu et al., 2008; Peng et al., 2009; Hochuli et al., 2016). If this
- proved to be a global phenomenon then the fortunes of plants would be fundamentally
- out of kilter with that of terrestrial tetrapods and the marine biosphere (cf. Stanley,
- 94 2009; Sun et al., 2012; Song H.J. et al., 2018; Allen et al., 2020; Romano et al., 2020).
- 95 It is notable that the onset of the "coal gap", marking the loss of wetland peat-forming
- ommunities, coincided with the PTME and not the end of the Griesbachian.
- 97 Following the PTME, floras were of low diversity in the Early Triassic (e.g., Yu et
- al., 2015; Feng et al., 2020). Unlike the prosperous Late Triassic floras such as the
- 99 Baoding, Jiuliang, Xujiahe, Shazhenxi floras in South China, the Early to Middle
- floras are poorly understood (Li, 1964; Xu et al., 1979; Huang and Lu, 1992; Meng et
- al., 1994; Li et al., 1995). Recent discoveries of Early to Middle Triassic floras have
- helped fill in the blanks at this time (Meng et al., 1995; Broutin et al., 2020). Clearly,
- we have yet to achieve a full understanding of the dynamics of terrestrial recovery
- following the PTME, and questions still remain: were plants marching to a different
- beat compared to the marine biota or are the major differences caused by a poor
- understanding of the floral record, and why is there a "coal gap" during the Early and
- much of the Middle Triassic?
- To address these questions, we here present a comprehensive review of the
- 109 Permian to Triassic fossil plant assemblages of South China supplemented with
- extensive additional data from our own field collecting. This region comprises the
- Kangdian Oldland to the west and the Cathaysia Oldland to the East that were
- separated by the upper Yangtze Platform. Intensive studies during the past few
- decades have provided a major increase in our understanding of the flora which have

substantially increased our knowledge of changes in low paleolatitudes. We also incorporate information from other proxies for the health of the terrestrial biosphere. These include levels of total organic carbon (TOC), weathering proxies, carbon isotopic variations and black carbon and charcoal concentrations (as wildfire indicators). Our synthesis of fossil plant occurrences is placed in a detailed stratigraphic framework and, where available, zircon-derived radiometric dating (Shen S.Z. et al., 1995, 2011; Burgess et al., 2014; He et al., 2017). Our synthesis of the floral species richness and abundance differs from previous analyses (e.g., Yu et al., 2015; Chu et al., 2016; Feng et al., 2020) by normalizing the plant fossil occurrences to remove duplications that are an artefact of paleobotanical systematics, preservational types, and taxon recording (see Hilton and Cleal, 2007; Cleal et al., 2021). This provides a more realistic measure of plant species richness in the fossil record and unsurprisingly produces a substantially lower estimate of floral species richness than previous accounts.

2. Materials and methods

We use a combination of approaches to evaluate plant species turnover from the Middle Permian to the early Late Triassic in South China. The successive floras are documented in stratigraphic order (Fig. 1) and for each we review their stratigraphical relationships and depositional sedimentary facies to provide a taphonomic context for the palaebotanical occurrences.

2.1. Sampling strategy

Due to an absence of marine fauna and diachronism of plant-bearing units, we use the concept of an end-Permian plant crisis (EPPC) to represent the point of disappearance of plant macrofossils in the terrestrial South China successions. The EPPC is composed of two phenomena. One is the gradual alternation of plant assemblages during the Changhsingian associated with a gradual decline of species richness. The second phenomenon was a much more rapid extinction interval, at the

end of the Changhsingian, when many long-ranging elements disappeared, and plant abundance declined markedly.

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For fossil plants we have adopted two different levels of stratigraphic 145 resolution. For formations recording the EPPC of the Dalong, Xuanwei and Kayitou 146 formations (Fig. 1), and its Lower to Middle Triassic aftermath in the Feixianguan, Lingwen and Badong formations (Fig. 1), we selected representative localities in 148 terms of their fossil plant composition. For the Xuanwei Formation, we systematically collected plants from the Chahe and Chinahe sections in Guizhou and Yunnan provinces, for the Dalong Formation the Xinmin and Duanshan A, B sections in Guizhou Province, for the Feixianguan Formation the Chinahe, Tucheng and Mide 152 sections in Yunnan and Guizhou provinces and the Pojiao and Lubei sections in 153 Yunnan Province, for the Lingwen Formation the Lingwen Section in Hainan 154 155 Province, and for the Badong Formation the Hongjiaguan and Furongqiao sections in Hunan Province. For each formation we have undertaken extensive fieldwork to 156 identify and collect fossil plants in situ on a bed-by-bed basis to investigate them in 158 stratigraphically high-resolution, with the floras of the Feixianguan being reported here in detail for the first time. For these formations all the fossils have been identified by the same individuals using reference materials such that the identifications are internally consistent and accurate, rather than being based solely on literature compilation. To reduce the influence of "Signor-Lipps" effect, all the 162 published plant fossil records from each studied flora in South China are collected and 163 used in calculating species richness and stratigraphical distributions. We have not applied statistical methods to correct for the Signor-Lipps effect (e.g. Marshall and 165 166 Ward, 1996; Wang et al., 2014) due to the non-uniform rates of deposition, diachroneity, depositional hiatuses and taphonomic controls in terrestrial strata that affect fossil plant distribution and preservation. The location of all the included 168 sections are marked on the paleogeographic map (Fig. 3, 8, 13). For the floras of the Liangshan, Maokou and Longtan formations that predate the 170 EPPC, and for the Upper Triassic floras of the Jiuligang, Daqiaodi and Dajing

formations (Fig. 1), we have conducted lower-resolution investigations and have

summarized the sedimentology and floral compositions, but do not provide detailed stratigraphic ranges for individual species as that information is not presently available. Data collected from these sections can only be evaluated at stage level rather than to show origination and extinction rates within the respective formations.

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2.2. Evaluating fossil plant species richness and normalizing data occurrences While ideally our study would seek to identify species diversity, it is rarely possible to determine species evenness from the fossil plant record (Cleal et al., 2012, 2021). Here we focus on species richness that denotes the number of species present in a particular bed, locality, formation or flora and is measurable from the plant fossil record (see supplementary dataset). To assess plant species richness, it is necessary to evaluate patterns of species change over time (e.g., Li et al., 1995; Peng et al., 2009; Yu et al., 2015; Chu et al., 2016; Feng et al., 2020). This is achieved by constructing stratigraphic range diagrams for each species showing their first appearance datum (= origination) and last appearance datum (= extinction) in each geological section, and then correlating using litho-, bio- or chemostratigrpahic methods to compile stratigraphic range charts. From this information the total number of species can be determined at particular time intervals, and origination and extinction rates calculated noting these are when species originate and go extinct in South China within the dataset. We do not extend the range of plant fossils by calculating confidence intervals or interpolation due to the complexity of terrestrial stratal deposition and taphonomy, utilizing the the stratigraphic range of the raw, plant fossil ocurrences as recorded in the field. In order to provide values for the Middle Permian Maokou Flora, we have also included data from the latest early Permian Qixia Flora to provide range-through data (e.g., Cleal et al., 2012, 2021). Likewise, to provide the same for the early Late Triassic Shazhenxi Flora, we have included data from latest Triassic (Rhaetian) Yangbaichong Flora (see supplementary data for additional details). To meaningfully extract plant species richness patterns from the fossil data, it is important to eliminate duplicates that are an artefact of paleobotanical nomenclature

(e.g., Hilton and Cleal, 2007; Cleal et al., 2012, 2021). A single reconstructed whole-

plant species in the fossil record includes numerous distinct organs (e.g., leaves, stems, roots, cones, seeds), each with their own generic and species name (see Chaloner, 1986; Bateman and Hilton, 2009). The solution we have adopted is to normalize the data (see Cleal et al., 2012, 2021) and evaluate only those organs whose fossil taxonomy is most likely to reflect the original whole organism taxonomy. This represents the first time this approach has been applied to plant species richness through the Permo-Triassic interval including the EPPC and its Triassic recovery. All previous studies have artificially inflated species richness by including names of organs from the same plant species. Species richness estimates for normalized taxa uses only one organ for each viable whole plant. The organ selected for normalization varies depending on the composition of individual fossil plant assemblages and requires critical evaluation. The process identifies from each systematic group the organ that and can be reliably identified and is the most diverse, selecting it as the most representative rather than other organs. Our approach includes omitting accounts of genera that lack species-level identifications (e.g., Tomiostrobus (=Annalepis) sp.) from assemblages in which one or more identified species of the same genus occurs (e.g., Tomiostrobus (=Annalepis) augusta, T. (A.) brevicystis). This assumes that the specimens identified as "sp." are likely poorly preserved or incomplete examples of named species. We have also omitted from species richness estimates fertile organs including gymnosperm seeds when other organs of the same plants are present in the same assemblage; in all cases where seeds are present, they co-occur with one or more species of gymnosperm leaf from which it is assumed that the seeds belonged to one or more of these plants. Lycopsid rootstock (Stigmaria), sporophylls (Lepidostrobophyllum) and cones (Lepidostrobus) are also omitted because, in all instances, these co-occur with stems (Lepidodendron) that are more distinctive and are typically identified to the species level (see Table 1). However, Mesozoic lycopods lack the leafy stems and branches of Paleozoic arborescent species, while their sporophylls tend to be well-preserved, systematically distinctive and diverse and so represent the best organ to measure richness. For Paleozoic sphenophytes we use their leaves (including species of

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Annularia) as they are systematically distinctive and the most diverse organ. However, leaves of Mesozoic sphenophytes are typically simpler, lack features to reliably distinguish species, and are of limited diversity so we use their stems and branches to measure richness as they are more readily distinguished from one another and more diverse. For "ferns" including members of the Marattiales, as well as gymnosperms including conifers, cycads and ginkgophytes, vegetative leaves have been used for normalization as they are reliably identified to species and genus level and in each case have the highest diversity. The only exceptions are rare instances where whole plants have been reconstructed in which, irrespective of whatever organ is selected for normalization, the name of the whole plant is used for that particular occurrence rather than the name of the isolated organ, following paleobotanical convention (see Bateman and Hilton, 2009). We accept that normalizing fossil plant data is, to some extent, subjective and cannot readily be tested for their robustness, but we consider these data to provide more realistic estimates of paleobotanical species richness than simple, raw-data based accounts. All data are presented in the supplementary dataset including information on normalization.

2.3. Fossil plant abundance

There is no robust method to quantitatively assess plant abundance in terrestrial settings (Cleal et al., 2021) unless fossil floras are preserved in-situ by obrution events such as volcanic ash-falls (e.g., Wang et al., 2012). In a broad sense, plant abundance may be indicated by a number of indirect measures including the number of locations that contain fossil plants as well as the paleobotanical richness in terms of numbers of specimens at each location, but these are subject to a variety of controls including collection intensity, spatial heterogeneity of plant distributions in contemporaneous settings, and a variety of physical (biostratinomic, sedimentary) and chemical taphonomic processes (Bateman, 1991; Allison and Bottjer, 2010). Here we focus on relative measure of abundance as well as using specific environmental proxies that provide crude insights into plant abundance in the environment in which they lived.

Firstly, having undertaken extensive fieldwork and collections-based

investigations on Permian–Triassic plant bearing sedimentary successions from South China, it is obvious that plant abundance varies considerably. In order not to overlook evidence for plants, we have considered all plant material, from fragmented plant debris to intact fossil plant organs because we are interested in assessing the presence of plants in the sedimentary system. As a crude measure of plant abundance, we use the relative descriptors absent, very rare, rare, common, abundant and very abundant to describe the amount of plant fragments encountered in each section in terms of (a) plant fossils on individual beds, and (b) the number of beds containing fossil plants.

Secondly, as coals represent accumulated peat, we consider coals to indicate high plant abundance for an extended time period. In contrast, the studied interval includes the Triassic "coal gap" (Retallack et al., 1996). While it is possible that the coal gap could represent widespread adverse preservation conditions for fossil plants (e.g., Vajda et al., 2020), in South China abundant waterlogged, fine grained paralic sedimentary facies occur in the early Triassic (Yu et al., 2008, 2010, 2015; Bercovici et al., 2015) that would have been suitable for preserving fossil plants if they were present in these settings. We therefore interpret the Triassic coal gap to indicate low plant abundance in paralic depositional sedimentary environments.

Thirdly, we consider environmental proxies related to soils on the basis that abundant vegetation cover is likely to bind soils together and diffuse water infiltration into the ground, thus having the net result of reducing surface water run-off (Zuazo and Pleguezuelo, 2009; Davies and Gibling, 2010). In contrast, bare, un-vegetated ground would be more susceptible to surface water run-off, physical weathering and erosion (Retallack, 2005; Algeo et al., 2011; Kaiho et al., 2016; Shen et al., 2015, 2022). We therefore use the presence of physical weathering in terrestrial settings as a rough proxy for plant abundance in terms of ground cover, highlighting the coevolution of plants and the environment. Finally, we consider Total Organic Carbon (TOC) levels in sediments as a crude proxy for terrestrial biomass and plant abundance with the caveat that it requires careful interpretation because of diagenetic controls on the value.

2.4. Evaluating plant ecology

The geological and paleoecological contexts in which fossil plants occur is important to their interpretation to provide links between fossil plants and the environment(s) in which they grew. It also allows greater understanding of the conditions in which extinction survivors lived, identifying features that may have contributed to their resilience as well as identifying the locations and environments of refugia.

Paleoecological assessments for plant taxa here come primarily from the available literature and is based on plant fossil anatomy and morphology and the sedimentary depositional environments in which they occur (Yao, 1978; Meyen, 1987; Bateman, 1991; Yang, 1993; Yang, 1994; Taylor et al., 2009). Fossil plant specimens were evaluated for their shape, size and completeness to elucidate their taphonomy in terms of being transported long, medium or short distances from their growth environment. For example, entire or almost entire organs lacking signs of taphonomic fragmentation, abrasion or size-sorting were interpreted as having undergone minimal transportation. This included in-situ plants and fossil plants found in paleosols, e.g., whole plants of *Lepacyclotes* (=*Annalepis*) in the Badong Formation (Meng, 1995). Greater levels of transport result in higher levels of fragmentation and size sorting and culminate in fine grained, well sorted plant debris beds such as layers of dispersed branches of *Neocalamites* in the Badong Formation (Meng, 1995). Depositional environments of the plant fossil were determined by sedimentary analysis and from the literature, as well as their paleogeographic location.

3. Fossil plant occurrences

3.1. Qixia Flora (Artinskian, Cisuralian)

This flora comes from the Liangshan Formation and other strata of the Qixia regional Stage in South China (Fig. 1), the age of which is roughly Artinskian (late Cisuralian, Shen S.Z. et al., 2019). Neither the Liangshan Formation nor its plant assemblage has received much study. This flora consists of a *Emplectopteris* triangularis - Taeniopteris multinervis assemblage but lacks gigantopterids,

representing the early stage of the Cathayian flora (Li et al., 1995). 323 324 325 3.2. Maokou Flora (Wordian–Capitanian) The Maokou Flora (Fig. 1) comes from the Maokou regional Stage in China, 326 327 the age of which is roughly Wordian to Capitanian in age (Li et al., 1995 and references therein). It mainly occurs in southeastern part of South China including 328 Fujian, Guangxi, Guangdong and Jiangsu provinces and seldomly occurs in eastern 329 330 parts of the Yangtze Platform. In the Maokouan (Capitanian Stage), the Cathaysian 331 (Gigantopteris) flora started to develop across South China. The dominant species are Gigantonoclea, Gigantopteris and Gigantopteridium, while Tingia, Asterophyllites 332 333 and Cordaites are common. 334 3.3. Longtan Flora (Wuchiapingian) 335 The Longtan flora (Fig. 1) from the Longtan Formation and lower Xuanwei 336 Formation is of Wuchiapingian age (Li et al., 1995). It belongs to the Cathaysian 337 338 (Gigantopteris) flora which steadily become more diverse and widespread during Wuchiapingian Stage (Li, 1997; Luo et al., 2021). In addition to southeastern parts of 339 340 South China, the Cathaysian (Gigantopteris) flora spread to the west of the Yangtze massif including the Xizang (Tibet) and Qinghai areas in the Wuchiapingian. 341 342 *3.4.* Xuanwei Flora (Changhsingian, latest Permian) 343 344 The Xuanwei flora occurs in the upper part of the Xuanwei Formation that crops out in western Guizhou and Eastern Yunnan provinces (Figs 1-3). It occurs in 345 346 siliclastic sediments and coals deposited in paralic settings (Li and Yao, 1980; Yu et al., 2015; Chu et al., 2016; Shen J. et al., 2019a). It rests stratigraphically on the 347 Emeishan Basalt (Fig. 2) and is dated to the late Changhsingian (He et al., 2017). In 348 the Chinahe section, in the Zhehai area (Fig. 2) ash bed zircons have yielded ages 349 252.4 ± 4.1 Ma and 252.30 ± 0.07 Ma (He et al., 2017; Shen S.Z. et al., 2011; Chu et 350 351 al., 2016). The Xuanwei Formation contains abundant fossil plants that represent the last occurrence of the Cathaysian tropical wetland flora (e.g., Shen G.L., 1995; Hilton 352

353	and Cleal, 2007; Feng et al., 2020). Of the 35 locations known to preserve fossil plant
354	assemblages in the Xuanwei Formation (Yu et al., 2015), the sections at Chahe
355	(Bercovici et al., 2015; Yu et al., 2015) and Chinahe (Fig. 3) are selected for study as
356	they are amongst the most continuous successions and have been studied in detail.
357	Fossil plant data from these sections provide statistical information on plant
358	distribution and evolutionary patterns through the Changhsingian, but information on
359	evolutionary rates also use data on all fossils from South China reported from this
360	time interval (see supplementary dataset).
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362	3.4.1. Chahe Section (Changhsingian, late Permian)
363	The Chahe section in Weining County, western Guizhou Province (Figs. 3, 4) has
364	been well studied (e.g., Yu et al., 2015; Chu et al., 2016). The section exposes the
365	Xuanwei Formation (Beds 1-70) and overlying Kaiyitou Formation (Beds 71-89) (Fig.
366	4) and is conformably overlain by the Dongchuan Formation. A zircon U-Pb date of
367	252.30±0.07 Ma from volcanic ash near the top of the Xuanwei Formation indicates a
368	latest Permian age (Shen S.Z. et al., 2011), suggesting the Permian-Triassic boundary
369	(PTB) is in the basal Kayitou Formation. Two layers of volcanic ash, with an
370	intervening layer of black mudstone, provide a common lithostratigraphic marker for
371	a level immediately below the Permo-Triassic Boundary in non-marine sections (Chu
372	et al., 2016).
373	The Xuanwei Formation at the Chahe section contains numerous plant fossils
374	including the Cathaysian wetland genera Gigantopteris, Lepdiodendron, Pecopteris,
375	and Fascipteris. Plant fossils are abundant in the 31 plant-bearing layers (Fig. 4)
376	before disappearing at Bed 69 in the latest Permian, with only <i>Peltaspermum</i> sp.
377	persisting in Bed 70 above the EPPC level. Fossil plants are absent in the overlying
378	Kayitou Formation in the Chahe section (Fig. 4).
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380	3.4.2. Chinahe Section (Changhsingian, late Permian)
381	The Chinahe section of eastern Yunnan Province (Fig. 3) starts with the Emeishan
382	Basalt (Bed 0) and is unconformably overlain by terrestrial facies of the Xuanwei

Formation (Beds 1–25). Paralic gray-blue-green mudstone facies of the Kayitou Formation (Bed 26) conformably overly the Xuanwei Formation (Fig. 5) which in 384 turn is overlain by the purple red Dongchuan Formation (Wignall et al., 2020). 385 The Xuanwei Formation in the Chinahe section commences with a thick layer of 386 gray-black mudstone (Bed 1), that contains large amounts of well-preserved plant 387 388 fossils (Fig. 5, 6). In Bed 1, Lobatannularia, Pecopteris and Gigantopteris are common, but the flora in this bed is diverse and also contains a range of sphenopsids 389 390 (Lobatannularia cathaysiana, L. heianensis, Paracalamites stenocostatus and 391 Schizoneura amnchuriensis), marattialean ferns (Pecopteris (Asterotheca) guizhouensis, P. (A.) orientalis, P. (A.) hemotelioides, P. sahnii, Fascipteris sinensis 392 and F. hallei), gigantopterids (Gigantonoclea guizhouensis, G. rosulata, Gigantopteris 393 394 dictyophylloides and Gigantopteris nicotianaefolia), ferns (Cladophlebis permica, C. ozakii) and occasional gymnosperm leaves (Neuropteridium sp., Peltaspermum sp., 395 Taeniopteris multinervis, Rhipidopsis panii). Above Bed 1, the lithology changes into 396 gray yellow or gray green, thin-bedded muddy siltstone and thin coals, pale gray or 397 398 gray blue, thin-bedded mudstone and yellow gray fine-grained sandstone (Bed 2–25). Thin-bedded mudstones developed above coals (in Beds 3, 12, 16, 25) contain some 399 400 plants fragments that are not identifiable to species level, but include fragments of Lepidostrobophyllum, gigantopterids, Compsopteris, Pecopteris and Taeniopteris. 401 402 Until Bed 25 there are more gray-black, medium bedded, siltstones mixed with shaly coals as well as gray yellow or green silty-mudstone. Above Bed 25, there are two 403 404 gray, thin-bedded mudstones together with a black mudstone layer, which may be correlative with the sandwich-like lithologies seen in the Chahe section at the 405 406 boundary of the Xuanwei and Kaiyitou formations. Above this "sandwich-like" 407 mudstone layer, a gray-yellow, thin-bedded sandstone and a layer of black siltstone contains numerous of fragmentary specimens of Peltaspermum, Lepidopteris, 408 Pecopteris and Giantopteridium. In the boundary of the Xuanwei and Kayitou 409 formations (Bed 26), there is a monotypic layer of dispersed *Tomiostrobus* 410 411 (=Annalepis) in the dark gray, thin-bedded siltstone (in the bottom of bed 26). After the layer of *Tomiostrobus* (=Annalepis) comes the blue to greenish blue, thick-bedded 412

siltstones of the Kavitou Formation, which contain no plants but abundant 413 414 conchostraca and a few horizons of marine bivalves (Fig. 6). 415 3.5. Dalong Flora (Changhsingian, late Permian) 416 417 The Dalong Formation formed in the western part of the Yangtze shallow sea and 418 is laterally equivalent to the Heshan Formation in eastern areas (Figs. 2). Fossil plants 419 from the Dalong Formation were documented by Liu et al. (2007), Song et al. (2013, 420 2015) and Li et al. (2019). The marine formation consists of thin-bedded cherts, siltstones, and clays (Shen J. et al., 2012a, 2013; Fig. 7). The chert beds contain both 421 plant fossils and a marine biota including radiolarian, brachiopods, bivalves, 422 conodonts, and foraminifera (Fig. 7) that enables correlation with the PTB GSSP at 423 424 Meishan (Yin et al., 2007; Shen J. et al., 2012a; Li et al., 2019). Based on conodont 425 occurrences, the Dalong Formation has been dated to the Changhsingian (Li et al., 2019). Although the Dalong and Heshan formations are of subtly different age (Figs. 426 1, 2), they share similar plant fossils and so are discussed together here. From them, 427 428 five locations contain plant fossils. 429 From the Dalong Formation we studied the plant fossils from three published sections: Xinmin, Duanshan A and Duanshan B sections (Fig. 7). In general, the 430 gymnosperms are well-preserved while the typical Cathaysian floral elements are 431 more fragmentary and abraded suggesting considerable transport into the depositional 432 setting. Xinmin is the longest section and, based on the presence of the conodont 433 434 Hindeodus parvus, is the only section where the Permian–Triassic boundary (PTB) is 435 well defined (Zhang et al., 2014). In the following account, the Xinmin section is illustrated as the main section, with the Duanshan A and B sections providing 436 437 additional data. 438 3.5.1. Xinmin Section 439 440 The Xinmin section in Jiaozishan town, Anshun City, southern Guizhou 441 Province is 105 km away from the Duanshan A and B sections of the Dalong Formation (Fig. 3). Lithologies in the Xinmin section (Fig. 7) comprise thin-bedded 442

443	chert and carbonates including micritic limestones, with siltstone interlayers and
444	occasional thin, gray-green, volcanic tuff beds (Beds 1-6, Shen J. et al., 2012b; 2021).
445	Plant fossils mainly come from the black, thin-bedded mudstone interlayers in the
446	middle of Bed 2 and the top of Bed 4. These comprise large conifer branches
447	including secondary or tertiary branches that can reach up to 50 cm long and
448	Taeniopteris leaves with good cuticle, amongst smaller, fragmentary fossils of
449	Lepidostrobophyllum, Paracalamites, Pecopteris, Gigantopteris, Cordaites and
450	Sphenobaiera (Li et al., 2019). The conifers with well-preserved cuticles have been
451	identified as Anshuncladus xinminensis, A. contiguous, A. aduncatus, Pseudoullmania
452	frumentarioides and Szecladia multinervis. Li et al. (2019) interpreted the conifers to
453	be preserved very close to where they grew, inhabiting coastal habitats, due to their
454	completeness. These conifers are only preserved in the terrestrial-marine interbedded
455	facies of the Dalong and Heshan formations and are absent in terrestrial facies rich in
456	fossil plants (Liu et al., 2007, 2013; Li et al., 2019). In contrast, the fragmentary
457	Gigantopteris and Pecopteris leaves in the Dalong Flora were likely transported
458	greater distances.
459	Based on the presence of the conodont Hindeodus parvus in the Dalong
460	Formation, the PTB is placed in the middle of the limestone at the bottom of Bed 5.
461	Beds 1 to 6 contain a Permian-Triassic fauna (Fig. 7) including bivalves
462	(Hunanopecten exilis - H. qujiangensis in the Permian and Claraia liuqiaoensis in the
463	Triassic), ammonites (Pseudotirolites - Sinoceltites of the Permian and Triassic
464	Xenaspis), and conodonts (Clarkina changxingensis, C. yini Permian and C.
465	meishanensis, Hindeodus changxingensis, H. parvus Triassic assemblages) (Zhang et
466	al., 2014; Yang, 2015; Li et al., 2019). The biostratigraphic evidence shows that this
467	section is complete (Li et al., 2019).
468	
469	3.5.2. Duanshan A Section
470	The Duanshan A section is in the same town as the Xinmin section in Huishui
471	County, Guizhou province (Fig. 3), and the two sections share similar plant as well as
472	marine faunal records. The section is lithologically divided into eight Beds (Fig. 7)

that comprise gray, siliceous mudstone, gray blue, thin-bedded micritic limestone, with gray black or gray—yellow mudstone interlayers. This is quite similar to the succession in the Xinmin section, except that Bed 3 is composed of gray yellow mudstone and white volcanic ash. Beds 1–2 and 4–6 contain the *Hunanopecten exilis* - *H. qujiangensis* bivalve assemblage, the *Pseudotirolites* - *Sinoceltites* ammonite assemblage, and the *Albaillella triangularis* - *A. yaoi* radiolarida assemblages (Fig. 7) and allow us to correlate Beds 3–8 in this section with the middle of Bed 3 to the top of Bed 4 in the Xinmin section (Fig. 7).

3.5.3. Duanshan B Section

The Duanshan B section is on the opposite side of the road to the Duanshan A section (Fig. 3) and is much shorter. It contains more grayish yellow mudstone and pelitic siltstone than Duanshan A and shares a similar plant fossil record (Fig. 7). According to the lithological successions and the location of the Duanshan A and B sections, Beds 1–6 of Duanshan B corresponds to Bed 3 in Duanshan A (Fig. 7).

3.6. Kayitou Flora (Griesbachian interval, Permian–Triassic transition)

The Kayitou Formation conformably overlies the Xuanwei Formation and represents paralic facies (Fig. 1). According to the plant fossils and conchostracan biostratigraphy and other criteria, the PTB occurs near the base of the Kayitou Formation (Yu et al., 2010; Chu et al., 2016; Wignall et al., 2020). The Kayitou Flora occurs in the bottom of the Kayitou Formation, but its exact age is contentious (Chen et al., 2011). The widespread appearance in south China of the lycopod *Tomiostrobus* (=*Annalepis*) has been suggested to coincide with the end of the EPPC and the beginning of Triassic (Yu et al., 2010, 2015). Six sections contain the Kayitou Flora in South China and have similar plant fossil compositions (Fig. 8). To eliminate the influence of diachroneity, representative sections containing datable marine biota and typical plant fossils are selected in this study: the Chinahe, Tucheng and Mide sections. In general, these sections contain limited occurrences of plant fossils, typically within single beds and abundance is low. This is noticeably different from

503 plant fossil occurrences prior to the EPPC. 504 505 3.6.1. Chinahe Section The Chinahe section in Xuanwei City, eastern Yunnan Province (Figs. 8, 9) 506 contains terrestrial facies of the Xuanwei Formation (Beds 1-25) and marine facies of 507 the Kayitou Formation (Bed 26–29). At Chinahe, the EPPC is defined at the level of 508 509 disappearance of the Xuanwei Flora together with the last coal line in Bed 25 and the 510 appearance of *Tomiostrobus* (=Annalepis) (Fig. 9) between Beds 25 and 26 at the lithologic boundary between the Xuanwei and Kayitou formations. The last coal in 511 512 the Xuanwei Formation marks the top of the Xuanwei Formation at Chinahe. Bed 26 at the base of the Kayitou Formation contains *Tomiostrobus* (=Annalepis) augusta, T. 513 (A.) brevicystis, T. (A.) latiloba, T. (A.) zeilleri, T. (A.) spp., Sphenopteris tenuis, 514 515 Fascipteris stena, Peltaspermum martinisii, and fragmentary remains of Gigantopteris spp. and Pecopteris (Asterotheca) orientalis that extend their range 516 from Beds 21 and 24 respectively (Fig. 9). Above this level, plant fossils are 517 518 extremely rare and limited to fragments of *Tomiostrobus* (=Annalepis) in Bed 27, which occur above the first occurrence of the bivalve Pteria variabilis at this location. 519 This bivalve also occurs above the EPPC boundary in the Tucheng and Mide sections 520 (Figs 8, 9). According to the presence of the *Pteria variabilis-Promyalina schamarae* 521 bivalve assemblage in Beds 26–28 (Song T. et al., 2018), the age of plant assemblage 522 in Beds 26–27 from the Chinahe section is considered to be Griesbachian. 523 524 525 3.6.2. Tucheng Section 526 The Tucheng section in Panxian County, western Guizhou Province (Fig. 8) exposes a thick coal (Bed 16) of the Xuanwei Formation, and the overlying Kayitou 527 Formation (Beds 17-23) (Fig. 9; Yu et al., 2015; Broutin et al., 2020). Beds 17-20 are 528 gray-yellow, thin-bedded mudstone except for Bed 19, a gray-yellow sandstone. 529 Fossil plants occur in the middle of Beds 17 and 18, and include Lepidodendron sp., 530 Tomiostrobus (=Annalepis) zeilleri, T. (A.) brevicystis, Pecopteris sp. and 531

Peltaspermum martini. The upper part of Bed 21 is gray-green, medium-bedded

533	siltstone interbedded with thin beds of mudstone. Bed 22 is a gray-brown, medium-
534	bedded siltstone, while Bed 23 changes into gray green siltstone and contains
535	Sphenopteris sp., Peltaspermum mattenii, P. lobutalum, P. sp., Pecopteris sp.,
536	Gigantonoclea guizhouensis, Gigantonoclea sp., Gigantopteris dictyophylloides and
537	Gigantopteris sp. (Fig. 9).
538	Beds 17–23 at Tucheng contain marine fossils including the bivalves <i>Unionites</i>
539	fassaensis, U. canalensis, U. sp., Leviconcha orbicularis, L. praeorbicularis, Pteria
540	ussurica variabilis and P. murchisoni leshanensis (Fig. 9). Correlation with other
541	sections (Fig. 9), places the termination of the EPPC in the lower middle of Bed 17,
542	below the first appearance of <i>Tomiostrobus</i> (= <i>Annalepis</i>).
543	
544	3.6.3. Mide Section
545	The Mide section in Xuanwei City, eastern Yunnan Province (Fig. 8) is
546	composed of thin-bedded, gray green siltstone (Beds 15-17 and 24-26), while Bed 18
547	is a thin, white volcanic ash (Fig. 9; Chen et al., 2011; Bercovici et al., 2015). Bed 19
548	is a yellowish green sandstone and Bed 20 a gray-yellow siltstone. Beds 21-23 are
549	pale yellow to gray-brown siltstone. Bed 16 contains Paracalamites stenocoastatus,
550	Gigantonoclea guizhouensis, Gigantopteris sp. Beds 20 and 21 contain Tomiostrobus
551	(=Annalepis) zeilleri, T. (A) brevicystis, Peltaspermum martensii, P. sp., Pecopteris sp.
552	and lycopsid roots of Stigmaria. Marine biota is abundant in Beds 18–25 (Fig. 9) and
553	includes the bivalves Unionites fassaensis, U. canalensis, U. sp., Leviconcha
554	orbicularia and L. praeorbicularis, the ammonite Ophiceras sp. and the ostracod
555	Langdaia suboblonga (Forel et al., 2020). The EPPC termination level occurs at the
556	beginning of Bed 19 prior to the appearance of <i>Tomiostrobus</i> (=Annalepis) (Fig. 9).
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558	3.7. Feixianguan Flora (Dienerian or Smithian, Early Triassic)
559	The lower part of Feixianguan Formation (Fig. 1) is correlative with the Triassic
560	portion of the Kayitou Formation (Tong et al., 2019) and conformably overlies the
561	Xuanwei Formation (Figs. 1, 2). Fossil plants occur at the top of the Feixianguan
562	Formation one metre below the boundary with the overlying Yongningzhen Formation

(Fig. 10). Bivalve data indicate that the age of the fossil plant assemblage in the Feixianguan Formation is Dienerian or early Smithian (Yin et al., 1985; Gou et al., 1996; Tong et al., 2019). In South China, only the Feixianguan Formation in the Pojiao, Lubei sections and the Dongchuan Formation in one section near Lubei Village (Feng et al., 2018) contain plant fossils of this age and all yield the same flora. Due to the similar age and floristic composition of the sections of the Dongchuan and Feixianguan formations, we combine them as one Feixianguan Flora and focus on describing the Feixianguan sections as much less is known about the flora of the Dongchuan Formation. In general, the Pojiao and Lubei sections contain limited plant fossil layers, and each layer yields a few fossil plant fragments. The sedimentary facies of the Feixianguan Formation comprise interacting terrestrial and marine facies according to the lithological succession and marine bivalve fossil yielded in the lower beds below the plant fossil horizons.

3.7.1. Pojiao Section

The Pojiao section is located in Pojiao village, Huize County, northeastern Yunnan Province (Fig. 8). It exposes the Feixianguan Formation (Beds 1–9) that comprises cyclic beds of reddish purple and gray black sandstone (Fig. 10), and the overlying Yongningzheng Formation (Beds 10–11) that gradually change from gray green thick-bedded calcareous sandstone (Bed 10), into dark gray limestone (Bed 11). In the Feixinguan Formation, fossil plants occur in at the top of Bed 9 in two layers of thin-bedded siltstone interbedded with gray green thick sandstone. These comprise *Phyllotheca* sp., *Neocalamites* sp., *Equisites* sp., *Cladophlebis* sp., *Todites* sp., *Voltzia heterophylla*, *Peltaspermum* sp., *Baiera* sp. and *Taeniopteris* sp. (Fig. 11). The Bed 9 plants are abraded preventing species-level identifications with the exception of *Peltaspermum* and *Voltzia* from which an almost fertile shoot is known. In this flora there are abundant gymnosperm seeds, but these are not listed in the range diagram (Fig. 10) as they most likely represent the seeds of one (or more) of the other gymnosperms in the flora (i.e., *Voltzia*, *Peltaspermum*, *Baiera*, *Taeniopteris*). The bottom and middle parts of the Feixianguan Formation at Pojiao are mostly marine

facies and contain two bivalve assemblages with the lower *Claraia wangi* assemblage

594 typical of the early Induan, and the upper Eumorphotis multiformis - Claraia aurita -

Claraia stachei assemblage typical of middle-late Induan of Feixianguan Formation

596 (Gou et al., 1996; Tong et al., 2019).

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3.7.2. Lubei Section

The Lubei section in Lubei village, Huize County, northeastern Yunnan Province

600 (Xu et al., 2017; Shen J. et al., 2019b) is close to Pojiao (Fig. 8). It exposes the late

Permian Xuanwei Formation (Beds 1–3), the early Triassic Feixianguan Formation

602 (Beds 4–8) and later Early Triassic Yongningzhen Formation (Beds 9–13) (Fig. 10).

The Feixianguan Formation comprises purple red to gray green, thin to medium-

bedded lithic sandstone and siltstone (Xu et al., 2017; Shen J. et al., 2019b). Fossil

plants occur in Beds 5–6 and comprise *Phyllotheca* sp., *Neocalamites* sp., *Equisites*

acanthodon, Equisites sp., Cladophlebis sp., Peltaspermum sp., Taeniopteris sp. and

607 *Voltzia* sp. (Fig. 11).

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3.8. Lingwen Flora (Spathian, Early Triassic)

The Linguen section in Hainan Province (Figs. 1, 8) was previously documented

by Zhou et al. (1979) and Zhang et al. (1992), but the flora has not been investigated

subsequently. Only one section contains the Lingwen Flora in South China. The

613 Lingwen Flora is considered to have been deposited during the Olenekian based on

plants and palynomorph assemblages that are similar to those from the French

Buntsandstein (Gall and Grauvogel-Stamm, 2005). The flora at Lingwen contains

elements including *Pleuromeia*? sp., *Todites shensiensis*, *Asterotheca szeiana*,

617 Ctenozamites cycadea, Leuthardtia ovalis, Vittaeohyllum sp., Albertia, Voltiza and

Pelourdea (=Yuccites) (See full fossil list in supplementary dataset; Fig. 12) that are

also common in the lower part of the Badong Flora (see below). The sediments of the

Lingwen Formation comprise terrestrial fluvio-lacustrine facies (Zhou et al., 1979;

621 Zhang et al., 1992).

- 3.9. Badong Flora (Anisian, Middle Triassic)
- Plant fossils in the Badong Formation (Fig. 1) were first reported by Ye et al.
- 625 (1979) and systematically investigated by Meng et al. (1993, 1995, 1996, 1998). The
- Badong Formation has a widespread distribution across the Upper and Middle
- Yangtze area in South China (Fig. 13). It conformably overlies the Jialingjiang
- Formation (Figs. 1, 14) and comprises five members, but in most areas only three or
- four are present due to erosion during Ladinian regression. In Hunan Province, the
- Sangzhi County area yields the longest stratigraphic sections especially around the
- village of Hongjiaguan (Fig. 13) where an almost continuous section occurs, although
- 632 it contains few plant fossils. In contrast, the nearby section in Furonggiao Village
- contains abundant plants but is less continuous. These two sections are correlated with
- each other and included as the combined Hongjiaguan and Furongqiao section (Fig.
- 635 14).

- 636 Lithologically the Badong Formation comprises pale gray, yellow to gray green
- calcareous mudstone to siltstone in Member 1 above a gypsum-karst breccia at the top
- of Jialinjiang Formation, and purple red, thick-bedded siltstone to sandstone with
- interbedded blue mudstone to siltstone in Member 2 (Fig. 14). Member 3 comprises
- gray-yellow calcareous mudstone to siltstone and limestone, and Member 4 purple red,
- thick-bedded siltstone to sandstone, while Member 5 comprises gray blue to yellow,
- thin-bedded siltstone and gray black, thin-bedded limestone to calcareous siltstone. In
- the Hongjiaguan and Furongqiao section, only members 1 and 2 are present (Fig. 14).
- The boundary between members 4 and 5 contains the first coal in South China after
- the PTB, and thus represents the end of the "coal gap" (Meng et al., 1995; Retallack et
- al., 1996). The sedimentary facies of the Badong Formation are coastal or tidal in
- members 1, 2, 4 and 5 and the abundant marine biota preserved together with in-situ
- 648 'mangrove-like' plants of Lepacyclotes (=Annalepis) and Pleuromeia (Meng et al.,
- 649 1995).
- Fossil plants in the Badong Formation mostly come from the boundary of
- members 1 and 2 (Fig. 14), although less common and more poorly preserved fossils
- also occur in green blue interlayers in members 2 and 4. We found eight plant fossil

653	locations at Hongjiaguan with four locations combined as one, and four in Furongqiao
654	Village (Fig. 15). The flora comprises $Lepacyclotes$ (= $Annalepis$) $brevicystis$, $L(A)$.
655	zeilleri, L(A). $sangzhiensis, L(A)$. $angusta, Pleuromeia sanxiaensis, P. marginulata, P.$
656	hunanensis, Equisites gracilis, Neocalamites shanxiensis, Todites shensiensis,
657	Peltaspermum multicostatum, P. miracarinatum, Yuccites vogesiacus, Y. anastomosis,
658	Yuccites sp., Voltzia heterophylla, V. curtifolia, V. sp., Willsiostrobus cordiformis,
659	Cardiocarpus triquestrus, and C. sp. (Fig. 15).
660	In Member 1 of the Badong Formation, typical Anisian bivalves of the
661	Leptochondria - Myophoria goldfussi mansuyi assemblage are preserved in siltstones-
662	mudstones, whilst Member 3 contains the bivalves Plagiostoma sp., Placunopsis sp.,
663	the conodonts Neospathodus sp. and Enantiognathus sp., and the ammonoid
664	Progonoceratites sp. that indicates an Anisian age (Meng et a. 1995).
665	
666	3.10. Carnian and Norian (Daqiodi and Shazhenxi) flora
667	Carnian and Norian floras are quite common in South China although the
668	boundary between these stages is poorly constrained. The Carnian/Norian floras are
669	known from the Daqiaodi Formation at Yongren on the border between Yunnan and
670	Sichuan Provinces, the Jiuligang Formation at Yuan'an in western Hubei Province,
671	and the Japeila Formation in eastern Xizang Province (Tibet), collectively from over
672	24 locations in South China; Xujiahe Formation at Guangyuan in Sichuan Province,
673	the Dajing Formation in the border area of Sichuan and Yunnan provinces, the
674	Anyuan Formation in Hunan and Jiangxi provinces, the Bagong Formation in
675	southern Fujian Province and the Malugou Formation at Tianqiaoling in Jilin province
676	(Xu et al., 1979; Li et al., 1995; Liu et al., 2009). The age of this flora is determined
677	by marine biostratigraphy (Li et al., 1995). The lower part of the flora comprises the
678	Abropteris - Pterophyllum longifolium Assemblage that includes Equisetites
679	arenaceus, Abropteris cottonii, Mixopteris intercaearis, Pterophyllum longifolium, P.
680	jaegeri, Angiopteris antiqua, Sagenopteris glossopteroides, Danaeopsis marantacea,
681	Ctenozamites chinensis, Stenopteris bifurcata and Ctenozamites chinensis and
682	resembles the early Late Triassic Lettenkohle Flora in western Europe (Li et al.,

683	1995). The upper part of the flora comprises the <i>Dictyophyllum - Drepanozamites</i> or
684	Dictyophyllum - Cycadocarpidium Assemblages, including Dictyophyllum nathorstii,
685	Clathropteris elegans, Reteophlebis simplex, Drepanozamites nilssonii,
686	Doratophyllum hsuchiahoense, Anomozamites loczyi, Podozamites (Cycadocarpidium)
687	gigantean, Cycadocarpidium swabii and Hausmannia ussuriensis, and thus
688	considered to be Norian (Li et al., 1995), comparable to the Nariwa Flora of Japan.
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690	3.11. Rhaetian (Yangbaichong) flora
691	The Rhaetian Flora (Fig. 1) is represented by the plant assemblage from the
692	Yangbaichong Formation at Hengyang in South Hunan province and comprises the
693	Ptilozamites - Anthrophyopsis Assemblage. The plant assemblage in the Anyuan
694	Formation probably belongs to this flora as well and comprises Ptilozamites chinensis,
695	Anthrophyopsis leeiana, Clathropteris meniscioides, Todites crenatus, Nilssoniopteris
696	oligotricha, N. xuiana, Pterophyllum ptilum, Podozamites distans, Cycadocarpidium
697	erdmannii and Stalagma samara (Li et al., 1995). It is comparable to the Lepidopteris
698	zone floras from eastern Greenland and Germany (Zhou et al., 1989), and is
699	considered to be of Rhaetian age (Li et al., 1995).
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702	4. Permo-Triassic vegetation change in South China
703	Artinskian (Middle Permian) to Rhaetian (Late Triassic) macro and micro-floral
704	stratigraphic range data have been compiled to determine the complete range of plant
705	taxa present during the End Permian to Middle Triassic before normalization (Figures
706	18, 19). Then normalized data from the fossil plant ranges in South China from the
707	Late Permian to the early Late Triassic are analyzed to show species richness and
708	origination and extinction taxa number for the entire flora (Fig. 18), and origination,
709	extinction rates of the entire flora together with the rates for individual plant group
710	(Fig. 19). These data show an increase in total species richness from the Artinskian
711	(late Cisuralian) and peak species richness in the Wuchiapingian, followed by a
712	decrease culminating in a diversity lowpoint either side of the PTB (Fig. 18). There is

no evidence for a floral crisis in the Capitanian although this may be because our data is compiled at the stage level, whilst in the marine realm extinction many losses were intra-Captanian (Bond et al., 2010). The Lower Triassic saw low species richness (typically <20 species), with values only increasing during the Carnian and Norian (Upper Triassic) before declining again in the Rhaetian.

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The plant diversity decline started from the lower Changhsingian with only species richness decline and flora alternation without apparent vegetation abundance reduction, characterized by the decline of gigantopterid flora in South China (Yu et al., 2015). The termination of the EPPC is marked by both these phenomena including the notable and abrupt drop of species together with cessation of coal formation, marking the collapse of terrestrial ecosystems (Fig. 22). The low-diversity survivors in the Kayitou Formation represents a holdover subset of the Cathaysian Flora including Lepidodendron, Paracalamites, Pecopteris, Sphenopteris, Fascipteris, Gigantopteris and Gigantonoclea, mixed with the opportunistic lycopod Tomiostrobus (=Annalepis). This floral assemblage is only present in the basal-most part of the Kayitou formation suggesting only a short survival interval. In this regard, they are closely comparable with the numerous, short-ranging holdover taxa in the marine record which thrived between two extinction pulses in the early Griesbachian (Song et al., 2012). Previous analyses of species richness within Paleozoic floras suggest Pennsylvanian wetlands of Europe (Cleal et al., 2012) had local standing diversity typically in the region of 40-60 species, but with regular species originations and extinctions (Cleal and Thomas, 2004; Cleal et al., 2012). From our analyses, species richness has been calculated using data compiled in stage-level time bins and from the pre-EPPC Xuanwei Flora values and is more comparable to Pennsylvanian regional-scale richness of the Variscan Foreland with standing diversity of >70 species (Cleal et al., 2012). Although not from the Permian, these data suggest a standing diversity of the pre-EPPC Longtan and Xuanwei floras was comparable to Euramerican Carboniferous peat forming communities, but the early Triassic diversity is significantly below this level (Fig. 18) and more comparable to diversity of the Devonian Rhyniophytic and Eophytic evolutionary floras (Cleal and Cascales-Miñana, 743 2014).

The species origination rate began to exceed the extinction rate in the Early
Triassic Feixianguan Formation, representing the earliest stage of the species richness
recovery following the EPPC. This interval is characterized by plant populations with
low abundances and an absence of coals. The first post-EPPC coals in South China
are in the Anisian, and their reappearance coincides with that of terrestrial herbivores
(*Lotosaurus*) in Member 2 of the Badong Formation (Meng et al., 1995; Hagen et al.,
2018). In the Upper Triassic, more floras are reported (Li et al., 1995) demonstrating
the continued diversification in humid conditions. Peak Triassic species richness
occurs in the Carnian and Norian when diversity attained a comparable level to that of
the Late Permian (Fig. 18).

Overall, late Permian to middle Triassic plant evolution patterns can be resolved

Overall, late Permian to middle Triassic plant evolution patterns can be resolved into four distinct phases according to the macro plant fossil data, palynology data, biomaker and wildfire proxies (Fig. 22). Phase 1 occurred during the Changhsingian and terminated at the end of the EPPC. Phase 2 consists of the holdover Permian flora found in the Permian–Triassic transitional Kayitou Formation, including the Chinahe, Tucheng and Mide sections (Changhsingian–Griesbachian interval). Phase 3 represents the recovery of species richness during the late Induan in the upper part of the Feixianguan section. Finally, Phase 4 represents substantial recovery after the Olenekian, as seen in floras of the Lingwen Formation at Lingwen and the Badong Formation, including the Hongjiaguan and Furongqiao sections.

4.1. Phase 1: Pre-extinction floras and the EPPC

The Changhsingian flora at Chahe (Fig. 3) and Chinahe (Fig. 4) has a typical Cathaysian character. The fragmented plant assemblage shows considerable turnover of short-ranging taxa and a final, rapid loss of almost all taxa, including long-ranging taxa, at the end of the phase. Plant taxa ranges vary between the sections. The minor losses during the initial episode of the EPPC can be considered a turnover of short-ranging taxa while the disappearance of long-ranging taxa marks a severe crisis. For example, the long-ranging elements both in the Chahe and Chinahe sections including

Lobatannularia multifolia, Pecopteris orientalis, P. arcuata, P. gracilenta, P. taiyuanensis, Rajahia guizhouensis, Fascipteris sinensis, Compsopteris contracta, C. punctinervis, Gigantopteris nicotianaefolia, Gigantonoclea largrelii, Gigantopteris dictyophylloides and Neuropteridium were common in every layer bearing fossil plants, until their disappearance at the end of the EPPC marked a dramatic ecological crisis.

Floral assemblages from the Chahe and Chinahe sections lack conifer macrofossil remains. This phenomenon is quite common in South China with Guadalupian—Cisuralian floras lacking conifers in terrestrial or terrestrial-marine sections. Conifers are only found in the fully marine facies of the Dalong Formation which yields both conifers and the cycadophyte *Taeniopteris* with well-preserved cuticles (Li et al., 2019). This pattern is in marked contrast to contemporaneous floras from Europe and North China where conifers occur in terrestrial and terrestrial-marine facies (Wang et al., 1985, 1996). According to taphonomic features of the fossils and the sedimentology, the conifers in the Dalong Formation are likely to have lived in coastal settings or on islands. Species of conifer in the Dalong Formation all disappear before or during the EPPC and do not reappear in the Triassic. Mesozoic-type conifers *Voltiza* and *Albertia* appear in South China after the EPPC. Due to lack of understanding of the evolutionary relationships within Paleozoic and Mesozoic conifers as whole plants, it is unknown whether stratigraphically younger Triassic conifers are closely related to the Late Permian conifers from South China.

The abrupt abundance reduction of the terminal EPPC is caused by elevated extinction and declining origination rates that are seen in the entire flora and amongst each plant group (Fig. 19). Most of the plants from Cathaysian floral communities disappeared during this event, including Paleozoic lycopods, noeggerathialean progymnosperms and cordaitalean coniferophytes, leaving only a few survivors. Five sections from different sedimentary facies allow the details of the plant evolution pattern to be deciphered: terrestrial facies from the Chahe section in the Xuanwei Formation, terrestrial-marine transitional facies from the Chinahe section in the Xuanwei Formation, and marine facies from the Xinmin and Duanshan A and B

sections in the Dalong Formation.

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The EPPC record is that of coal-swamp flora rather than upland taxa (Yu et al., 2015). Xeric upland taxa took over the empty space after the extinction of coalswamp taxa a trend that began slightly before the terminal crisis, as shown by the gradually increasing proportion of pollen before the EPPC (Yu et al., 2008). Moreover, the palynological record from the Changhsingian paralic Kayitou Formation in Yunnan Province (Ouyang, 1991) is similar to the marine Yinkeng Formation in Meishan section of Zhejiang Province, and other sections in South China (Zhang et al., 2004). There were few fungal/algal spores before the lithological boundary of the Permian and Triassic. The palynological record from before the crisis in terrestrial locations in Xinjiang differs from that in South China, being mainly composed of gymnosperm pollen, thereby showing an earlier transition from Paleophytic to Mesophytic flora in this northern area (Qu et al., 1986) (Fig. 22). The crisis in the Guodikeng Formation in Xinjiang was coupled with an abundance of lycopod spores (Qu et al., 1986; Chu et al., 2015). Black carbon (BC) content and biomarkers for combustion process such as polynuclear aromatic hydrocarbons (PAHs) are useful tools for indicating wildfires (Shen W.J. et al., 2011, 2012; Xie et al., 2007; Chu et al., 2020). In the paralic Chinahe section, charcoal content peaks in the upper part of Bed 25 to Bed 26 with a similar peak also seen in the coeval beds 23–24 at Meishan (Fig. 20), although PAHs peak later in Bed 26 in Meishan (Xie et al., 2007; Shen W.J. et al., 2011) suggesting there was still fuel for combustion, after the main plant crisis, albeit for a short period of time (Shen W.J. et al., 2011).

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4.2. Phase 2: Griesbachian interval extinction and survival

The Griesbachian plant assemblage from the Kayitou Formation is characterized by pioneering or opportunistic taxa comprising Mesozoic-type lycopods *Tomiostrobus* (=Annalepis), mixed with holdover taxa including lycopods (Lepidodendron), sphenophytes (Paracalamites), ferns (Pecopteris) and gigantopterids (Gigantopteridium). This composition is mirrored in palynological data and marked

as a distinct, Permian-Triassic transitional flora (Chen et al., 2011; Yu et al., 2015). 833 The palynological record in the Dalongkou section in Xingjiang (NW China) also 834 comprises a transitional assemblage, with a mix of Permian and Triassic species, but it 835 differs from the South China record in having a higher percentage of gymnosperm 836 pollen (Qu et al., 1986; Ouyang, 1991, 2007). Although holdover elements persist into 837 the Kayitou Formation, they rarely range more than 30 m above the base in all 838 sections, whilst most disappear within 5 m. The palynology record confirms the short 839 840 duration of survival of the holdover elements, whilst the pioneers persisted for much longer into the Late Triassic (Ouyang, 1991; Grauvogel-Stamm and Ash., 2005; Yu et 841 al., 2008, 2010). The survival elements are restricted to refuges, for instance swamp 842 facies preserved in the paralic Kayitou Formation (Li et al., 1995; Grauvogel-Stamm 843 and Ash., 2005; Yu et al., 2008). 844 Post EPPC opportunistic plant species abruptly occupied empty niches but 845 disappeared shortly afterwards following the initial stages of ecosystem recovery. In 846 addition to holdover and opportunist taxa, the early Induan flora also includes the 847 848 surviving seed fern *Peltaspermum* that rapidly recovered after the EPPC in terms of both its species richness and abundance, together with conifers that gradually radiated 849 (Whittaker & Goodman, 1979; Huston and Smith, 1987; Glenn-Lewin et al., 1992; 850 Ren et al., 2001). The success of *Peltaspermum* and conifers marks forest ecosystem's 851 re-establishment and shows spatial and ecological species richness, such as those from 852 the late Early Triassic Feixianguan and Lingwen Floras that contain various conifers 853 854 (Zhang et al., 1992; Li et al., 1995). Following the dominance of spores over pollen in the early Griesbachian 855 856 substage, the percentage of pollen increases in the South China palynological record (Zhang et al., 2004; Yu et al., 2008; Ouyang and Zhu, 2007). Although there is only a 857 single, rare megafossil species of *Peltaspermum* in the Kayitou Formation, 858 palynology indicates the widespread presence of seed plant groups that are absent 859 from the macrofossil record (Fig. 17). 860 861 The majority of the plant taxa in Phase 2 are inherited from the Permian

suggesting the Kayitou Flora belongs to the survival stage rather than the recovery

during this interval (Xie et al., 2007; Shen W.J. et al., 2011, 2012; Yin et al., 2012; Fig. 864 20), 865 During Phase 2 in the Kayitou Formation, the plant extinction rate is higher than 866 the origination rate, due to the demise of holdover taxa. This is followed by an 867 868 absence of plants fossils from the middle Kayitou Formation to the middle of the Dongchuan Formation (Feng et al., 2018) (Fig. 1). 869 870 4.3. 871 Phase 3: Late Griesbachian–Smithian species richness recovery Phase 3 marks the start of the post-EPPC recovery and is recorded in the 872 Dienerian-Smithian Feixianguan Formation. The Feixianguan Formation contains 16 873 genera and 17 species, thus with low species numbers within individual genera. The 874 origination rate in the Feixianguan Formation exceeds the extinction rate for the first 875 time since the EPPC, although extinctions still occurred but at low levels (< 20 876 species extinctions per time bin). Within individual plant groups (Fig. 19), the 877 878 origination rate of Mesozoic lycopod, sphenophyte, cycads, ferns and conifers noticeably exceeds their extinction rates, whilst the origination rates of seed plants for 879 880 example seed ferns, ginkgophytes and some gymnosperms, such as Pelourdea 881 (=Yuccites), are similar to the extinction rates. Plant taxa in the Feixianguan Formation consists of small numbers of Mesozoic-882 type lycopods, abundant sphenophyll branches (including Neocalamites and 883 884 Equisites), and includes Mesozoic-type ferns (Todites spp., Anomopteris, 885 Dictyophyllum (=Thaumatopteris)), seed ferns (Peltaspermum sp.), cycads 886 (Taeniopteris sp.), ginkgophyte (Baiera sp., Sphenobaiera sp.) and some conifers 887 (Albertia sp., Voltzia heterophylla, Voltzia sp.) (Zhou et al., 1979; Figs. 10). Most of the plants, especially *Peltaspermum* and the conifers, are considered to be floral 888 elements adapted to dry and hot climates based on their thick cuticles (Poort and Kerp, 889 1990; Huang and Lu, 1992; Taylor et al., 2009). 890 891 Early Triassic (Late Griesbachian to Smithian) palynological data from South China is absent whilst the Xinjiang record from North China is derived from from the 892

stage. Wildfire proxies, such as black carbon, show that wildfires were still prevalent

893 Jiucaiyuan and Shaofanggou formations (Qu et al., 1986). Spores dominate over pollen in both formations although the younger levels of the Shaofanggou Formation 894 have slightly higher pollen content (Qu et al., 1986). The abundance of the Triassic 895 lycopod spore Lundbladispora in those two formations denotes the success of 896 897 lycopods during the Griesbachian and Smithian (Qu et al., 1986). Using biomarker proxies for plant abundance, retene, simonellite and dehydroabietane, which are likely 898 to be derived from herbaceous rather than woody plants because of low C/N ratios 899 900 (<10), suggests recovery was underway during the Smithian (Saito et al., 2013). Nonetheless, plant fossils are rare and coal formation is still absent during Phase 3. 901 902 4.4. Phase 4: Spathian abundance recovery 903 904 The Spathian Lingwen flora and the Anisian Badong flora are dominated by Mesozoic-type lycopods Lepacyclotes (=Annalepis) and Pleuromeia alongside 905 sphenopsids (Equisites and Neocalamites), while gymnosperms (Taeniopteris, 906 907 Peltaspermum (Vittaephyllum), Pelourdea (=Yuccites) and the conifer Voltzia) are 908 common (Figs. 14, 15). Diversity within the Lingwen and Badong formations are broadly similar to that of the Early Triassic Feixianguan Formation with 22 genera 909 910 and 29 species in Lingwen flora and 17 genera and 27 species in Badong flora present. However, plant abundance is much higher in the Badong Formation, as reflected by 911 912 fossil abundance and the development of peat (Meng et al., 1995). Significantly, 913 Member 2 of the Badong Formation yields the terrestrial tetrapod *Lotosaurus* (Figs. 914 14, 15), showing there was sufficient vegetation to support large herbivores (Hagen et al., 2018). 915 916 According to palynological data in South China, gymnosperm pollen is the major 917 constituent, especially in the Spathian Lingwen Formation (Zang et al. 1992; Meng et al. 1995). In the Xinjiang area, gymnosperm pollen content slightly exceeds that of 918 spores in the Shaofanggou Formation and is increasingly common in the Karamay 919 Formation of the Xinjiang Dalongkou section. This is similar with the Badong 920 921 Formation of South China in the Hongjiaguan section (Qu et al., 1990; Meng et al., 1995) and indicates progressive vegetation changes during the Spathian. Increased 922

923 abundance of the conifer derived biomarker pimanthrene during the Spatian in South 924 China denotes a vegetation change from the Griesbachian to Smithian 925 lycopods/herbaceous bryophytes to conifer-dominated floras (Saito et al., 2013) and correlates well with gymnosperm pollen dominance of over 90% in Lingwen flora 926 927 (Zhang et al., 1992) (Fig. 22). In addition, rapid increases of C/N ratios (>10, reaching 28) after the Smithian-Spathian boundary in South China from the Chaohu section, 928 reveals the flourishing of vascular land plants (Saito et al., 2013). Plant macro fossils, 929 930 palynology and biomaker data all record species richness and abundance recovery 931 happened after the Spathian.

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5. Discussion

934 *5.1 Distinctions between macro- and micro-floral (palynology) data*935 In paleobotanical studies across the PTB in South China, the study of Xiong and

Wang (2011) stands out for documenting a gradual, stepwise loss in plant megafossil

species richness in the run up to the EPPC while concurrent palynological records

only recorded a minor fluctuation in species richness. Their study was based on an

uncritical compilation of data from literature with identifications that were not

verified by examination of original materials, and species were not collected in a

detailed, bed-by-bed stratigraphic framework as undertaken here for the EPPC

interval. In our study, as well as having the extinction level, our megafossil data also

shows a gradual, stepwise loss of megafossil species richness (Figs. 4, 5, 7, 9),

presumably related to gradual facies and/or environmental changes (MacLeod, 1997;

Stevens et al., 2011) in the run up to the extinction level. These environmental or

946 facies changes adversely affected plants in wetland, peat forming clastic settings (e.g.,

Wang et al., 2011; Yan et al., 2019; Feng et al., 2020), but they do not provide insights

into the vegetation from contemporaneous upland, extrabasinal settings (see

DiMichele et al., 2020) such as the Khangdian Oldland in South China (Fig. 1; Wang

950 et al., 2020).

We consider this is a probable consequence of taphonomic megabias in which the microfossil record potentially samples a larger geographical source area including

954 lowland depositional settings (e.g., Looy et al., 2004; Neregato et al., 2016; DiMichele et al., 2020; Cleal et al., 2021). Support for this comes from the 955 characteristically Mesozoic palynomorphs *Wilsonisporites* (unknown affinity), 956 957 Neoraistrickia (putative isoetalian lycopsid; Singh 1971) and Pteruchipollenites 958 (corystosperm gymnosperm) found in conglomerates from the basal Xuanwei Formation (Neregato et al., 2016) whilst the plants that produced them are absent 959 960 from the megaflora. This shows their parent plants persisted in South China outside 961 the coastal wetlands preservation window and survived the EPPC in this region. In addition, xerophyte palynomorphs recorded in the Chahe section (Yu et al., 2008) are 962 distinct from the contemporaneous wetland megaflora and include disaccate striatiti 963 (Coniferopsida), Protohaploxypinus and Vittatina (Peltaspermales, Ginkgopsida; 964 Balme, 1995), Lueckisporites (Majonicaceae, Coniferopsida; Clement-Westerhof, 965 1974), Striatopodocarpidites (Glossopteridales, Ginkgopsida; Pant, 1977; Balme, 966 1995) and Taeniaesporites (=Lunatisporites: Podocarpaceae, Coniferopsida; Clement-967 968 Westerhof, 1974). While first appearing in the late Permian Xuanwei Formation, these palynomorphs became dominant in the Early Triassic Kayitou Formation where they 969 played significant roles in post-EPPC floras and the EPPC recovery (Fig. 17). 970 971 In the Changhsingian, spore producing plants including lycopods, sphenophytes 972 and ferns were dominant in the megafossil record, while Paleozoic lycopod spores are 973 absent from the palynology record (Fig. 16, 17). This discrepancy indicates that to 974 fully characterize the flora information from both sources are required. Palynological data likely samples flora from a wider setting than just the lowland depositional 975 976 environments of the Xuanwei Formation but has limits in reconstructing the affinity diversity and abundance as it is often hard to correlate palynological species with 977 parent plants. In contrast, plant macrofossil data tends to record more localized areas 978 979 in the Xuanwei Formation in detail, but it does not necessarily represent the vegetation from the entire basin. The combined macro- and micro-floral data indicates 980 981 that the end Permian Changhsingian lowlands of the Xuanwei Formatoin were occupied by Paleozoic lycopods, sphenophytes, fern, progymnosperms gigantopterids 982

uplands compared to megafossil assemblages that are extensively restricted to

and seed ferns while the uplands were dominated by other gymnosperms including conifers, ginkgophytes, cycads and peltaspermalean seed ferns. The proliferation of fungal spores indicates a widespread land ecosystem crisis in South China during the EPPC. After the EPPC, lowland floras were left with only a few Paleozoic holdover taxa and pioneering Triassic lycopods growing in coastal areas, while the uplands saw the persistence of gymnosperm-like peltasperms, cycadophytes, ginkgophytes and conifers: a Permo-Triassic transitional flora. Soon after the early Griesbachian, the survival flora died out in lowland areas, and gymnosperms previously occupying upland habiats gradually occupied the empty niches and formed what became typical Mesozoic gymnosperm-dominated floras (Fig. 17). The palynological record shows this transformation may have started in the late Changhsingian, although it is only seen in the macrofloral record after the Griesbachian (Fig. 16, 17).

Ouyang (1991) noted that about 30–50% of palynology species from the Permian–Triassic transitional flora at the bottom of the Kayitou Formation in Yunnan province were holdovers from Permian or older ages and comprised exclusively of gymnosperm pollen. Of these only 15–17% extended into the later Early Triassic (Fig. 17). We consider these gymnosperm pollen as Methuselah taxa (see Looy et al., 2004; Blomenkepmer et al., 2018; DiMichele et al., 2020) with unexpectedly early stratigraphic occurrences that were living outside the preservationally biased wetland settings in ecological niches such as upland fluvial and lacustrine systems less affected by the EPPC extinction mechanisms. Further study is required to evaluate the taphonomic nature of late Permian palynofloras to confidently identify Methuselah taxa, and where possible, match the dispersed spore and pollen accounts to plant groups to characterize in detail for the first time the composition of these cryptic upland floras.

5.2 Permian–Triassic extinction on land and in ocean

As to the age of the terminal phase of the EPPC, the Hg/TOC spikes and the carbon isotope trends in Chinahe and many other terrestrial sections can be correlated with the marine GSSP at Meishan (Shen J. et al., 2019b; Chu et al., 2020): a peak of

Hg/TOC was recorded in Meishan Bed 24, and in Bed 26 at Chinahe (Fig. 20). The
former records the first, severe phase of the marine mass extinction but at Chinahe the
main plant mass extinction occurs below this in Bed 25 indicating an earlier terrestrial
crisis. The pioneer lycopod genus <i>Tomiostrobus</i> (=Annalepis) occurs immediately
below the Hg/TOC peak and maybe used for correlation due to its stratigraphically
short-ranging and geographically widespread distribution in South China (Yu et al.,
2010). Further evidence for this earlier crisis comes from radiometric dating at the
Chahe section, where the loss of plants occurs in Bed 69. This level is constrained by
a zircon age from Bed 68 of 252.30 \pm 0.07 Ma. This is close to the age of 252.104 \pm
0.089 Ma in Bed 22 and 251.941 \pm 0.037 Ma in Bed 25 at Meishan (Shen S.Z. et al.,
2011; Burgess et al., 2014). The first phase of the marine extinction at Meishan
therefore lagged behind the floral crisis by tens to hundreds of thousands of years (Fig.
20; Yin et al., 2012; Cui et al., 2017; Dal Corso et al., 2022; Wang Y. et al., 2022).
This conclusion is supported by the two fungal spore peaks in Bed 66 and 68, and the
proliferation of gymnosperm pollen in Bed 70 and 78 of the Chahe section (Yu et al.,
2008). In the marine facies of the Meishan Section, the end Permian to early
Griesbachian palynological record is more continuous and shows the increasing
dominance of gymnosperm pollen from Bed 27 (Yu et al., 2008; Zhang et al., 2004)
after the Permian-Triassic Boundary in the early Griesbachian.
Based on moretane/hopane (C ₂₉ -M/C ₃₀ -HP, C ₃₀ -M/C ₃₀ -HP) ratios and the
biomarker DBF index (DBF/DBF+DBT+F), anomalously high terrestrial organic C
inputs occurred in the latest Permian (Beds 25-26) at Meishan, before gradually
decreasing in the Early Triassic (Beds 27–30) before increasing again in Bed 34 (Xie
et al., 2007, 2009; Wang, 2007). According to this timescale, the marine faunal
extinction episode at the base of Bed 25 at Meishan (Fig. 20) and the marine
productivity decline before Bed 25 (Song et al., 2012; Shen J. et al., 2015) occurs
after the terrestrial EPPC, whilst plant abundance declined to its lowest level after
Meishan Bed 24.

To evaluate the paleoenvironmental influences of the floral changes evaluated here, a timescale for environmental events has been compiled through the end Permian to Middle Triassic. This is divided into two phases: the Changhsingian to Griesbachian interval set against the timescale of the marine Meishan section from which precise zircon ages have been determined (Fig. 20), and, with lower stratigraphic resolution, the Induan to Anisian interval (Fig. 22).

The EPPC is thought to coincide with a long-term aridification associated with the formation of Pangaea beginning in the Middle Permian (Kidder et al., 2004; Roscher et al., 2011; Benton et al., 2014; Blomenkemper et al., 2018). Rising extinction rates in South China began in the Changhsingian and coincide the start of a trend that saw pollen percentages climb (see supplementary dataset for the macro and micro spore and pollen plant percentage from Wuchiapingian to Ladinian) (Figs. 16, 19). Increased charcoal concentrations in the latest Changhsingian suggest aridity intensified, likely seasonally, as the climax of the EPPC developed (Shen W.J. et al., 2011; Yan et al., 2019; Chu et al., 2020; Cai et al., 2021). The increase of the chemical weathering index (CIA) in South China at the same time (e.g. Xu et al., 2017) could reflect the loss of plant cover. The increase of fungi could also be caused by more prolonged arid episodes (Berdugo et al., 2020).

Increasing drought and climbing temperature could all have weakened the gigantopterid (Cathaysian) flora in South China and lead to the the mass extinction that marked the culmination of the EPPC. Whether this was a culmination of stresses that began in the early Changxingian or if the terminal EPPC was a distinct event, with a separate causation, can be debated. However, prior to the rapid warming of equatorial, ocean surface-waters in the latest EPPC, temperatures were stable and rather cool during the Changxingian (Joachmiski et al. 2020) which argues against the notion of progressive temperature rise reaching a lethal threshold at the end of the EPPC. Instead, the effect of rapid warming appears to have been impact a South China flora that was already experiencing diversity decline perhaps due to increasing seasonal aridity. Siberian volcanism is generally apportioned the blame for the rapid warming episode and other consequences of the eruptions may have been acid rain

and depletion of the ozone layer resulting in increased UVB radiation (Benca et al., 2018; Black et al., 2018; Cai et al., 2021; Fig. 21).

Volcanism-induced weathering on land and increasing terrestrial inputs play significant role to the marine ecosystem (e.g., Shen J. et al., 2022; Huang et al., 2022). Modelling of factors such as volcanism, tectonism, marine redox and acidification, cannot explain the extreme hothouse climate in Early Triassic without including the terrestrial biome (Mills et al., 2021, Fig. 21). We inferred the enhancement of climate instability, seasonal aridity and following loss of lowland peat vegetation during EPPC caused the drop of terrestrial biomass storage, probably contributing to the carbon cycle fluctuation, while this requires further study of land biomass and global carbon cycle.

Wildfire proxies probably indicate the disappearance of the Griesbachian interval vegetation after the early Griesbachian (Fig. 20). The initial loss of the holdover flora might result in a temporary increase of soil erosion (Fig. 20). The flora of this interval was dominated by the herbaceous lycopods *Tomiostrobus* (=*Annalepis*) and *Pleuromeia* which, with their shallow rooting systems (Retallack et al., 1975; Yu et al., 2010), were likely insufficient to effectively bind soils (Algeo et al., 2011; Boyce et al., 2016; Fig. 22).

From our data, plant species richness recovery occurred during the Griesbachian to the Smithian stage, while plant abundance indicated by the palynology data, biomakers, TOC and C/N ratio data this aspect began to recover in the Spathian (Saito et al., 2013). The first post-EPPC herbivorous tetrapods appeared and coal accumulation re-commenced in the Anisian, indicating a return of diverse and productive terrestrial ecosystems. The soil erosion proxy also indicates the restabilization of land surface system after Spathian (Algeo et al., 2011). Diverse marine ecosystems were also reestablished at Anisian but full recovery to a pre-extinction level was not until the Late Triassic (Song H.J. et al., 2018).

5.4 Comparison of floristic patterns between low latitude South China and other geographical areas

Both the North and South China plates occupied low–middle latitude positions
during the Late Permian and experienced tropical-subtropical climates (Nowak et al.,
2020). The Late Permian vegetation from North China was a mixed Cathaysian,
Euramerican and Angara flora, whilst a typical Cathaysian flora occupied South China
(Wang et al., 1985; Yu et al., 2015; Wu et al., 2021). In North China terrestrial
depositional facies have made identification and correlation of the PTB and PTME
difficult, with recent investigations using radiometric ages from ash beds to confirm
the End Permian Plant Crisis predates the PTME which concludes with the PTB in the
uppermost Sunjiagou Formation from the Liujiang Coalfield (Wu et al., 2021; Wang Y
et al., 2022). The terrestrial ecosystem collapse in North China commenced
approximately 270±150 kyrs before the marine crisis (Guo et al., 2022), but occurs
approximately 310 kry later than the terrestrial crisis in high southern latitudes in
Australia (Lu et al. 2022). Prior to the PTME, the plant macrofossil extinction and
origination rates in North China are comparable to those of South China and indicate
significant floral turnover (Xiong et al., 2021). In both areas plant extinction rates
exceeded origination rate before the EPPC, but in North China the severest plant crisis
event, which is shown by the biggest value difference between extinction and
origination rates, occurred before the PTB boundary and may been earlier than South
China (Xiong et al., 2021), although Lu et al. (2022) considered they may be
synchronous. After the terrestrial plant crisis and PTME, the earliest Triassic flora in
North China comprised similar pioneering Triassic isoetalean and Pleuromeia
lycopods and later in the early Triassic conifer dominated floras (Yu et al., 2015;
Xiong et al., 2021).
The fossil record from Australia in high latitude Gondwana reveals that the
Glossopteris flora suffered abrupt extinction due to rapid warming and increased
seasonality somewhat before the Permian-Triassic Boundary (Vajda et al., 2020;
Frank et al., 2021; Fielding et al., 2022). This ecological disaster reset Paleozoic
terrestrial phytogeographic provincialism and marked the end to the former separation
of floras into the low-mid latitude Euramerican and Cathaysian floras and the high-
latitutde Gondwana floras. The peak of plant species richness decline, last coal seam,

and ecosystem collapse indicated by fungal spike in Australia, South and North China 1133 1134 all denote the onset and main peak of the land plant crisis occurred tens of thousands of years before the marine crisis (Yu et al., 2008, 2015; Xiong et al., 2021; Fielding et 1135 1136 al., 2022). Spikes of fungal spores are common in Australia, South China, and the 1137 Karoo Basin where they occur at several levels at this time (Visscher et al., 1996; Steiner et al., 2003; Ouyang and Zhu, 2007; Yu et al., 2008; Fielding et al., 2022). 1138 Early Triassic floras from low to high latitudes comprised a uniform lycopod 1139 1140 dominated flora (e.g. in Australia and South and North China). In the Kuznetsk Basin in Russia, increased aridity may have affected the 1141 composition of the Angaran flora, but this region saw floral turnover and migration in 1142 response to changing climate rather than an extinction event (Davydov et al., 2021). 1143 The regional extinction of the humidity-adapted, cordaites-dominated flora happened 1144 approximately 820 kyrs earlier than the PTME marine extinction event in South China 1145 (Davydov et al., 2021). Following the floral turnover, plants subsequently diversified 1146 1147 across the Permian-Triassic transition when mixed fern (Cladophlebis, 1148 Kovuntschania, Katasiopteris, Kchonomakidium, Todites, Kedroviella and Prynadaeopteris), sphenophyte (Neokoretrophyllites, Schizoneura, Paracalamites) 1149 peltasperm (Lepidopteris), seed fern (Tersiella and Madygenia), cycad (Tomia and 1150 Glossozamites), Ginkgoales (Rhipidopteris and Glossophyllum), conifer 1151 1152 (Quadrocladus) and Triassic lycopods (Tomiostrobus, Mesenterihyllum) characterized 1153 the flora (Davydov et al., 2021). Comparison between floras in different latitudes and in various distances from 1154 continental interiors shows that climate instability and expansion of seasonal aridity 1155 1156 was a significant control on floral composition and distribution through the Permian and Triassic transition. The plant mass extinction level occurred over wide areas with 1157 only the Siberian region recording a diverse flora in the aftermath of the crisis and 1158 1159 turnover (Davydov et al., 2021). Given the proximity of this region to the flood basalts of the Siberian Traps it is ironic that the flora of the Phase 2 interval was so 1160 1161 diverse. It could be argued that factors that are at their most intense adjacent to volcanism, such as acid rain, may not therefore have been an important factor in the 1162

floral mass extinction. Other factors such as a relatively muted temperature rise, in the high northern Siberian latitudes, and a persistent humid climate may all have favoured this region as a refuge.

In tropical areas such as South China, a rapid temperature rise of over 15°C proved fatal, resulting in ocean surface temperatures > 35°C degrees, and possibly > 42°C on land; such levels are likely to have been directly responsible for the extinction losses (Sun et al., 2012). In higher latitudes, the peak temperatures would have been lower whilst still exceeding the tolerance of indigenous plans (Fielding et al., 2022), although perhaps not in the Siberian region (Davydov et al., 2021). Plants living in higher altitudes may also have been more resilient to extreme temperatures due to temperatures typically decreasing adiabatically with height. Consequently, upland floras were able to colonize lowland settings after the EPPC once competition pressures (and temperatures) in these settings were lower following extinction.

6. Conclusions

Investigation of plant macrofossil occurrences from the Artinskian to Rhaetian in South China has shown that floral species richness declined after the Wuchiapingian and experienced a distinct species richness and abundance drop in the Changhsingian that we term the End Permian Plant Crisis (EPPC). During the EPPC plant extinction rates overtook origination rates, with this scenario continuing into the early Triassic although the gap narrowed after the EPPC. The culmination of the EPPC was marked by a major extinction with losses of coal-swamp taxa including tree lycopods (*Lepidodendron*), sphenopsids (*Lobatannularia*, *Annularia*), Noeggerathiales progymnosperms (*Tingia*), Marattiales ferns (*Pecopteris*), gigantopterids (*Gigantopteris*) and cordaites gymnosperms (*Cordaites*) which flourished during the Late Paleozoic (Phase 1). There seems to have been a hidden upland gymnospermdominated flora at this time which is not preserved in the macrofossil record but is evident in palynological data. Other evidence of terrestrial plants, such as wildfire and terrestrial input proxies obtained from marine sections, indicate the climax of the terrestrial EPPC predated the marine PTME extinction.

In the early Triassic Kayitou Flora, surviving end-Permian elements were mixed with Triassic opportunist herbaceous lycopods with low species richness and abundance. We term this the Griesbachian interval flora (Phase 2). Origination rates only began to exceeded extinction rates in the late Induan, and this trend continued into the Olenekian. The recovery of abundant plant biomass happened later than the diversity recovery, and probably commenced during the Spathian as indicated by increasing conifer biomarker concentrations and C/N ratios, and continued into the Anisian. Triassic floras only attained a comparable species richness in South China to the pre-EPPC flora after the Carnian (early Late Triassic). According to the origination and extinction rates of each plant family, the flora overall changed from a Paleozoic Cathaysian peat forming type into Mesozoic seed plant-dominant type which likely reflects an adaption to drier climate.

Declaration of Competing Interest

The authors declare they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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1799	
1800	Table and figure captions
1801	Table 1. Example of normalized occurrences for species richness estimates within the
1802	arborescent lycopsids showing the distribution of organ taxa for stems
1803	(Lepidodendron), rootstock (Stigmaria), sporophylls (Lepidostrobophyllum) and
1804	cones (Lepidostrobus). In each formation, stems represent the best measure of species
1805	richness as they present the most reliable features to distinguish species, and in each
1806	case show the largest species richness compared to the other organs present. While
1807	more species of rootstock are present in the Xuanwei Formation, we consider this an
1808	unreliable measure of species richness. Species delimitation is less reliable in species
1809	of Stigmaria that have few distinguishing features that may vary in different positions
1810	across the rooting system. Furthermore, one of the rootstock accounts from the
1811	Xuanwei Formation has not been identified to the species level (Stigmaria sp.) and
1812	most likely represents a poorly preserved or incomplete specimen of one or more of
1813	the other species present. In all cases non-normalised estimates significantly inflate
1814	species richness estimates.

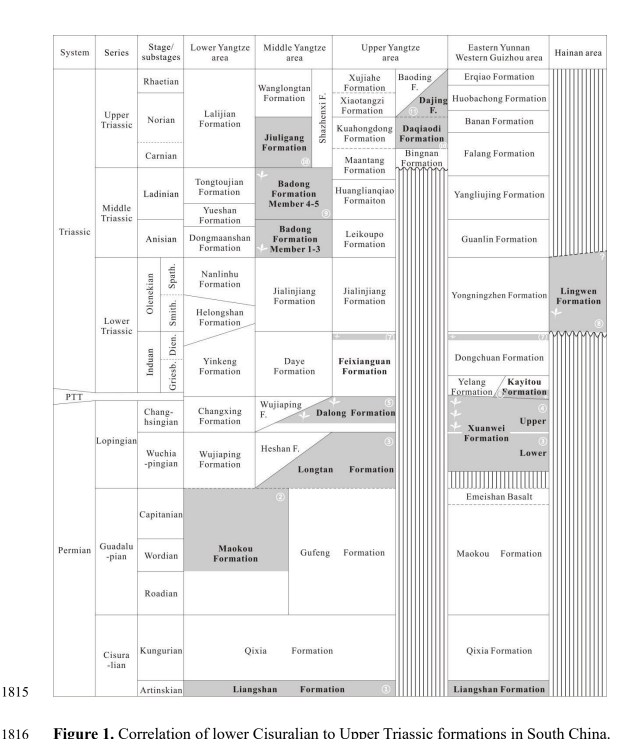


Figure 1. Correlation of lower Cisuralian to Upper Triassic formations in South China. gray units contain plant fossils, with leaf representing position of separate beds containing plants. Numbers in formations represent: 1. Liangshan section; 2. Maokou section; 3. Longtan and lower Xuanwei section; 4. Chahe and Chinahe sections; 5. Xinmin, Duanshan A and B sections; 6. Chinahe, Mide and Tucheng sections; 7. Lubei, Pojiao and Dongchuan sections; 8. Lingwen section; 9. Hongjiaguan and Furongqiao section; 10. Jiuligang and Daqiaodi sections; 11. Xujiahe, Dajing, Anyuan, Bagong sections. PTT = Permo-Triassic transition. Figure modified from Yang et al.

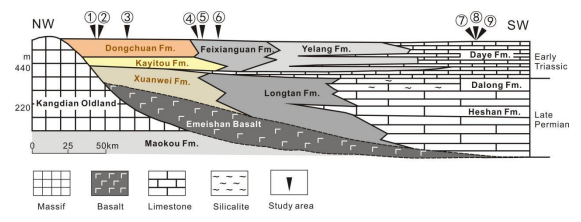


Figure 2. Sketch map of the Permian Changsingian lithofacies in Western Guizhou and Eastern Yunnan, southwestern China. 1. Pojiao section; 2. Lubei section; 3. Chahe section; 4. Chinahe section; 5. Mide section; 6. Tucheng section; 7. Xinmin section; 8. Duanshan A section; 9. Duanshan B section; modified from Yu et al. (2015), Wignall et al. (2020).

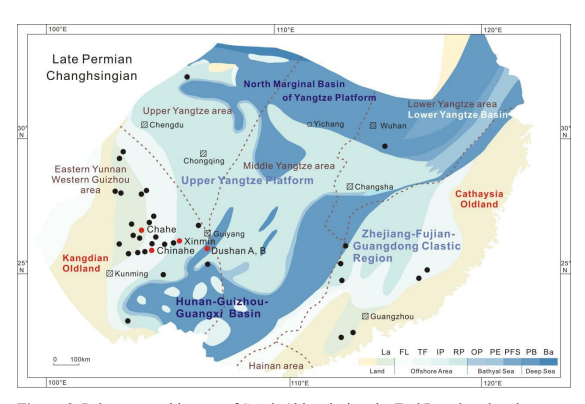


Figure 3. Paleogeographic map of South China during the End Permian showing positions of sections studied from the Changhsingian aged Xuanwei and Dalong formations. Red point: studied sections in this paper; Black point: supplementary

sections from literature; La = Lacustrine; FL = Flood land; TF = Tide flat; IP = Isolate platform; RP = Regional platform; OP = Open platform; PE = Platform edge; PFS = Carbonate platform fore slope; PB = Platform basin; Ba = Bathyal sea; modified from Zheng et al. (2011), Yin et al. (2014) and Yu et al. (2015).



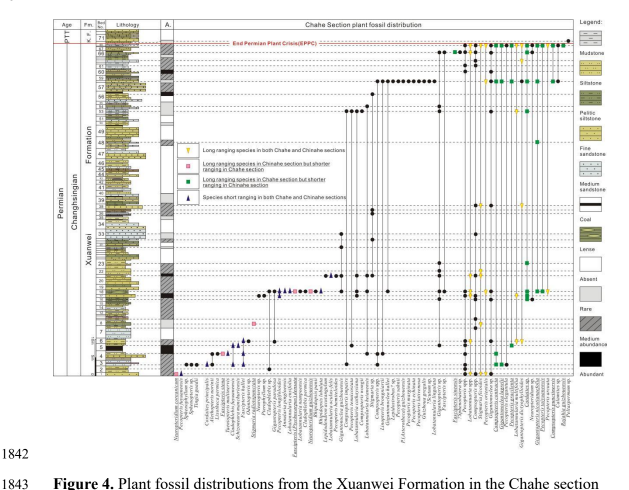


Figure 4. Plant fossil distributions from the Xuanwei Formation in the Chahe section (unnormalized). PTT = Permo-Triassic transition; Grib. = Grisbachain; K. F. = Kayitou Formation; A. = Abundance of plant fossil. The color of the lithology column shows the real rock color in the field.

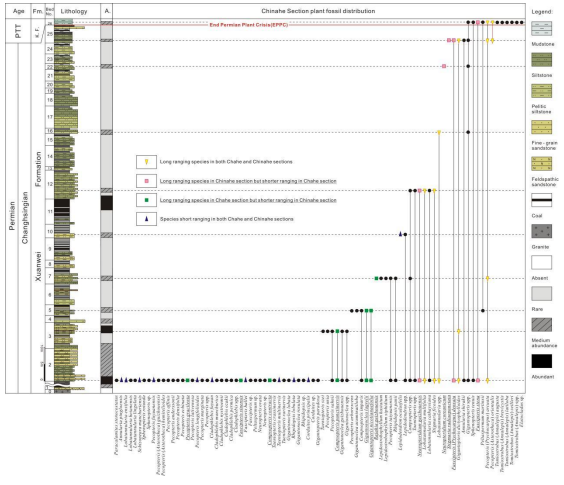


Figure 5. Plant fossil distributions from the Xuanwei Formation in the Chinahe section (unnormalized). PTT = Permo-Triassic transition; K. F. = Kayitou Formation; A. = Abundance of plant fossil. The colour of the lithology column shows the real rock colour in the field.



Figure 6. Field panorama, sedimentology and representative plant fossils showing preservation condition from the Chinahe section. 1. Strata of the Kayitou Formation; 2. Strata of the upper Xuanwei Formation; 3. Emeishan Basalt with vesicular structure; 4. Boundary between the Emeishan Basalt and the Xuanwei Formation; 5. Panorama of the Chinahe section ranging from the Emeishan Basalt (right) to the Kayitou Formation (left); 6. Conchostraca in the Kayitou Formation; 7. *Tomiostrobus* (=Annalepis) layer in the bottom of Kayitou Formation (Bed 26); 8. Abraded Gigantopteris dictyophylloides fragments together with Tomiostrobus (=Annalepis) (Bed 26); 9. Small Peltaspermum martinsii together with Tomiostrobus (=Annalepis) (Bed 26); 10. Broken Compsopteris leaf in the upper part of Xuanwei Formation (Bed 22); 11. Layer of Lepidodendron oculus-felis in middle of Xuanwei Formation (Bed 10); 12. Gigantopteris fragments occurring from the lower to middle of the Xuanwei Formation (Beds 3–10); 13. Well-preserved leaves with insect feeding trace fossils (bed 2); 14. Complete fern branches in the bottom of Xuanwei Formation (Bed 1). PTT = Permo-Triassic transition; P = Permian.

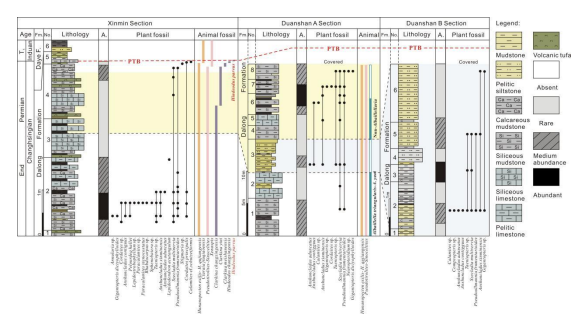


Figure 7. Plant and marine animal fossil distributions from the Dalong Formation in the Xinmin, Duanshan A and Duanshan B sections (unnormalized). T₁ = Early Triassic; Fm. = Formation; NO. = Bed number; A. = Abundance of plant fossil; PTB = Permian Triassic boundary. The color of the lithology column shows the real rock color in the field.

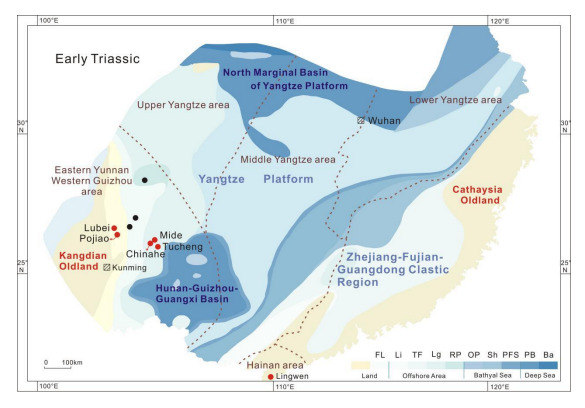


Figure 8. Early Triassic paleogeographical map of South China showing locations of

sections containing the Induan Kayitou Formation, Olenekian Feixianguan and Lingwen formations. Red point: studied sections in this paper; Black point: supplementary sections from literature; FL = Flood land; TF = Tide flat; Lg = Lagoon; RP = Regional platform; OP = Open platform; Sh = Shallow sea; PFS = Carbonate platform fore-slope; PB = Platform basin; Ba = Bathyal sea; modified from Zheng et al. (2011) and Yin et al. (2014).



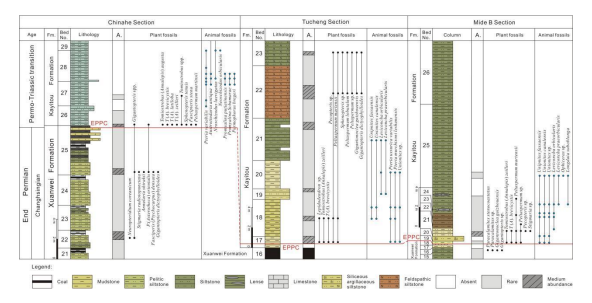


Figure 9. Plant and marine animal fossil distributions from the Kayitou Formation in the Chinahe, Tucheng and Mide sections (unnormalized). Fm. = Formation; NO. = Bed number; EPPC = End Permian Plant Crisis; A. = Abundance of plant fossil. The color of the lithology column shows the real rock color in the field.

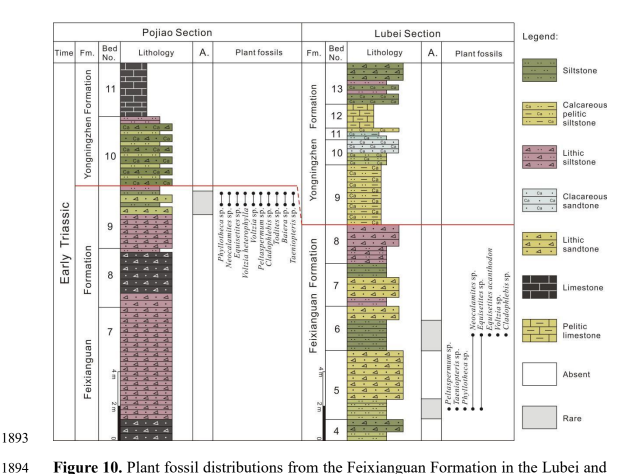


Figure 10. Plant fossil distributions from the Feixianguan Formation in the Lubei and Pojiao sections (unnormalized). Fm. = Formation; NO. = Bed number; A. = Abundance of plant fossil. The color of the lithology column shows the real rock color in the field.

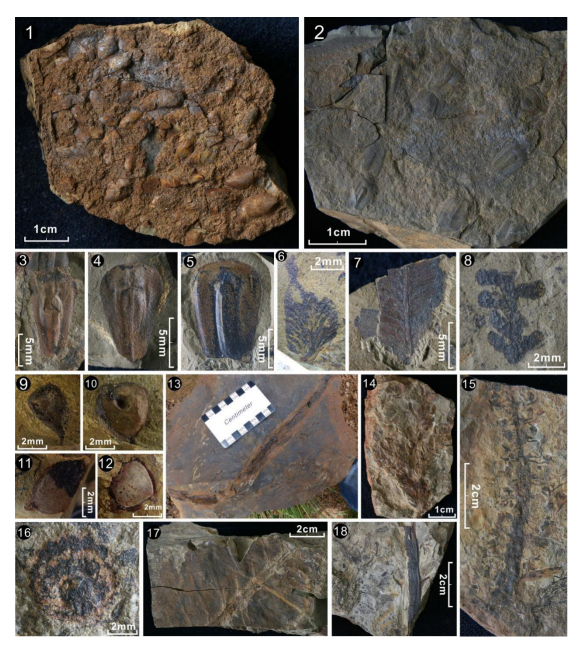


Figure 11. Representative plant fossils in Kayitou Formation of Chinahe section (1–8) and Feixianguan Formation of Lubei and Pojiao section (9–15). 1. Bivalves; 2. *Tomiostrobus (=Annalepis)* spp.; 3. *T. (A.) zeilleri*; 4. *T. (A.) augusta*; 5. *T. (A.) latiloba*; 6. Unkown index; 7. *Fascipteris stena*; 8. *Peltaspermum martinsii*; 9–12. *Carpolithus* spp.; 13. *Neocalamites* branches, common in both Lubei and Pojiao sections; 14. *Voltzia* sp.; 15. Possible fertile spike?; 16. *Peltaspermum* sp.; 17. Fern; 18. *Todites* sp.

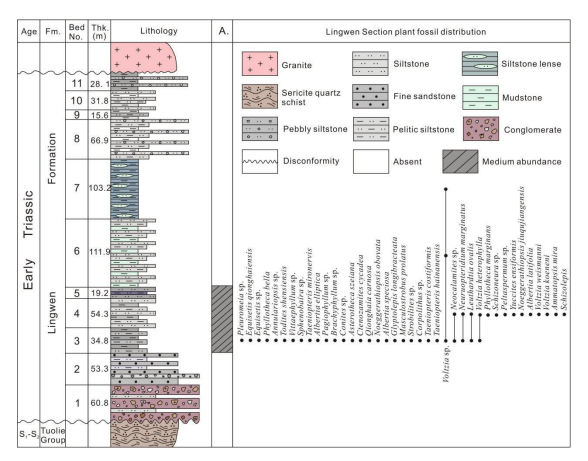


Figure 12. Plant fossil distributions from the Lingwen Formation in the Lingwen section (unnormalized). Fm. = Formation; NO. = Bed number; Thk. = Thickness; A. = Abundance of plant fossil. The color of the lithology column shows the real rock color in the field.

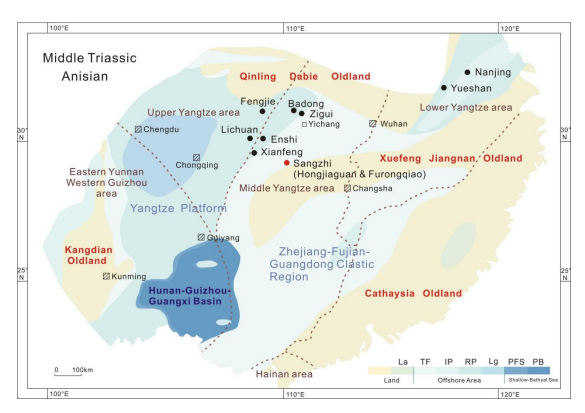


Figure 13. Middle Triassic paleogeographic map of South China showing locations for sections of the Badong Formation. Red point: studied sections in this paper; Black point: supplementary sections from literature; La = lacustrine; TF = Tide flat; IP = Isolated platform; RP = Regional platform; Lg = Lagoon; PFS = Carbonate platform fore slope; PB = Platform basin; modified from Zheng et al. (2011).

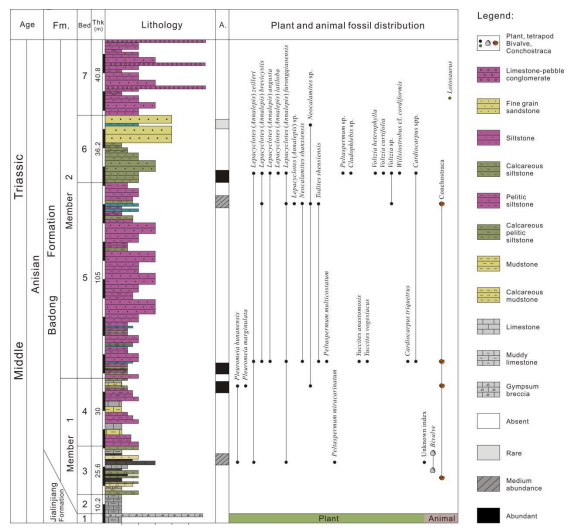


Figure 14. Plant fossil distributions from the Badong Formation in the Hongjiaguan and Furongqiao sections (unnormalized). Fm. = Formation; NO. = Bed number; Thk. = Thickness; M. = Mineral; J. = Jialinjiang Formation; A. = Abundance of plant fossil. The color of the lithology column shows the real rock color in the field.

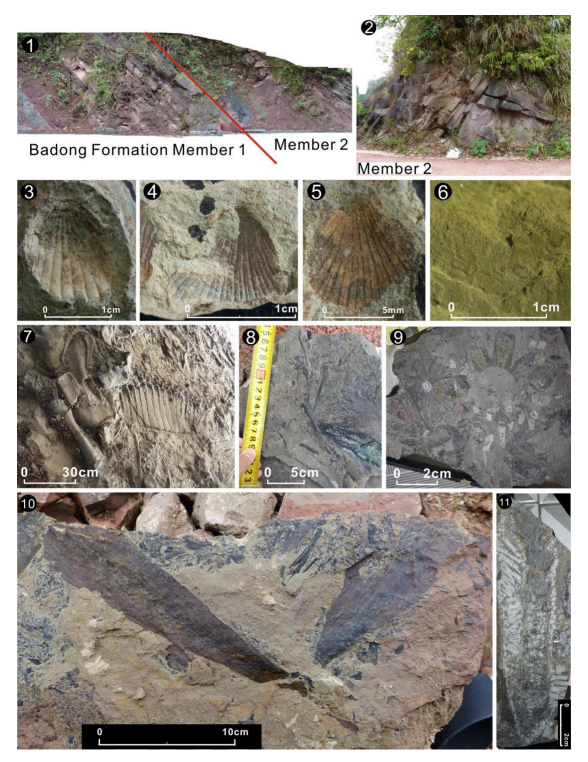


Figure 15. Field panorama, sedimentology, representative animal and plant fossils of the Badong Formation from Hongjiaguan and Furongqiao sections. 1. Lithological boundary of Badong Formation Member 1 (left) and Member 2 (right); 2. Thickbedded sandstone Member 2 (Bed 6 in figure 14); 3. *Myophoria (Costatoria) goldfussi*; 4. *Myophoria (Costatoria) goldfussi mansuyi*; 5. *Leptochondria albertii*; 6. *Euestheria* sp.; 7. *Lotosaurus* in Member 2 in Furongqiao village; 8. Long-distance

transported plant fragments in sandstone from Member 2; 9 *Lepacyclotes (=Annalepis)* sangzhiensis in the Hongjiaguan section (collected by Fansong Meng); 10. *Yuccites* sp.; 11. Preserved in-situ *Pleuromeia sanxiaensis* in the Dawotang section, Fengjie, Sichuan (collected by Fansong Meng).

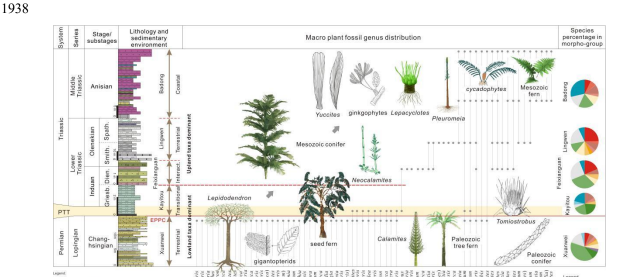


Figure 16. Lithology, sedimentary, macro plant fossil distribution range, floral composition from End Permian Changhsingian to Middle Triassic Anisian in South China area. Kayitou Formation conformably overlies on Xuanwei Formation, while Kayitou, Feixianguan, Lingwen and Badong formation does not directly connect with each other and are divided by dash line. Xuanwei Formation: terrestrial facies; Kayitou Formation: terrestrial marine transitional facies; Feixianguan Formation: terrestrial marine interacting facies; Lingwen Formation: terrestrial facies; Badong Formation: coastal facies. Legend of macro plant morpho group: 1. conifer, 2. gymnosperm, 3. peltasperm, 4. seed fern, 5. cordaites, 6. ginkgophyte, 7. cycadophyte, 8. Noeggerathiales, 9. gigantopterid, 10. fern, 11. fern or seed fern, 12. sphenophyte, 13. Paleozoic lycopod, 14. Triassic lycopod. All the plant reconstructions are not to scale. Reconstruction of *Lepidodendron*, *Lepacyclotes*, Paleozoic conifer, Paleozoic tree fern and *Calamite* are drawn by Huisu studio, *Tomiostrobus* reconstruction comes from Naugolnykh (2012), ginkgophytes is modified after Zhou (1990), gigantopterids

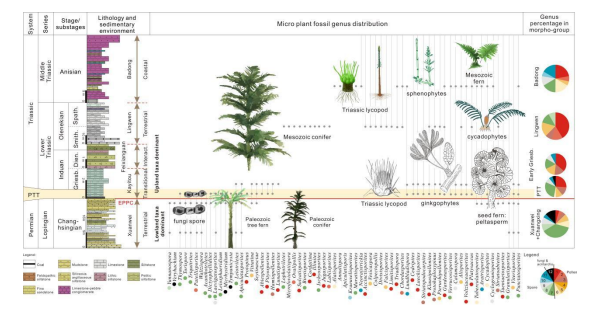


Figure 17. Lithology, sedimentary, micro plant fossil distribution range, floral composition from End Permian Changhsingian to Middle Triassic Anisian in South China area. Kayitou Formation conformably overlies on Xuanwei Formation, while Kayitou, Feixianguan, Lingwen and Badong formation does not directly connect with each other and are divided by dash line. Xuanwei Formation: terrestrial facies; Kayitou Formation: terrestrial marine transitional facies; Feixianguan Formation: terrestrial marine interacting facies; Lingwen Formation: terrestrial facies; Badong Formation: coastal facies. Legend of micro plant morpho group: 1. conifer, 2. gymnosperm, 3. peltasperm, 4. cordaites, 5. ginkgophyte, 6. cycadophyte, 7. fern, 8. fern or seed fern, 9. sphenophyte, 10. Paleozoic lycopod, 11. Triassic lycopod, 12. fungi or acritarchs spore. All the plant reconstructions are not to scale. Reconstruction of Middle Triassic lycopod, Paleozoic tree fern are drawn by Huisu studio, Early Triassic lycopod reconstruction comes from Naugolnykh (2012), ginkgophytes is modified after Zhou (1990), peltasperm cone comes from Naugolnykh (2000), Paleozoic conifer comes from Corey A. Ford, others come from Zhen Xu.

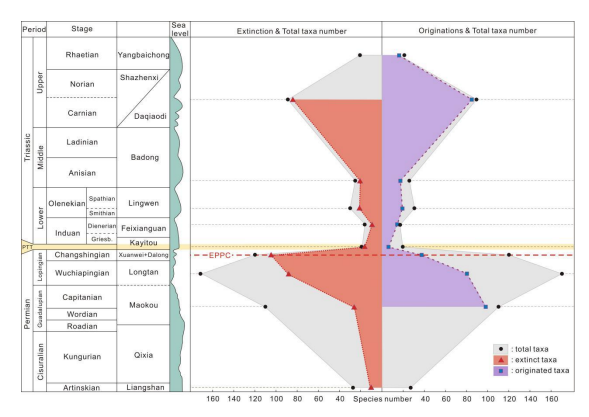


Figure 18. Diversity trends for fossil plant species from the Middle Permian Qixia Formation to the Late Triassic Yangbaichong Formation showing originations, extinction and total taxon number. PTT = Permo-Triassic transition; Griesb. = Griesbachian; EPPC = End Permian Plant Crisis.

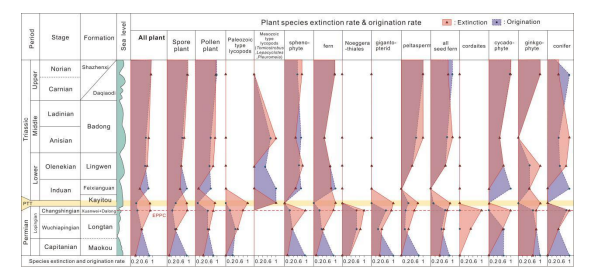


Figure 19. Extinction and origination rates from the Middle Permian Maokou Formation to the Late Triassic Dajing Formation showing origination and extinction rates for individual plant groups (PTT = Permo-Triassic transition; EPPC = End

Permian Plant Crisis; Red solid line and red area denote extinction rate; purple dashed line and blue–purple area denote origination rates).

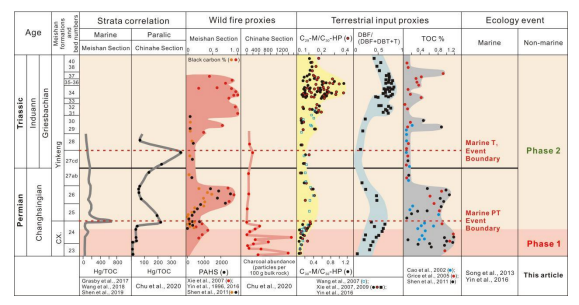


Figure 20. High-resolution comparison between terrestrial events and marine feedback from the End-Permian GSSP Meishan section Bed 23 to Early Triassic Bed 40. C. X. = Changhsingian; modified from Yin et al. (2016).

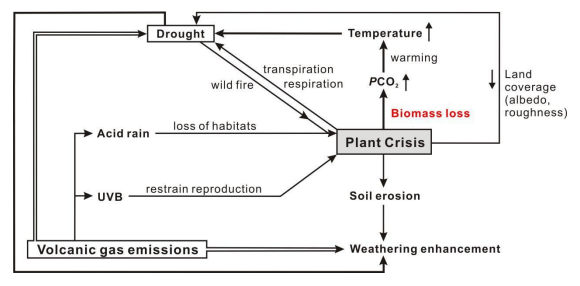


Figure 21. Hypotheses for the relationships between environmental changes and plant distribution emphasizing how changes contribute to episodes of plant species richness crisis, and in return causes environmental change.

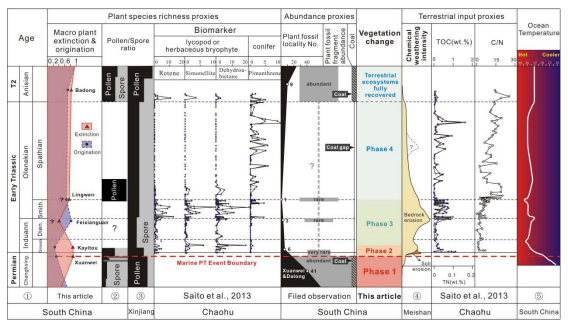


Figure 22. Comparison between floral change pattern proxies from macro plant fossils, coal, palynology, biomarker and environment event such as terrestrial input, marine feedback, marine temperature through the End-Permian Changhsingian to Middle Triassic Anisian. Changhsing. = Changhsingian; Griesb. = Griesbachian; Dien. = Dienerian; Smith. = Smithian; T2 = Middle Triassic. (1). Stratal data from Burgess et al. (2014); (2) Palynology data from Changhsingian to Induan of South China from Zhang et al. (2004), Yu et al. (2008), Ouyang et al. (2007), Olenekian Lingwen of South China from Zhang et al. (1992) and Middle Triassic Anisian of South Chin from Qu et al. (1990) and Meng et al. (1995); (3) Palynology data of Dalongkou section, Xinjiang Province, from Changhsingian to Anisian from Qu et al. (1986); (4) Chemical weathering rate from Algeo et al. (2011); (5) Oceanic temperature from Sun et al. (2012).

Formation	Stem	Root	Megasporophyll	Cone	Species richness	
	(Lepidodendron)	(Stigmaria)	(Lepidostrobophyllum)	(Lepidostrobus)	Non-normalised	Normalised
Xuanwei	L. acutangulium	S. ficoides	L. xiphidum		8	3
	L. lepidophylloides	S. rugulosa				
	L. oculus-felis	S. radiatopunctata				
		S. sp.				
Longtan	L. lepidophylloides	S. ficoides	L. caudatun	L. acutisquanmus	13	5
	L. polygonale	S. rugulosa	L. hastum			
	L. xuanweiense	S. sp.	L. junlianense			
	L. emeishanense		L. mucronatum			
	L. oculus-felis					
Maokou	L. asymetricum	S. ficoides	L. caudatun		5	2
	L. oculus-felis	S. sp.				
Qixia	L. asymetricum	S. ficoides			5	3
	L. oculus-felis	S. sp.				
	L. cf. szeianum					

2013 Table 1