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EARLY FLOWERS OF PRIMULOID ERICALES FROM THE LATE CRETACEOUS OF PORTUGAL AND THEIR ECOLOGICAL AND PHYTOGEOGRAPHIC IMPLICATIONS

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Abstract: A distinctive feature of the major eudicot diversification that occurred through the Late Cretaceous is the unequivocal presence of Cornales and diverse Ericales. Here we describe well-preserved fossil flowers from the Mira locality in western Portugal, of Campanian-Maastrichtian age, that we assign to a new extinct genus of Ericales with two new species; *Miranthus elegans* gen. et sp. nov. and *Miranthus kvacekii* sp. nov. The fossil flowers are pedicellate, structurally bisexual, actinomorphic, pentamerous and isomerous, with five narrowly triangular persistent calyx lobes, a five-lobed corolla, five antepetalous stamens, five staminodes alternating with the petals and a semi-inferior, unilocular ovary. The ovary consists of five carpels and has a raised nectariferous ring with stomata-like openings above the insertion of the perianth, and a long five-angled style. A key feature, which confirms a relationship with Primulaceae s. l., is the free, central dome-shaped placenta that bears numerous, densely spaced ovules. The ovary matures into a capsule containing many, minute, reticulate seeds. Flowers of *Miranthus* are especially similar to those of extant *Samolus*, a genus of about twelve species that is sister group to other genera of subfamily Theophrastoideae and that has a disjunct distribution mainly in the Southern Hemisphere. *Miranthus* also appears to have grown in environments influenced by marine conditions, an ecological preference also seen in *Samolus*. *Miranthus* expands the diversity of Ericales known from the Late Cretaceous, and together with previously described fossils provides further evidence that the diversification of Ericales was already underway by the Campanian-Maastrichtian stages of the Late Cretaceous.

Key words: Campanian, Ericales, fossil flowers, Maastrichtian, Primulaceae s. l., SRXTM, synchrotron radiation X-ray tomographic microscopy

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and ovule attachment, which allow the fossils to be placed in Late Cretaceous vegetation. The Ericales are known from more than 35 fossil species of flowers, fruits and seeds from Late Cretaceous sediments in Europe and North America (e.g., Friis 1985, Knobloch and Mai 1986, Nixon and Crepet 1993, Keller et al. 1996, Schönenberger and Friis 2001, Martinez-Millán et al. 2009, Friis et al. 2011, Schönenberger et al. 2012, Crepet et al. 2013, 2018). Some of these fossil species are based on fragmentary or limited material, such as fruits or isolated seeds, and while their relationship to Ericales appears well-founded, their systematic position within the order is not always clear. However, several highly informative flowers, as well as isolated seeds with characteristic morphology and seed coat structure, can be placed unequivocally in extant families (e.g., Parasaurauria Halloniensis J.A. Keller, Herend. et P.R. Crane, Actinidiaceae) and in some cases even in extant genera (e.g., Eurya crassitesta Erw. Knobloch, Pentaphylaceae). The available palaeobotanical data, taken as a whole, indicates that three of the major lineages of the Ericales; the Pentaphylaceae-Sladeniae clade, the Theaceae-Symphlocaceae-Styrracaceae-Diapetraceae clade and the Sarraceniaceae-Actinidiaceae-Roridulaceae-Clethraceae-Cyrrlidaeace-Ericaceae clade, were well-established and diverse in the Late Cretaceous, although we exclude the highly speculative report of a Sarracenia-like pitcher plant, Archaeamphora longicervar H.Q. Li, from the Early Cretaceous Yixian Formation of northeastern China (Li 2005), which was based on poorly preserved material.

In this paper, we expand the Late Cretaceous fossil record of Ericales with the description of a distinctive new genus and two new species based on minute coalified fossil flowers that can be assigned securely to the order. As with other coalified plant mesofossils (e.g., Friis et al. 2007, 2014, 2015), the application of synchrotron-radiation X-ray tomographic microscopy (SRXTM) reveals astonishingly fine details, including features of placentation and ovule attachment, which allow the fossils to be placed unequivocally among the Primulaceae s. 1.

Material and methods

The fossil flowers studied here were collected between 1989 and 1992 by EMF and KRP from the Mira locality (40°25′N, 08°44′15″W), an abandoned clay pit in the southern outskirts of the small town of Mira, about 25 km south of Aveiro, in the District of Coimbra, western Portugal. The pit is on the outskirts of the Mira locality contain diverse angiosperms (Friis et al. 2010), including several rosid taxa, of which three have been described formally. Esqueiria miraensis E.M.Friis, K.R.Pedersen et P.R.CRane (Friis et al. 1992) is of myrtalean (combreteaceous) affinity, while Normanthus miraensis Schönenb., K.R.Pedersen et E.M.Friis (Schönenberger et al. 2001) and Endressianthus miraensis E.M.Friis, K.R.Pedersen et Schönenb. (Friis et al. 2003) both belong to the Normapolles complex (Fagales). The Mira assemblage is also interesting for the presence of two monocot taxa, Spirematospermum sp. (Zingiberales) and seeds of Araceae (Friis et al. 2006a) as well as several undescribed monocot flowers.

Mesofossil assemblages from the Mira locality were isolated from the sediments and cleaned of adhering mineral matrix following standard techniques (see Friis et al. 2011, 2018). For SEM, specimens were mounted on aluminium stubs with nail polish, coated with gold (thickness not measured), and studied using Jehol JSM 840 (S101267, S101268) and Phillips 515 scanning electron microscopes (S100732, S101266) or a Hitachi S-4300 field emission microscope (S153144–S153146, S170106, S170123, S170125, S170155) at the Swedish Museum of Natural History.

Ten specimens (S100732, S153144–S153146, S156331, S170106, S170123, S170155, S170157, S266059) were studied using synchrotron-radiation X-ray tomographic microscopy (SRXTM) at the TOMCAT beamline, Swiss Light Source at the Paul Scherrer Institute, Villigen, Switzerland (Stampanoni et al. 2006). Specimens S153144–S153146 were analysed using phase contrast imaging at 20 keV. The other specimens were studied using absorption imaging at 10 keV. All specimens were analysed with rotation over 180° and most specimens were examined using a 10× objective with isotropic voxel size of 0.74 μm (S100732, S153144–S153146, S170106, S170123, S170155, S170157) or 0.65 μm (specimens S15633, S170106, S266059). Specimens S170123 and S170106 were studied using 20× objective with isotropic voxel size of 0.37 μm (S170123) or 0.325 μm (S170106). For details of SRXTM work on Cretaceous plant mesofossils at TOMCAT see Friis et al. (2014). Data derived from the SRXTM scans were analysed and manipulated using Avizo software (9.1.1-9.5; Thermo Scientific, Waltham, MA, USA) for computed tomography. Photoshop CS5 Adobe (San Jose, CA, USA) was used to create an even black background for SEM images and to label SRXTM images.

An assessment of the phylogenetic position of the fossils was performed using Mesquite software version 3.2 (Maddison and Maddison 2017) using the Parsimony Ancestral State feature with all characters treated as unordered. Taxa and morphological character states were selected from the comprehensive study of relationships among extant primuloids by Anderberg and Stähl (1995).
based on morphology, and include representatives from the immediate sister groups to Primulaceae (Ebenaceae: Diospyros spp., Sapotaceae: Manilkara zapota (L.) P.ROYEN) as well as genera selected to represent the four subfamilies of the Primulaceae (Maesoideae: Maesa subg. Maesa, Theophrastoideae: Samolus valeradi L., Theophrasta americana L., Clavija spp., Primuloideae: Douglasia nivalis Lindl., Soldanella alpina L., Primula veris L., Cortusa mathioli L., Myrsinoideae: Cyclamen coum MILLER, Lysimachia glutinoso ROCK, Ardisiandra sibthorpioides Hook.f., Anagallis monellii L., Myrsine africana L.). A backbone tree was constructed for these taxa based on topologies supported by molecular data for Eriques (Rose et al. 2018) and for Primulaceae (Morozowska et al. 2020). The number of character state changes for each potential position of Miranthus was recorded by linking the fossils manually to all possible branches of the backbone tree.

All specimens are deposited in the palaeobotanical collections of the Swedish Museum of Natural History, Stockholm (S). Raw data from the SRXTM analyses are available at the PSI Public Data Repository, Paul Scherrer Institute, Villigen, Switzerland (Friis et al. 2021). The names assigned to the new genus and species described here, are each registered with a unique registry number in the Plant Fossil Names Registry, which is hosted and operated by the National Museum, Prague for the International Organisation of Palaeobotany (IOP).

Characters scored for the phylogenetic analyses

Characters scored for extant primuloid flowers and fossils of Miranthus are based on the morphological dataset (characters 24–82) of Anderberg and Ståhl (1995). Thirty characters were scored for the fossil flowers. 25: Pedicel (0) without paired bracteoles. 26: Flower (1) perigynous. 27: Flowers (0) homostylous. 28: Flowers (1) bisexual. 29: Flowers (2) haplostemonous with the functional stamens opposite the corolla lobes. 30: Perianth (0) 5-merous. 33: Corolla (1) with staminodes. 34: Corolla aestivation (0) imbricate or quincuncial. 37: Corolla texture (1) thin. 38: Sepals (1) united, often cup-shaped. 40: Petals (0) entire. 41: Petals (0) longer than sepals or subequal. 42: Petals (1) lanceolate-ovate, more or less acute, and usually distinctly longer than wide. 45: Flower (3) with somatic nectaries. 46: Anther filaments (0) without long sepal trichomes. 47: Anther filaments (0) free from each other. 49: Anther filaments (0) eglanular. 57: Anthers (0) not forming a cone. 61: Pollen (0) 3-, 4-, or 5-coltorpe, or colporoidate. 62: Style (0) included, shorter than or slightly longer than the anthers. 64: Style (1) entire. 66: Fruit (2) a capsule. 67: Fruit (0) normally many-seeded. 68: Fruit opening (1) with teeth or valves. 69: Fruit (1) globose to subglobose. 70: Placenta (0) normally many-seeded. 72: Ovules (1) bitegmic. 73: Ovules (0) not immersed in the placenta. 74. Ovules (0) in two or more series, usually many.

Systematic palaeobotany

Subdivision Angiospermae Lindl., 1830
Order Ericales Bercht. et J.PRESL., 1820
Family Primulaceae Batsch ex Borkh., 1774 nom. cons.
Text-fig. 1. SEM images of flowers of *Miranthus elegans* gen. et sp. nov.; Mira locality, Portugal. a, b: Flowers in lateral view showing elongated pedicel, narrowly triangular sepals and elongated protruding style (a); note the large openings in the floral tissue and pedicel (asterisks) interpreted as schizogenous secretory cavities. c: Flower in lateral view with portion of the calyx missing exposing the ovary wall and slightly raised nectariferous ring with probable stomata-like secretory structures (arrow). d: Flower in lateral view showing long pedicel and three of the five tepals; note the elongated narrowly triangular form of the sepals. e: Flower in oblique lateral view with portion of the calyx missing exposing the ovary and elongated style. f, g: Flowers in apical view showing the bases of five sepals (f) and apex of the five-parted ovary; note larger openings in the floral tissue (asterisk) interpreted as schizogenous secretory cavities. Specimens, Mira 100-S170155 (a, holotype), Mira 100-S153145 (b, c, g), Mira 100-S101267 (d), Mira 105-S100732 (e), Mira 100-S101268 (f). Scale bars = 600 µm (a–g).
Text-fig. 2. SEM images of *Miranthus elegans* gen. et sp. nov.; Mira locality, Portugal. a, b: Flowers in oblique lateral view showing remains of calyx and slightly semi-inferior ovary with elongated apical style (a); note larger openings in the floral tissue (asterisk) interpreted as schizogenous secretory cavities and the stomata-like secretory structures on the upper portion of the ovary (arrows) that are interpreted as nectariferous (b). c: Detail of ovary surface showing secretory stomata-like structures (arrows). d: Flower in lateral view showing fragmentary calyx and broken slightly semi-inferior ovary with secretory stomata-like structures; note the point of attachment of the central placenta (pl). e: Cluster of seeds removed from the ovary in (d) showing reticulate surface. f: Outer (abaxial) surface of calyx lobe showing the slightly pointed papillae and scattered, fine trichomes (arrows). g: Triaperturate pollen grains from the ovary surface. Specimens, Mira 100-S153146 (a, b), Mira 100-S1700155 (c), Mira 100-S101266 (d, e), Mira 105-S100732 (f), Mira 100-S170125 (g). Scale bars = 600 μm (a, b, d), 300 μm (f), 100 μm (c, e), 10 μm (g).
Text-fig. 3. SRXTM images of *Miranthus elegans* gen. et sp. nov.; Mira locality, Portugal. a, b: Volume renderings of flower bud in two different lateral views showing long pedicel, distinct calyx (ca) with almost equiaxial epidermal cells and corolla (co) with nearly smooth surface. c–e: Transverse sections (c, orthoslice xy1500; d, orthoslice xy1760; e, orthoslice xy1850) through flower bud at levels below the anthers showing stamen filaments (yellow) opposite the corolla lobes (co) and smaller staminodes (orange).
Type stratum and age. “Argilas de Vagos”, Late Cretaceous (Campanian/Maastrichtian).

Diagnosis. As for the genus with the following addition. Epidermis of pedicel and outer (abaxial) epidermis of calyx composed of cells that each have a central pointed papilla. Trichomes scattered and thin.

Dimensions. Length of flower without pedicel and corolla: 0.8–1.0 mm; diameter: 0.7–1.0 mm. Length of flower bud with corolla and without pedicel: 1.5 mm.

Description and comments on the species. The fossil flowers were briefly described and illustrated by Friis et al. (2010, 2011), but not formally named. More material has subsequently been discovered, including a small, informative flower bud. In most flowers the corolla and androecium have been shed (Text-figs 1a–g, 2a–d) and the flowers are probably fossilized at a post-anthetic stage. The flower bud has remains of corolla and stamens preserved (Text-fig. 3a–f), but unfortunately, internal structures, including the ovary, ovules and parts of the androecium, are not well preserved. These internal structures were partially destroyed during fossilization and are only preserved on one side of the flower, which complicates interpretations of their organisation. The flower bud is assigned to the same species as the post-anthetic flowers based on features of the calyx. A second flower bud from the Mira mesofossil flora is described as a separate species, Miranthus kvacekii, based on epidermal differences from Miranthus elegans.

The flowers are borne on a long, slender pedicel (Text-figs 1a–d, 2d, 3a, b) and lack bracts or bracteoles immediately below the calyx. The pedicel is up to about 0.9 mm long (Text-fig. 1d). All of the fossil flowers are isolated and there is no information on inflorescence structure, or how the pedicel may have been attached to the plant.

The flowers are structurally bisexual, actinomorphic, pentameric and isomerous, about 0.8–1.0 mm long, excluding the pedicel and corolla, and 0.7–1.0 mm in diameter. Including the corolla, but excluding the pedicel, the flower bud is about 1.5 mm long. The calyx is persistent and consists of five imbricate sepals in a single whorl that are fused at the base (Text-figs 1a–g, 5a, b). The sepals are narrowly triangular, with the free portion up to about 0.9 mm long, which tapers into a long tip (Text-fig. 1a–g). Each sepal is supplied by three distinct vascular bundles, each of which typically appears as a hollow space in transverse sections (Text-figs 5a, 6a, b). In some specimens, minor vascular bundles are seen close to the sepal margins. The epidermis of the pedicel, and the abaxial (outer) epidermis of the calyx, is covered by a thick cuticle. The epidermal cells are small, almost equiavial, thick-walled and typically with a pointed papilla that gives the surface a faint verrucate to spiny appearance. Similar epidermal features are also observed on the outer surface of the young ovaries (Text-figs 1a–g, 2f, 3c–e, 5a–d, 6a). Very slender trichomes occur scattered on the outer surface of the sepals (Text-fig. 2f). The epidermal cells of the adaxial (inner) epidermis of the calyx lobes are small, equiavial, thin-walled and covered with a thin cuticle (Text-fig. 3c–e). The calyx of the single flower bud is very similar in shape, size and epidermal features to the calyx post-anthetic flowers (Text-fig. 3a, b) and the anatomy of all preserved organs is also identical.

There is no trace of corolla and androecium in the supposed post-anthetic flowers. Corolla and androecium were most likely shed together after flowering, which would also be consistent with stamens that were fused to the corolla. The corolla is five-lobed, with imbricate lobes that are free for most of their length (Text-fig. 3c–e). They appear united at the base, but the preservation does not allow a fully secure conclusion that they were sympetalous.

The internal structures are preserved on one side of the flower bud, but based on the symmetry of the structures that remain, the androecium can be reconstructed as pentamorous and obhapolostemonous with five antepetalous stamens, which are seen near the base of the flower as stout filaments positioned in front of the petal lobes (Text-fig. 3c–e) and near the bud apex as remains of thecae. Smaller structures, interpreted as staminodes, are seen alternating with the petals (Text-fig. 3c–e). Both stamens and staminodes appear to be attached to the corolla near the base (Text-fig. 3f), which is also consistent with the absence of stamens and stamen bases in the post-anthetic flowers.

Normalpolls-type pollen, which is very common on the surface of other Mira mesofossils, is also present on Miranthus flowers. However, in addition, two flowers have another kind of pollen on their surface. In both cases, the pollen is about 16 µm long and triaperturate with very long colpi that reach almost to the poles (Text-fig. 2g). Unfortunately, the grains are folded and it cannot be established whether the grains are tricolpate or tricolporate. The tectum is foveolate.

The ovary is semi-inferior and unilocular (Text-figs 2d, 4a–d, 5a, b, 6a, b, e). The apical portion of the ovary has a slightly protruding ring with stomata-like openings (Text-figs 1c, 2b, c). We think it likely that this zone was nectariferous and that the openings were nectar secreting. In some specimens there appear to be remains of sepals in the apical part of the ovary, but these may be invaginations of the ovary wall and they do not reach the centre of the ovary (Text-fig. 5a). The style is long, slender (Text-figs 1e, 4c), and distinctly five-angled in cross-section with a vascular bundle that extends for the length of the style in each of the angles (Text-fig. 6c, d). The style is hollow and the stylar canal is wide along its full length (Text-fig. 6d), except near the tip where it is closed (Text-fig. 6c). In some specimens, the style is broken, but the specimens available give no indication of different style lengths and the flower is interpreted as homostylous. Placentation is free and central, with a dome-shaped, almost globose placenta, alternating with the corolla lobes; note verrucate abaxial surface of calyx lobes (ca) and ovary wall (ow) partly distorted and displaced to one side. f: Longitudinal section (orthoslice yz0567) of flower bud showing ovary wall (ow) and insertion of calyx (ca), corolla (co) and stamens (st) on a hypanthism rim. Specimen, Mira 100-5266059 (a–f). Scale bars = 600 µm (a, b), 300 µm (c–f).
borne on a central column (Text-figs 4a–d, 5a–b, 6a, b, e). Numerous ovules are densely spaced on the placenta, but not immersed in its surface (Text-figs 2e, 4a–d, 5a–d, 6b, e). Ovules are bitegmic and anatropous, about 0.14 mm long, with a reticulate seed surface (Text-fig. 2e).

Among the post-anthetic flowers, none has a fully developed fruit. However, some are clearly five-parted, have five valves and are interpreted as young capsules (Text-fig. 1e, f). In one specimen the ovary is split open along one of the valves (Text-fig. 2d). The epidermal cells of the young capsules are small, equiauxial, and often with tiny papillate projections. Stomata occur scattered over the surface of the calyx lobes and pedicel together with smaller openings that are probably trichome bases. Larger openings are irregularly scattered and are interpreted as schizogenous secretory cavities that are sometimes burst (Text-figs 1b, c, 4b).
**Miranthus kvacekii** E.M.Fris, P.R.Crane et K.R.Pedersen sp. nov.

Text-fig. 7a–e

**Holotype.** S170157 (Mira sample 100; figured Text-fig. 7a–e).

**Plant Fossil Names Registry Number.** PFN002673 (for new species).

**Repository.** Palaeobotanical Collections, Swedish Museum of Natural History, Stockholm, Sweden (S).

**Etymology.** In commemoration of Zlatko Kvaček and his outstanding contributions to Cretaceous and Cainozoic angiosperm palaeobotany.

**Type locality.** Mira (40° 25’ N; 08° 44’ 15ʺ W), about 25 km south of Aveiro, District of Coimbra, Portugal.

**Type stratum and age.** “Argilas de Vagos”, Late Cretaceous (Campanian/Maastrichtian).

**Diagnosis.** As for the genus with the following addition: Epidermis of pedicel and outer (abaxial) epidermis of calyx with closely spaced, robust, stiff trichomes.

**Dimensions.** Length of flower excluding pedicel and corolla: 0.8–1.0 mm; diameter: 0.7–1.0 mm. Length of flower bud with corolla, but excluding the pedicel: 1.5 mm.

**Description and comments on the species.** The species is based on a single flower bud.
discovered with *Miranthus elegans* in the Mira mesofossil flora (Text-fig. 7a–e). The individual floral parts are strongly appressed and somewhat distorted, which impedes a full understanding of the floral structure and organization (Text-fig. 7b–e). However, the androecium clearly consists of five stamens. The stamens also apparently alternate with the sepals and are obhaplostemonous (Text-fig. 7d, e). The stamen filaments appear to be fused to the corolla for part of their length, while the anthers are free from the corolla and from each other. Alternating with the filaments are filament-like structures. These are not as distinct as the stamen filaments and interpreted as staminodes (Text-fig. 7e). The flower bud is similar to *Miranthus elegans* in its characteristic free, dome-shaped placenta (Text-fig. 7b, c) and semi-inferior ovary (Text-fig. 7b, c), but differs in having short, densely spaced trichomes on the outer surface of the calyx lobes, particularly along the margins (Text-fig. 7a). Although scattered trichomes and trichome basis are observed in *Miranthus elegans*, the trichomes are much finer and less densely spaced.

**Five-carpellate capsule assigned to Miranthus**

Text-fig. 8a–d

Description. A single five-carpellate capsular fruit (S156331) was discovered with the *Miranthus* flowers in the Mira flora (Text-fig. 8a–d). The fruit is subglobose, about 1.6 mm in diameter with remains of the perianth near the base. The fruit contains many angular seeds, up to about 0.8 mm long with a bitegmic seed coat with a thin testa and a membranous tegmen (Text-fig. 8c). The placenta is central, free and borne on a central column (Text-fig. 8c, d). The seed surface is reticulate with lumina arranged in longitudinal rows and delimited by slightly raised periclinal walls (Text-fig. 8a, b), similar to those of the ovules in
Text-fig. 7. SEM (a) and SRXTM (b–e) images of *Miranthus kvacekii* sp. nov.; Mira locality, Portugal. a: Lateral view of flower bud showing corolla lobes extending beyond calyx; note surface of pedicel, calyx and corolla with small equiaxial epidermal cells and indumentum of densely spaced, short stiff trichomes. b, c: Longitudinal sections through floral bud in two directions perpendicular to each other (a, orthoslice yz1024; b, orthoslice xz0950) showing corolla (co), calyx (ca), stamens (st) and semi-inferior ovary with thin ovary wall (ow) and central mushroom-shaped globose placenta (pl) bearing numerous ovules (ov). d, e: Transverse sections through floral bud above placenta (d, orthoslice xy0915; e, orthoslice xy1095) showing calyx (ca), corolla (co), ovary wall (ow) and ovules (ov); yellow outlines indicate the positions of anthers (d) and filaments (e); orange outlines indicate the position of three of the possible staminodes. Specimen, Mira 100-S170157 (a–e, holotype). Scale bars = 600 µm (a–c), 300 µm (d, e).
flowers of *Miranthus*. Because of this similarity and because *Miranthus* is the only flower in the Mira mesofossil flora with five carpels, the capsule is interpreted as a mature fruit of either *Miranthus elegans* or *M. kvacekii*. Unfortunately, details of the epidermis of the calyx are not known, which precludes assignment at the species level.

**Discussion: Systematic relationships of *Miranthus***

Characters of the *Miranthus* flowers clearly place the fossil taxon in the primuloid clade, which is firmly nested among the Ericales (e.g., Anderberg et al. 2002, Schönenerberger et al. 2005). In most classifications the primuloid clade comprises four groups recognised either as families of the order Primulales (Maesaceae, Theophrastaceae, Primulaceae, Myrsinaceae; Anderberg and Ståhl 1995, Anderberg et al. 1998a, 2000a, Källersjö et al. 2000, Schönenerberger et al. 2005) or as subfamilies of the family Primulaceae (Maesoideae, Theophrastoideae, Primuloideae, Myrsinoideae; APGIII 2009). A fifth, monogeneric lineage, Samolaceae/Samoloideae, is now usually treated as a tribe, Samoleae, within the Theophrastaceae/Theophrastoideae group (e.g., Källersjö et al. 2000, Caris and Smets 2004). We here use the inclusive definition of Primulaceae s.l.

Extensive studies on extant primuloids by Anderberg and colleagues (e.g., Anderberg and Ståhl 1995, Anderberg et al. 1998a, b, 2000a, b, 2007, Källersjö et al. 2000, Källersjö and Ståhl 2003, Wannop and Anderberg 2011), as well as by other researchers (e.g., Caris et al. 2000, Ma and Saunders 2003, Caris and Smets 2004, Morozowska et al. 2020), have been pivotal for our comparative studies and for recognizing the affinities of the fossil flowers. Relationships among
the main primuloid lineages are well supported based on phylogenetic analyses of both morphological and molecular data. The monogeneric Maesoideae is resolved as the sister lineage to the remaining taxa, with Theophrastoideae resolved as sister to a well-supported Primuloideae-Myrsinoideae clade (Källersjö et al. 2000, Bremer et al. 2002, Schönenberger et al. 2005). Within Theophrastoideae, the monogeneric Samoleae are sister to all other members of the subfamily.

Shared features for *Miranthus* and the Primulaceae include the pentamerous and isomerous organisation of the flowers, persistent calyx, haplostemonous androecium and the highly characteristic free central and dome-shaped placenta. Among Ericales, a free central placenta is known only for the Primulaceae and is a clear synapomorphy for the group (Schönenberger et al. 2005).

Few floral features distinguish the various primuloid lineages and a separation of the lineages based on floral morphology is only poorly supported. As noted by Caris and Smets (2004) “it seems that no unambiguous morphological synapomorphies can be found that justify a position of *Samolus* in either Primulaceae or Theophrastaceae”, but a position as sister to Theophrastoideae is well supported by molecular data (Källersjö and Ståhl 2003). Nevertheless, the semi-inferior ovary of *Miranthus* flowers, and the presence of staminodes alternating with the corolla lobes, are important distinctive floral features. Furthermore, among extant primuloids, a semi-inferior ovary is known only for *Maesa* Forssk. and *Samolus* L., and staminodes are present only in *Maesa*, the genera of Theophrastoideae (including *Samolus*), and *Soldanella* L. of the Primuloideae (Anderberg and Ståhl 1995, Anderberg et al. 1998a). A nectariferous zone on the upper part of the ovary, referred to as “nectarostomata on the flanks of the ovary” by Caris and Smets (2004), is a further feature shared with *Maesa* and *Samolus* that is not present in other primuloid taxa (Caris et al. 2000, Caris and Smets 2004). *Miranthus* thus appears to be nested securely in the primuloid clade.

*Maesa* comprises about 150 species of tropical and subtropical vines, shrubs, and trees in the Old World (Anderberg et al. 2000a), whereas species of *Samolus* are herbaceous and less diverse with about twelve species. Four or five species of *Samolus* occur in the southern United States, Cuba and Mexico, and one species (*Samolus valerandi* L.) is cosmopolitan in distribution. The remaining species are restricted to the Southern Hemisphere, with three species in South America, four in Australia and Tasmania, and one in South Africa (Ståhl 2004a, Wanntorp and Anderberg 2011). *Maesa* is distinguished from *Miranthus* and all other primuloid taxa by the presence of two bracteoles immediately under the calyx. *Miranthus* and other primuloids display the likely plesiomorphic condition in which such bracteoles are lacking (Anderberg et al. 2000a). *Miranthus* is also more like *Samolus* in having a long and slender pedicel.

The close relationship of *Miranthus* and *Samolus* suggested by their morphological similarities is supported by the results of the phylogenetic analysis (Text-fig. 9). The most parsimonious position of *Miranthus* is as sister to *Samolus* in the Theophrastoideae (131 steps). However, a sister group relationship to Theophrastoideae, or a position as sister to *Clavija Ruiz et Pav.* + *Theophrasta L.*, is only one step longer (132 steps), while a sister group relationship...
Conclusions: phytogeography and ecology

The subfamily Theophrastoideae includes mainly shrubs and small trees distributed from northern Mexico and southern Florida to southern Brazil and northern Paraguay, with *Samolus* as the principal exception (see above and Ståhl 2004b, Wanntorp and Anderberg 2011). Based on a fossil calibrated phylogenetic analysis Rose et al. (2018) suggested that Theophrastoideae originated about 70 million years ago in the Neotropics, and that occurrences outside the Neotropics, including the distribution of *Samolus*, reflect subsequent and probably relatively recent migration. The discovery of *Miranthus* from Portugal in the Late Cretaceous, combined with the disjunct distribution of *Samolus* in the Southern Hemisphere, suggests a more complex biogeographic history, which also recalls the likely close relationship between the Late Cretaceous fossil flower *Silvianthemum* E.M.Friis from southern Sweden and the Southern Hemisphere genus *Quintinia* A.DC. (Friis 1990, Friis et al. 2013). Such results suggest the need for caution in interpreting the origin of present-day phytogeographic patterns without the benefit of palaeobotanical data. *Miranthus*, and other fossils imply significant extinction in the early history of Ericales during the Late Cretaceous. The role that such extinctions may have played in the origin of present day biogeographic patterns deserves more detailed consideration.

A probable close relationship between *Miranthus* and *Samolus* is also of ecological interest. Most species of *Samolus* inhabit wetlands, and many grow near the seashore in salt marshes or brackish ponds (Ståhl 2004a). The quality of preservation of *Miranthus*, including little damaged, delicate floral structures, suggests that the fossils were deposited close to the environment in which they lived. The flowers of *Miranthus* were discovered in sediments with clear marine influence. Like *Samolus*, *Miranthus* may have been growing in coastal plains or estuaries partially influenced by marine conditions. *Miranthus* thus provides an example not only of relative morphological stasis, but may also indicate the long-term persistence of a particular habitat preference established early in the diversification of Ericales.

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