ABSTRACT

This study reports the results of an examination of about 1,500 fossil leaf impressions from the Early Cretaceous (Albian) strata at the Hoisington III locality, Kansas, USA. We recognize 24 fossil angiosperm leaf species, of which seven are assignable to the modern orders Illiciales (1), Laurales (4) and Magnoliales (2). Two species show close relationship with modern family Nymphaeaceae. Three species are possibly related to the Chloranthaceae. One species shows affinity with the Nelumbonaceae. Three species are closely related to the Platanaceae. One new genus, Wingia gen. nov., is established. A total of seven new species and new combinations are proposed. They include: Longstrethia aspera (Lesquereux) comb. nov., Sapindopsis powelliana (Lesquereux) comb. nov., Anisodromum schimperi (Lesquereux) comb. nov., Wingia expansolobum (Upchurch and Dilcher) comb. nov., Sapindopsis retallackii sp. nov., cf. Anisodromum upchurchii sp. nov., and Dicotylophyllum skogii sp. nov. Comparisons with the angiosperm leaf assemblages from other localities indicate that the Hoisington locality represents the most species-rich leaf assemblage from the Dakota Formation along the east side of the Cretaceous Western Interior Seaway (KWIS) and equivalent units. The result provides new information for understanding the early diversity and evolution of the angiosperms during the mid-Cretaceous (late Albian – early Cenomanian), a critical time during which angiosperms began a rapid adaptive radiation.

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INTRODUCTION

Angiosperm leaves are abundant in the Dakota Formation and its equivalent units along the eastern margin of the Cretaceous Western Interior Seaway. These fossils have been studied since the late 19th century (Lesquereux, 1868, 1872, 1873, 1874, 1876a, 1876b, 1878, 1883, 1892; Newberry, 1868, 1895, 1898; Berry, 1911a, 1911b, 1911c, 1911d, 1916, 1920, 1922a, 1922b, 1923). However, these early publications are biased against mudstone and shale specimens because the specimens studied were primarily collected from various sandstone units of the Dakota Formation. More recent works (Upchurch and Dilcher, 1990; Wang and Dilcher, 2006a, 2009) have turned attention to the fossil plant material from the mudstone and shale units of the Dakota Formation collected by David Dilcher, colleagues, and students since the 1960s. The extensive collection of angiosperm leaf megafossils from many localities in Iowa, Kansas, Nebraska, and Minnesota during the past fifty-five years enables the comprehensive survey of the mid-Cretaceous Dakota Formation angiosperms. In this paper, we focus on the systematics of angiosperm leaves from the Lower Cretaceous (upper Albian) Dakota Formation at the Hoisington III locality, Kansas (Figure 1). Many publications have been generated based upon material from this locality (Dilcher, 1979; Crane and Dilcher, 1984; Dilcher and Crane, 1984; Dilcher and Kovach, 1986; Skog et al., 1992; Skog and Hill, 1992; Skog and Dilcher, 1994; Dilcher, 2000). In this study, we examined approximately 1,500 specimens and report 24 fossil species of angiosperm leaves. This research represents the third in a series of detailed studies of the Dakota flora from localities in Kansas, Nebraska, Iowa, and Minnesota by the authors (Wang and Dilcher, 2006a, 2009).

MATERIAL AND METHODS

All specimens discussed in this paper were collected from the Hoisington III locality (UF15706), Barton County, Kansas (Figure 1). The clay pit is about 3.2 kilometers south of Hoisington on State Route 281 at latitude 38°28.31' N and longitude 98°46.92' W. Specimens were collected from the claystone overlying interbedded sandstone and shales of the Terra Cotta Clay Member of the Dakota Formation (Figure 2) that outcropped at the south end of the north clay pit (Figure 3; also see Retallack and Dilcher, 1981b, text-figures 6-8 and Retallack and Dilcher, 2012, figure 2) by David Dilcher and colleagues Greg Retallack, Karl Longstreth, Carolyn Bagley, and Frank Potter during the field excursions in 1978 and 1979. Specimens have been assigned the locality number UF15706 and are deposited in the Paleobotany and Palynology collection at the Florida Museum of Natural History (FLMNH), University of Florida, Gainesville, Florida, USA. The locality number is followed by a specimen number (for example, UF15706-8263).

Most specimens are impressions except for a few compressions with poorly preserved cuticle. Descriptions are based on direct observation under a dissecting microscope. Details were enhanced in photographs by using a Nikon D200 digital camera under extreme oblique lighting to highlight the venation patterns.

Farley and Dilcher (1986) interpreted the age of the plant fossil-bearing strata at the Hoisington III locality as Cenomanian (early Late Cretaceous). However, based upon results of palynostratigraphic and sedimentologic analyses, Brenner et al. (2000)
proposed that the age of this sequence at this locality is late Albian (latest Early Cretaceous). Retallack and Dilcher (1981a, 1981b) interpreted the sedimentary environment at this locality as a brackish water lagoon or fresh water lake with river influence. The interpretation of a predominately fresh water lake was confirmed by the presence of aquatic ferns (Skog et al., 1992; Skog and Hill, 1992; Skog and Dilcher, 1994) and aquatic angiosperms (Wang and Dilcher, 2006b).

We follow Upchurch and Dilcher's (1990) scheme for specific and generic classification and the scheme of Angiosperm Phylogeny Group (2009, 2016) for familial and ordinal classification. Comparisons with other fossil taxa are based primarily on published figures, except most of those published by Lesquereux (1892) which were examined at the Paleobiology Department of the National Museum of Natural History, Smithsonian Institution, Washington D.C., USA. All the type specimens from the Rose Creek locality (Upchurch and Dilcher, 1990), Braun Ranch locality (Wang and Dilcher, 2006a), and Courtland I locality, Minnesota (Wang and Dilcher, 2009) were examined at the Paleobotany and Palynology collection in the Florida Museum of Natural History. Comparisons with extant taxa are based on published illustrations and direct observation on modern leaf collections at the Paleobotany and Palynology collection at the Florida Museum of Natural History.

In the synonym list, we follow Matthews (1973, p. 718) and use the following two symbols to indicate the degree of confidence with which particular items (specimens) in the list are referred to the taxon under discussion: "*" in front of the year indicates that with the publication of this paper, the species can be regarded as valid under the terms of the ICBN; "v." in front of the year indicates that we have checked the deposited specimens that relate to the work cited and because of the evidence of the deposited specimens we are able to take responsibility for this assignment.

**FIGURE 2.** Chart shows the Albian, Cenomanian and Turonian stratigraphic units in Kansas, the Tri-state area (Iowa, Nebraska, and South Dakota; modified from figure 4 of Brenner et al., 2000), and Minnesota (modified from figure 8 of Setterholm, 1994). 1, Courtland I clay pit, Minnesota; 2, Rose Creek, Nebraska; 3, Braun Ranch, Kansas; and 4, Hoisington III, Kansas.

**FIGURE 3.** Cross section of the plant-bearing layer at the southwest end of the north clay pit at the Hoisington III locality, Kansas (Modified from figure 2 of Retallack and Dilcher, 2012).
Leaf architectural terminology mostly follows Dilcher (1974), Hickey (1973, 1979), Wolfe et al. (1975), and Ellis et al. (2009) with the following exceptions. We use the term leaf and/or leaflet when describing specimens only if the specimens demonstrate enough features to show leaf organization (i.e., simple or compound). Otherwise we use the term lamina in our descriptions. We follow Upchurch and Dilcher (1990) and Upchurch et al. (1994) by using “cf.” to indicate minor differences between our specimens and a previously published taxon and a fossil taxon’s similarities to modern families. The following abbreviations are used: L/W represents the ratio of leaf, leaflet, or lamina length.

**TABLE 1.** Occurrences of angiosperm leaf megafossils at the four localities of the Dakota Formation, Braun’s ranch, Kansas (Wang and Dilcher 2006a), Hoisington, Kansas, Rose Creek, Nebraska (Upchurch and Dilcher 1990), and Courtland, Minnesota (Wang and Dilcher 2009). A plus sign “+” indicates the presence of a taxon at a locality.

| Taxon Genus/species | Hoisington | Braun Ranch | Rose Creek | Courtland |
|---------------------|------------|-------------|------------|-----------|
| **Crassidenticulum decurrens** | + | + | + |   |
| **Crassidenticulum cracendentis** |   |   |   | + |
| **Crassidenticulum trilobum** | + |   | + |   |
| **Crassidenticulum cf. trilobum** | + |   |   |   |
| **Crassidenticulum landisiae** |   |   | + |   |
| **Yangia glandifolia** |   |   | + |   |
| **Densinervum kaulii** |   | + | + |   |
| **Landonia calophylla** |   |   | + |   |
| **Landonia callii** |   |   | + |   |
| **Aquatifolia fluitans** | + |   |   |   |
| **Brasenites kansense** | + |   |   |   |
| **Longstrethia varidentata** |   |   | + |   |
| **Longstrethia aspera** |   |   | + |   |
| **Pabiania variloba** | + |   |   | + |
| **Setterholmia rotundifolia** |   |   | + |   |
| **Setterholmia deleta** |   |   | + |   |
| **Manchesteria macrophylla** |   |   | + |   |
| **Pandemophyllum kvacekii** | + |   | + |   |
| **Pandemophyllum attenuatum** | + |   | + |   |
| **Pandemophyllum sp.** | + |   |   | + |
| **Rogersia dakotensis** | + |   | + |   |
| **Rogersia parlatorii** | + |   | + |   |
| **Rogersia lottii** |   |   | + |   |
| **Wolfiophyllum daphneoides** | + |   |   |   |
| **Wolfiophyllum heigii** |   |   | + |   |
| **Wolfiophyllum pfaffianum** | + |   | + |   |
| **Jarzenia kanbrasota** | + |   | + |   |
| **New genus A** |   |   | + |   |
| **Liriophyllum siemii** | + |   |   |   |
| **Liriophyllum kansense** |   |   | + |   |
| **cf. Paleonelumbo macroloba** | + |   |   |   |
| **Credneria quadrata** |   |   | + |   |
| **Credneria cyclophylla** | + |   |   | + |
| **Dischidus quinquelobus** |   |   | + |   |
| **Eoplataurus serrata** |   |   | + |   |
to width. When describing the angle of origin of tertiary veins, we use A, R, and O to represent acute, right, and obtuse angles respectively (Dilcher, 1974). For example, AO represents tertiary veins originated from the exmedial (lower or proximal) side of the secondary veins at acute angles (A) and at obtuse angles (O) from the admedial (upper or distal) side of the secondary veins.

We only provide diagnoses for newly established taxa in this report. For new occurrences of previously published taxa we only provide descriptions. Table 1 is a list of taxa reported from the Braun (Wang and Dilcher, 2006a) and Hoisington
III localities, Kansas (this paper), Courtland I locality, Minnesota (Wang and Dilcher, 2009), and Rose Creek locality, Nebraska (Upchurch and Dilcher, 1990).

**SYSTEMATIC PALEOBOTANY**

Family cf. CHLORANTHACEAE R. Br. ex Sims, 1820

Genus CRASSIDENTICULUM Upchurch and Dilcher, 1990

Type species. *Crassidenticum decurrens* (Lesquereux) Upchurch and Dilcher, 1990

*Crassidenticum decurrens* (Lesquereux)
Upchurch and Dilcher, 1990

Figure 4.1

*1892 Celastrophyllum decurrens Lesquereux, p. 172, plate 36, figure 1.

**Description.** Leaflet margin toothed. Primary venation pinnate; primary vein massive, course straight. Secondary venation mixed craspedodromous (most of the secondary veins terminating at the margin and the rest brochidodromous); secondary veins thin, angle of divergence wide acute at 80°; course of secondary veins irregular, straight, slightly curved or recurved; secondary veins densely arranged, 5 to 10 pairs per cm at middle portion of lamina, spacing irregular; intersecondary veins common. Veins of higher order not observed.

**Number of specimens examined.** 2.

**Specimens illustrated.** UF15706-24648 (Figure 4.1).

**Occurrences.** Rose Creek locality, Nebraska; Braun Ranch and Hoisington III localities, Kansas.

**Remarks.** The venation pattern and tooth type of the two leaflet fragments are consistent with those described by Upchurch and Dilcher (1990) and Wang and Dilcher (2006a). Therefore, we interpret this specimen as conspecific and the blade as a leaflet. The chloranthoid tooth type (tooth with a prominent, often thickened gland, a medial vein and a pair of accessory veins that run along the tooth margin and fuse with the gland) and the secondary venation pattern of *Crassidenticum* show its closest similarities with extant Chloranthaceae (Upchurch and Dilcher 1990).

*Crassidenticum decurrens* Dilcher and Wang, 2006a

Figure 4.2

**Description.** Leaf simple, five-lobed; lobe shape lanceolate; medial lobe base symmetrical while lateral lobe base strongly asymmetrical; margin of lobes toothed except on the extreme base of lamina; axes of serrations inclined to the tangent of the margin, apical angle acute; serration type convex (basal side)-convex (apical side); sinus of tooth rounded; eight to 10 teeth per cm, regularly spaced; teeth simple (all of one size); sinus between lobes rounded, bracing accomplished by primary veins (lamina structure lacking or less than 0.5 mm wide on the sinus). Petiole thin; observed petiole 5 cm long and 0.5 mm wide. Secondary venation of lamina lobes pinnate; secondary veins thin. Veins of higher order not observed.

**Number of specimens examined.** 1.

**Specimens illustrated.** UF15706-24677 (Figure 4.2).

**Occurrences.** Braun Ranch and Hoisington III localities, Kansas.

**Remarks.** This species is common in the Braun Ranch locality, Kansas (Wang and Dilcher, 2006a). Though only one specimen is observed from the Hoisington III locality, a suite of characters, including trilobate leaf, massive multi-stranded primary vein, secondary venation pattern, and its chloranthoid tooth type are consistent with those described by Wang and Dilcher (2006a).

*cf. Crassidenticum trilobum* Dilcher and Wang, 2006a

Figure 4.3-4

**Description.** Leaf simple, five-lobed; lobe shape lanceolate; median lobe base symmetrical while lateral lobe base strongly asymmetrical; margin of lobes toothed except on the extreme base of lamina; axes of serrations inclined to the tangent of the margin, apical angle acute; serration type convex (basal side)-convex (apical side); sinus of tooth rounded; eight to 10 teeth per cm, regularly spaced; teeth simple (all of one size); sinus between lobes rounded, bracing accomplished by primary veins (lamina structure lacking or less than 0.5 mm wide on the sinus). Petiole thin; observed petiole 5 cm long and 0.5 mm wide. Secondary venation of lamina lobes pinnate; secondary veins thin. Veins of higher order not observed.

**Number of specimens examined.** 1.

**Specimens illustrated.** UF15706-24684 (Figure 4.3-4).

**Occurrences.** Hoisington III locality, Kansas.

**Remarks.** The single specimen examined lacks the details of higher order venation pattern. The five-lobed leaf appears to be compound, but there is lamina tissue connecting adjacent lobes (Figure 4.4). The secondary venation pattern of the lamina
FIGURE 4. 1. *Crassidenticulum decurrens* (Lesquereux) Upchurch and Dilcher, 1990, UF15706-24648, fragment of leaf lamina. Note fine teeth on margin (indicated by arrow). Scale bar equals 2 mm. 2. *Crassidenticulum trilobum* Dilcher and Wang, 2006a, UF15706-24677, a trilobed leaf. Note thin and long petiole. Scale bar equals 1 cm. 3-4. cf. *Crassidenticulum trilobum* Dilcher and Wang, 2006a, UF15706-24684, a five lobed leaf (scale bar equals 1 cm) (3) and an enlargement of Figure 4.3 to show decurrent lamina tissue between two adjacent lobes (indicated by arrow) (scale bar equals 1 mm) (4).
lobes and the tooth type on the margin are similar to those of *Crassidenticulum trilobum* Dilcher and Wang (2006a). It is possible that this species is a variant of *C. trilobum*.

This species differs from those compound leaves with several petiolate leaflets from the mid-Cretaceous Cedar Mountain Formation, Utah, i.e., Morphotype CM15 (Harris and Arens, 2016, p. 654-655, figure 7.1-4) and Europe, i.e., *Debeya insignis* (Hosius and von der Marck) Knobloch from southern Poland and western Ukraine (Halamski, 2013) and Austria (Herman and Kvaček, 2007a, 2007b, 2010), in that the leaf from the Hoisington III locality is simple with five lobes connected at the base with lamina tissue.

Family cf. NYMPHAEACEAE Salisbury, 1805 (including CABOMBACEAE A. Richard ex A. Richard, 1822)

Genus AQUATIFOLIA Wang and Dilcher, 2006b

**Type species.** *Aquatifolia fluitans* Wang and Dilcher, 2006b

*Aquatifolia fluitans* Wang and Dilcher, 2006b

**Description.** Leaf shape wide ovate (length/width ratio 1.2:1) to very wide ovate (L/W ratio 1:1 or less). Leaf base strongly cordate, with basal lobes overlapping or wide obtuse. Observed petiole length 28 mm long and 1.5 mm wide, containing a spherical float at a distance of ca. 10 mm from the leaf base (measured from the center of the spherical float to the leaf base). Float present on the petiole, spherical, sometimes appearing to be bilobed, or fusiform with some fine reticulate venation. Leaf margin entire, crenate, or strongly crenate. Typically, three veins entering a tooth with one medial vein parallel to the leaf margin and two flanking lateral veins (tertiary or quaternary) running along the margin before joining the medial vein. Primary venation is actinodromous or basal actinodromous; course of primary veins sinuous or straight; typically, one medial and four lateral primary veins (in two pairs) present in one leaf lamina. Primary veins and lateral secondary veins may fork up to three to four times, and adjacent branches join to form five - to six - sided elongate meshes oriented perpendicular to the leaf margin. All lateral primary veins and exmedial branches of lateral primary veins apically curved. Veins of higher order (tertiary or quaternary) orthogonally reticulate, forming quadrangular or pentagonal small meshes within those meshes formed by primary and secondary veins. Near the leaf margin, the veins dichotomize into veins of tertiary or quaternary order to form anastomosing loops from which arise one medial and two lateral veins that enter the tooth. Quaternary veins also orthogonally reticulate, forming the smallest polygonal meshes.

**Number of specimens examined.** 75.

**Specimens illustrated.** UF15706-8263, 8263’ (Figure 5.1); 24120 (Figure 5.2).

**Occurrences.** Hoisington III locality, Kansas.

**Remarks.** Two specimens from Wang and Dilcher (2006b) are illustrated in this report. Figure 5.1 is the holotype specimen of *Aquatifolia fluitans*, showing the spherical float on the petiole. Figure 5.2 is a paratype specimen showing actinodromous primary venation and ovate leaf shape.

Genus BRASENITES Wang and Dilcher, 2006b

**Type species.** *Brasenites kansense* Wang and Dilcher, 2006b

*Brasenites kansense* Wang and Dilcher, 2006b

**Description.** Leaf margin entire; shape suborbiculate (L/W ratio 1.2:1) to orbiculate (L/W ratio 1:1); lamina slightly funnel form at the position where the petiole is attached; leaf base peltate central (petiole attached within the boundaries of the leaf margin and near the center of the leaf). Primary venation actinodromous, consisting of 11 primary veins in one leaf lamina; five major primary veins present, with three of them directed oppositely (apically directed) from the other two; three minor primary veins radiate from the leaf center toward each lateral side; central major primary vein of the three extending nearly to the leaf margin, producing pinnate secondary veins; two apically directed and the two basally directed primary veins having strong exmedial branches, which fork repeatedly. Secondary veins produced from forked primary veins join adjacent branches to form polygonal (five- to six-sided) meshes both variable in shape and size, intergrading with higher-order (tertiary or quaternary) veins at a fourth to a third of the distance of the radius near the margin. Numerous small rounded protuberances present on the entire lamina.

**Number of specimens examined.** 4.

**Specimens illustrated.** UF15706-14806 (Figure 5.3-4).

**Occurrences.** Hoisington III locality, Kansas.

**Remarks.** One specimen (Figure 5.3) and its line drawing (Figure 5.4) are presented in this report. Figures 5.3 and 5.4 show the typically suborbiculate leaf shape, entire leaf margin, centrally peltate base, and major primary veins of this species. Similar leaves have also been reported from the early
FIGURE 5. 1-2, *Aquatifolia fluitans* Wang and Dilcher 2006b, UF15706-8263’, leaf showing the spherical float on the petiole (1) and UF15706-24120, leaf showing cordate base and thin high order venation (2). Scale bars equal 5 mm. 3-4, *Brasenites kansense* Wang and Dilcher, 2006b, UF15706-14806, specimen (3) and line drawing (4) to show suborbiculate leaf shape, entire leaf margin, peltate central base, major primary veins. Scale bars equal 1 cm.
Campanian Grünbach flora from the Grünbach Formation of the Grünbach-Neue Welt Basin, Austria (Herman and Kvaček, 2007a, 2007b), and the Late Cretaceous Arman Formation Northeastern Russia (Herman et al., 2016). Brasenites kansense differs from Exnelumbites callejasiae, a fossil taxon described from the Late Cretaceous McRae Formation, Mexico, in having much stronger bilateral symmetry to the venation, an entire margin, and suborbicular lamina (Estrada-Ruiz et al., 2011). For differences among Brasenites, Exnelumbites and other fossil members of the Nelumboaceae, and other aquatic plants, see Estrada-Ruiz et al. (2011, table 1).

A phylogenetic analysis that includes three Early Cretaceous fossil species (including Brasenites kansense) with peltate leaves (Taylor et al., 2008) indicates their affinities to Cabombaceae. However, in a recent analysis by Taylor and Gee (2014), adding the fossil leaves of Brasenites kansense (Wang and Dilcher, 2006b), Pluricarpellatia peltata (Mohr et al., 2008) and Scutifolium jordanicum (Taylor et al., 2008) results in the loss of most of the monophyletic groups found in the living-taxa analysis.

cf. Family SCHISANDRACEAE Blume, 1830 (including ILLICIACEAE A.C. Smith, 1947)
Genus LONGSTRETHIA Upchurch and Dilcher, 1990

Type species. Longstrethia varidentata Upchurch and Dilcher, 1990

Longstrethia aspera (Lesquereux) comb. nov. Figure 6

*1892 "Myrica" aspera Lesquereux, p. 66, pl. 2, fig. 11.

Emended specific diagnosis. Lamina margin toothed; serrate axes inclined to the tangent of margin, serration type straight or convex-straight or convex; tooth simple, spacing irregular; sinus rounded; three veins entering each tooth with one medial secondary vein or branch from secondary vein, two accompanying veins from basal and apical side, these two veins joining superadjacent veins of the same order to form loops before entering tooth. Primary venation pinnate; primary vein stout, multi-stranded, course curved. Secondary venation mixed craspedodromous (most of the secondary veins terminating at the margin and the rest brochidodromous); secondary veins thin relative to primary vein, subopposite; originating from primary vein at moderate acute angles, uniformly apically curved before entering the tooth or running along the basal side of tooth before terminating on the tooth with superadjacent secondary veins. Intersecondary veins present, composite. Tertiary veins thin, orthogonal reticulate, forming irregular meshes. Quaternary vein orthogonal reticulate, forming imperfect areoles. Veinlets simple, linear or curved.

Description. Observed lamina 9 cm to 15 cm long and 1.5 cm to 2.5 cm wide. Margin toothed; serrate axes inclined to the tangent of margin, serration type straight or convex-straight or convex; tooth simple, spacing irregular, one per cm; extending on all margin observed; sinus rounded, up to 1.5 mm deep (vertical distance from adjacent tooth apex to bottom of sinus); three veins entering each tooth with one medial secondary vein or branch from secondary vein, two accompanying veins from basal and apical side, these two veins joining superadjacent veins of same order to form loops before entering tooth. Primary venation pinnate; primary vein stout, multi-stranded, course curved. Secondary venation mixed craspedodromous (most of the secondary veins terminating at the margin and the rest brochidodromous); secondary veins thin relative to primary vein, one pair per cm, subopposite; originating from primary vein at moderate acute angles, uniformly apically curved before entering the tooth or running along the basal side of tooth before terminating on the tooth with superadjacent secondary veins. Intersecondary veins present, composite. Tertiary veins thin, orthogonal reticulate, forming meshes irregular in shape and size. Quaternary vein orthogonal reticulate, forming imperfect areoles. Veinlets simple, linear or curved.

Number of specimens examined. 2.

Lectotype. Longstrethia aspera Lesquereux (1892, p. 66, plate 2, figure 11; designated here).

Paratypes. UF15706-24578 (Figure 6.1, 6.3); 24650 (Figure 6.2).

Occurrences. Hoisington III locality, Kansas.

Remarks. Two specimens with middle portion of lamina are examined. Lesquereux (1892, p. 66, plate 2, figure 11) reported two specimens from the Pipe Creek locality, Cloud County, Kansas and assigned them to the extant genus Myrica. However, leaf architecture and cuticular characters of these leaves show a possible relationship to Illiciaceae or Trimeniaceae but differ from all extant members of the Magnoliidae in at least several characters (Upchurch and Dilcher, 1990, p. 33-34). We assign these specimens to the fossil genus Longstrethia established by Upchurch and Dilcher (1990) and set up a new combination for the specimens from the Hoisington III locality, Kansas.
FIGURE 6. *Longstrethia aspera* (Lesquereux) comb. nov. 1, UF15706-24578, middle and lower portion of lamina. Scale bar equals 1 cm. 2, UF15706-24560, specimen showing secondary veins and toothed margin. Scale bar equals 5 mm. 3, Enlargement of Figure 6.1 (left middle portion of lamina; indicated by arrow) to show fine venation. Scale bar equals 2 mm.
Longstrethia aspera is similar to Longstrethia varidentata Upchurch and Dilcher in that both have linear leaf shape and toothed margin, but they differ in that (1) leaf margin of Longstrethia varidentata varies from entire to coarsely toothed, (2) secondary venation varies from brochidodromous with strongly flattened brochidodromous loops, to pinnate with an intramarginal vein, and (3) relatively thin and numerous secondary veins. Future collection of specimens with well-preserved cuticle from the Hoisington III locality may provide information to clarify the relationship between the specimens from the Hoisington III locality, Kansas and those from the Rose Creek locality, Nebraska described by Upchurch and Dilcher (1990).

Clade MAGNOLIIDS Angiosperm Phylogeny Group, 2009
Order LAURALES Perleb, 1826
Genus PABIANIA Upchurch and Dilcher, 1990

**Type species.** Pabiania variloba Upchurch and Dilcher, 1990

_Pabiania variloba_ Upchurch and Dilcher, 1990

- Figures 7-8

v. 1981a _Acerites multiformis_ Lesquereux; Retallack and Dilcher, 1981a, p. 38, fig. 2.4.

**Description.** Leaf commonly trilobate, rarely bilobate or unlobed; base ranging from acute to obtuse and tending towards cuneate, with the basalmost portion of the lamina decurrent on the petiole; margin entire; apex of lobes acute to rounded; unlobed portion of the lamina decurrent on the petiole; narrowly tending towards cuneate, with the basalmost primary venation to form an open venation. Arealation poorly developed. Marginal venation consisting of a thin fimbrial vein.

**Number of specimens examined.** 90.

**Specimens illustrated.** UF15706-24423 (Figure 7.1-3); 30154 (Figure 7.4-5); 14832 (Figure 7.6); 14823 (Figure 8.1); 24464 (Figure 8.2, 8.4); 24587 (Figure 8.3).

**Occurrences.** Rose Creek locality, Nebraska and Hoisington III locality, Kansas.

**Remarks.** In Upchurch and Dilcher’s (1990) diagnosis and description of this species, the primary venation was described as palinactinodromous. Observations based on Hoisington III locality specimens indicate that the primary venation of _P. variloba_ can vary from basal actinodromous (Figure 8.1), suprabasal actinodromous (Figure 7.6), to suprabasal palinactinodromous (Figure 7.1-3). The leaves from the Rose Creek locality, Nebraska described by Upchurch and Dilcher (1990) and stored at the Paleobotany collection in the Florida Museum of Natural History (more than 100 specimens) are smaller in size compared with the specimens from the Hoisington III locality, Kansas. The specimens from the Hoisington III locality also show that the petioles are long and they have winged lamina tissue along both sides. The petiole base may be ocreate (Figure 7.4, 7.5). Other variations of leaf morphological characters include: (1) lamina from unlobed (Figure 8.3) to trilobed (Figures 7.1, 7.6, 8.1-2); leaf base from acute (Figure 7.6) to rounded (Figure 8.1); (2) apex of lamina lobes from acute (Figure 7.6), rounded (Figure 7.1), to mucronate (Figure 8.2). All other leaf morphological characters are consistent with those described by Upchurch and Dilcher (1990).

Heer (1883, plate 38, figure 3) and Lesquereux (1892, plate 54, figures 1-3) described a few specimens with similar leaf morphology and assigned them to _Aralia greenlandica_ Heer. Unfortunately, only primary and secondary venation are observed on these leaves, which makes it difficult to compare them with the specimens from the Rose Creek and Hoisington III localities.

-cf. Family LAURACEAE de Jussieu, 1789
Genus ROGERSIA Fontaine, 1889

**Type species.** Rogersia longifolia Fontaine, 1889

_Rogersia dakotensis_ Wang and Dilcher, 2009

- Figures 9.1, 9.3, 10.1
FIGURE 7. Pabiania varioloba Upchurch and Dilcher, 1990. 1-3, UF15706-24423, specimen showing suprabasal actinodromous primary venation (scale bar equals 1 cm) (1), enlargement of a sinus area to show sinus bracing by secondary veins (scale bar equals 2 mm) (2); and enlargement of basal portion of leaf to show two pairs of basal secondary veins (scale bar equals 2 mm) (3). 4-5, UF15706-30154, specimen showing a long petiole and ocrea-like structure at the base (scale bar equals 5 mm) (4) and enlargement of the petiole to show the ocrea-like structure (scale bar equals 2 mm) (5). 6, UF15706-14832, specimen showing suprabasal actinodromous primary venation and acute apices of the lobes. Scale bar equals 1 cm.
FIGURE 8. *Pabiania variloba* Upchurch and Dilcher, 1990. 1, UF15706-14823, leaf showing basal actinodromous primary venation and basal secondary veins. Scale bar equals 1 cm. 2, UF15706-24464, leaf showing rounded lobe apex. Scale bar equals 1 cm. 3, UF15706-24587, specimen showing a small lobe on the left and entire margin on the right of the leaf. Scale bar equals 5 mm. 4, Enlargement of Figure 8.2 to show straight primary vein extending to lobe apex and two series of loops in the excostal region. Scale bar equals 1 mm.
Description. Leaf simple; whole lamina and base symmetrical; form linear oblong, L/W >9, 7.5 cm to 9 cm long and 0.5 cm to 1 cm wide. Apex attenuate. Base acute, decurrent. Margin entire. Petiole short, stout. Primary venation pinnate; primary vein stout, multi-stranded; course straight or slightly curved. Secondary venation brochidodromous; secondary veins thin relative to primary vein; sub-opposite, decurrent on primary vein; angle of divergence narrow acute (ca. 30º), spacing irregular; secondary veins uniformly curved to join exmedial branches of superadjacent secondary veins or superadjacent secondary veins at a distance of one fifth to one fourth of half lamina to form an

FIGURE 9. 1 and 3, Rogersia dakotensis Wang and Dilcher, 2009, UF15706-24798, general leaf shape, note thin secondary veins forming intramarginal veins (scale bar equals 5 mm) (1) and enlargement to show intramarginal veins formed by looping secondary veins (scale bar equals 1 mm) (3). 2 and 4, Rogersia parlatorii Dilcher and Wang, 2006, UF15706-7529, specimen showing leaf shape (scale bar equals 5 mm) (2) and enlargement to show secondary venation (scale bar equals 5 mm) (4).
intercostal region with its long axis oblique primary vein (ca. 30°), then continuing to join exmedial branches of adjacent secondary veins to form two series of loops; these loops enclosing elongate areas with long axis almost parallel to leaf margin; secondary veins running along the margin appearing to be intramarginal veins. Intersecondary veins present, one per intercostal region, simple, extending about one half of the distance from primary vein to leaf margin and then intersect with superadjacent secondary veins. Tertiary veins thin, irregular, percurrent with retroflexed or straight course. Veins of higher order not observed.

**Number of specimens examined.** 200.

**Occurrences.** Hoisington III locality, Kansas and Courtland I locality, Minnesota.

**Specimens illustrated.** UF15706-24798 (Figure 9.1, 9.3); UF15706-24620 (Figure 10.1).

**Remarks.** Rogersia dakotensis is similar in leaf shape and angle of divergence of secondary veins to Wolfiophyllum heigii from the Braun’s Ranch locality of Kansas, but they differ in that W. heigii has eucamptodromous secondary venation, fork-
ing secondary veins near margin, and percurrent or exmedially ramified tertiary veins (Table 2).

**Rogersia parlatorii** Wang and Dilcher, 2006a

**Description.** Leaf simple; whole lamina and base symmetrical; form narrow elliptic, 7.5 cm long and 1.8 cm wide. Apex probably attenuate. Base acute, decurrent. Margin entire. Petiole normal, 1.1 cm long and 1 mm wide. Primary venation pinnate; primary vein stout, multi-stranded, course straight. Secondary venation festooned brochidodromous; ca. 10 pairs per lamina, opposite or subopposite, decurrent; angle of divergence narrow acute (ca. 30º), with lowest two or three pairs more acute than pairs above; spacing of secondary veins irregular, course uniformly curved; secondary veins joining exmedial branches of superadjacent secondary veins to form two series of loops. Intersecondary veins present, simple. Tertiary veins thin, predominately percurrent; angle of origin AO (acute on the lower side of the secondary and obtuse on the upper side), primary vein-tertiary vein angle oblique; course slightly wavy. Quaternary veins thin, orthogonal reticulate, anastomosing to form pentagonal or quadrangular meshes. Veins of higher order not observed.

**Number of specimens examined.** 2.

**Occurrences.** Braun Ranch and Hoisington III localities, Kansas.

**Specimens illustrated.** UF15706-7529 (Figure 9.2, 9.4).

**Remarks.** This species is abundant at the Braun Ranch locality, Kansas. See Table 2 for the differences between this species and other Dakota Formation entire margined leaves.

**Genus WOLFIOPHYLLUM** Dilcher and Wang, 2006

**Type species.** *Wolfiophyllum heigii* Dilcher and Wang, 2006

**Wolfiophyllum pfaffianum** (Heer) Wang and Dilcher, 2009

**Description.** Leaf simple; whole lamina and base symmetrical; form very narrow elliptic to narrow oblong, L/W 4 to >7, 10 cm to 12 cm long (estimated maximum length) and 1.3 cm to 2.6 cm wide. Apex missing. Base acute, decurrent. Margin entire. Petiole normal, short, stout, up to 1 cm long and 1.5 mm wide, with decurrent lamina tissue on both sides, curved to one side. Primary venation pinnate; primary vein stout, multi-stranded, course straight or slightly curved. Secondary venation eucamptodromous, secondary veins upturned and gradually diminishing apically near the margin, connected to the superadjacent secondary veins by a series of cross veins without forming prominent marginal loops; secondary veins up to 12 pairs per lamina; angle of divergence uniform, narrow acute (less than 45º); secondary veins uniformly curved and gradually diminishing apically inside the margin, connected to the superadjacent secondary veins without forming prominent loops; secondary vein course apically curved, occasionally forking near the margin; intersecondary veins common, one per intercostal region, extending half to almost the same distance as adjacent second-

### TABLE 2. Comparisons of leaf shape and venation of *Rogersia* and *Wolfiophyllum* species (modified from table 1 of Wang and Dilcher, 2009).

| Taxon            | Leaf shape       | 2° venation     | Number of 2° veins | Loops near margin | Intercostal region | Intersecondary veins | Tertiary veins       |
|------------------|------------------|-----------------|--------------------|-------------------|-------------------|----------------------|----------------------|
| *Wolfiophyllum*  |                  |                 |                    |                   |                   |                      |                      |
| *pfaffianum*     | very narrow      | eucamptodromous | 8                  | no                | no                | common               | reticulate           |
| *heigii*         | lorate or linear | eucamptodromous | 10 to 12           | no                | no                | present              | percurrent or exmedially ramified |
| *daphneoides*    | narrow ovate to  | eucamptodromous | 5 to 7             | no                | no                | common, simple       | Not observed         |
| *Rogersia*       |                  |                 |                    |                   |                   |                      |                      |
| *dakotensis*     | linear oblong    | brochidodromous | >10                | two series, elongate area | well defined | present              | percurrent, more or less irregular |
| *lottii*         | narrow elliptic  | festooned       | 10                 | one series, elongate area | well defined | present, simple     | random reticulate    |
| *parlatorii*     | oblong, lorate to linear | brochidodromous | >10                | irregular          | elongate and admedially oriented | common               | intergrading with 2° and 4° |

**Note:** The differences between this species and other Dakota Formation entire margined leaves.
ary veins; intersecondary veins simple. Tertiary veins predominately percurrent, course straight; angle of origin AO (acute on lower side of secondary veins and obtuse on upper side of secondary veins), forming cross veins between adjacent secondary and intersecondary veins. Veins of higher order poorly preserved.

**Number of specimens examined.** 1.
**Specimens illustrated.** UF15706-14815 (Figure 10.2-3).

**Occurrences.** Hoisington III locality, Kansas and Courtland I locality, Minnesota.

**Remarks.** The suite of characters, especially the combination of narrow elliptic simple leaf and eucamptodromous venation is different from any other angiosperm leaf megafossils from the Hoisington III locality, Kansas. The distinction between *Wolfiophyllum pfaffianum* with other similar leaves from the Dakota Formation is presented in Table 2.

**Order MAGNOLIALES** Bromhead, 1838
**Genus JARZENIA** Wang and Dilcher, 2009

*Jarzenia kanbrasota* Wang and Dilcher, 2009

**Description.** Lamina elliptic, L/W 2.5, 3.5 to 4 cm wide and 9 to 10 cm long (estimated length). Apex missing. Base acute, decurrent. Margin entire. Petiole missing. Primary venation pinnate; primary vein stout, multi-stranded, course straight. Secondary venation brochidodromous; secondary vein moderate relative to primary vein. >eight pairs per lamina, opposite to subopposite, decurrent; spacing irregular; angle of divergence narrow acute (<45º), uniformly curved to join exmedial branches of superadjacent secondary veins to form one series of loops. Intersecondary veins present, course simple. Tertiary veins not well preserved.

**Number of specimens examined.** 2.
**Specimens illustrated.** UF15706-3171 (Figure 11.1).

**Occurrences.** Hoisington III locality, Kansas and Courtland I locality, Minnesota.

**Remarks.** *Jarzenia kanbrasota* differs from *Setterholmia rotundifolia* (Lesquereux) Wang and Dilcher, 2009 in that *J. kanbrasota* has oblanceolate or narrow obovate shape, more cuneate leaf base, lowest one or two pairs of secondary veins originating at more acute angles than those above, not well defined intercostal regions, less percurrent tertiary veins, and the absence of glands on leaf lamina and veins. *Jarzenia kanbrasota* is very common at several localities of the Dakota Formation in Kansas and Nebraska based upon observation of specimens stored in the Paleobotany and Palynology collection at the Florida Museum of Natural History.

**Genus LIRIOPHYLLUM** Lesquereux, 1878

**Type species.** *Liriophyllum populoides* Lesquereux (designated by Berry 1902)
*Liriophyllum kansense* Dilcher and Crane, 1984

**Description.** Leaf petiolate, bilobed and deeply divided. Leaf 13.5 cm long and 16 cm wide; petiole 13 cm long and 3 mm wide. Primary vein stout, 3 mm wide, extending to the base of the sinus and forking at about 45° into two prominent veins forming the leaf margin in the lower part of the sinus. Apex of the lobes broadly rounded; leaf base straight. Secondary venation pinnate, camptodromous; secondary veins alternately arranged; angle of divergence of secondary veins gradually decreasing apically; secondary veins branching and gradually becoming thinner close to the margin to form weak, camptodromous loops, typically of tertiary order. Tertiary veins more or less decurrent where they join the secondary veins and the primary vein. Toward the leaf margin, tertiary, quaternary, and quinqueratary veins forming polygonal areolae.

**Number of specimens examined.** 30.
**Specimens illustrated.** UF15826-3188 (Figure 11.2).

**Occurrences.** Linnenberger Ranch and Hoisington III localities, Kansas.

**Remarks.** The primary vein running to the apex and dividing into a pair of prominent veins contiguous with the lamina margin in the sinus is unique in fossil and extant angiosperm leaves. Based upon the evidence of co-occurrence and common presence of distinctive resin-bodies in both the leaves and fruits, Dilcher and Crane (1984) suggested that *Liriophyllum kansense* leaves and *Archaeanthus*, multifollicular angiosperm fruiting axes, are from the same plant species. This plant is closely related to *Liriophyllum* in Magnoliaceae (Romanov and Dilcher, 2013).

**Clade EUDICOTS** Doyle and Hotton, 1991
**Order PROTEALES** Jussieu ex von Berchtold and J. Presl, 1820
**Family cf. NELUMBONACEAE** von Berchtold and J. Presl, 1820

**Genus PALEONELUMBO** Knowlton, 1930
FIGURE 11. 1. *Jarzenia kanbrasota* Wang and Dilcher, 2009, UF15706-3171, showing elliptic leaf shape and secondary venation. Scale bar equals 1 cm. 2. *Linophyllum kansense* Dilcher and Crane, 1984, UF15826-3188, showing deeply lobed leaf, long petiole, and secondary venation. Scale bar equals 1 cm. 3. *Credneria cyclophylla* (Heer) Wang and Dilcher, 2009, UF15706-14821, leaf showing craspedodromous venation. Note all secondary veins and their exmedial branches terminating on leaf margin, resulting in a wavy appearance of leaf margin. Scale bar equals 1 cm.
Type species. *Paleonelumbo macroloba* Knowlton, 1930

*Paleonelumbo* cf. *macroloba* Knowlton, 1930

**Figure 12**

**Description.** Leaf incomplete. Base peltate central; margin lobed with glandular teeth. Primary venation actinodromous with at least eight primary veins extending directly into individual lobes; sinus depth varying from one tenth to one half length of primary vein. Pinnate secondary veins present on the distal portion of lamina, brochidodromous. Tertiary and quaternary veins random reticulate. Vein of higher order not observed.

**Number of specimens examined.** 1.

**Specimens illustrated.** UF15826-24649 (Figure 12).

**Occurrences.** Hoisington III locality, Kansas.

**Remarks.** One incomplete specimen is observed. This specimen is different from other Cretaceous and Cenozoic species that are assigned to Nelumboaceae (Saporta, 1894; Berry, 1911d; Knowlton, 1922, 1930; Brown, 1954, 1962; Upchurch et al., 1994; Johnson, 2002; Barclay et al., 2003; Gandolfo and Cúneo, 2005; Estrada-Ruiz et al., 2011) in that the specimen from Hoisington III locality is lobed with varying depth of sinus and toothed margin. This specimen resembles *Paleonelumbo macroloba* Knowlton (Knowlton, 1930, plate 39, figure 3, plate 42, figures 3, 4; Brown, 1963, plate 35, figure 7; Johnson, 2002; Barclay et al., 2003, figure 9A) and *Nelumbo tenuifolia* (Lesquereux) Knowlton (Knowlton, 1922, plate 26, figure 7) in having a lobed lamina. They differ in that (1) the Hoisington III specimen has fewer primary veins; (2) it has veins of primary to tertiary order entering the marginal teeth; and (3) it does not possess transverse tertiary veins.

This leaf appears to be sagittate in shape but it only has one pointed lobe. The direction of this lobe is unknown since the leaf is incomplete. The deep sinus (Figure 12.1) does not seem to be the result of insect or mechanical damage since there are consistent looping veins along the margin. We tentatively assign this specimen to *Paleonelumbo macroloba* because its morphology is more similar to this species than other species assigned to Nelumboaceae.

Family cf. PLATANACEAE T. Lestiboudois, 1826

Genus CREDNERIA Zenker, 1883

Type species. *Credneria denticulata* Zenker, 1883

*Credneria cyclophylla* (Heer) Wang and Dilcher, 2009

**Figure 11.3**

**Description.** Whole lamina and base symmetrical, very wide ovate, L/W <1, 8 cm long (estimated length) and 8.5 cm wide. Apex missing. Base obtuse, decurrent. Margin toothed; tooth simple, extending on upper two-thirds of the margin, spacing irregular, two or three teeth per cm on middle portion of margin; dentate axes approximately perpendicular to the tangent of the margin; dentate apex obtuse (>90°), mucronate; toothed type concave on both sides; sinus rounded, shallow (less than 1 mm deep-vertical distance from tooth apex to bottom of sinus). Observed petiole 3 cm long and 1 mm wide. Primary vein stout, multi-stranded, course straight. Secondary venation simple craspedodromous (all of the secondary veins and their branches terminating at the margin); secondary veins thick relative to primary veins, multi-stranded, ca. five pairs per leaf lamina, opposite or subopposite; angle of divergence narrow acute (<45°) with uniform variation; secondary vein course straight or slightly curved, all terminating at the margin; two or three exmedial branches from basal two pairs of secondary veins also terminating on the margin. Intersecondary vein absent. Tertiary veins thick, percurrent, course convex (middle portion of vein curve away from the center of the leaf); angle of origin AA (acute on both sides of secondary veins); arrangement on secondary veins close (interval between veins less than 0.5 cm). Quaternary veins percurrent (perpendicular to tertiary veins), course straight; quaternary veins orthogonal reticulate, forming quadrangular well-developed areoles. Veinlets simple, linear or curved.

**Number of specimens examined.** 3.

**Specimen illustrated.** UF15706-14821 (Figure 11.3).

**Occurrences.** Hoisington III locality, Kansas and Courtland I locality, Minnesota.

**Remarks.** The specimens from Hoisington III locality are less variable in leaf shape and they are less abundant than those from the Courtland I locality, Minnesota.

Genus SAPINDOPSIS (Fontaine) Dilcher and Basson, 1990

**Type species.** *Sapindopsis magnifolia* Fontaine, 1889 (designated by Dilcher and Basson, 1990)

*Sapindopsis powelliana* (Lesquereux) comb. nov.

**Figures 13-16**
**FIGURE 12.** *Paleonelumbo cf. macroloba* Knowlton, 1930. 1, UF15826-24649, specimen showing primary venation (scale bar equals 5 mm) and 2, enlargement to show high order venation and toothed margin (scale bar equals 1 mm).

**Basionym.** *Rhus powelliana* Lesquereux (1892, p. 155, plate 56, figures 4, 5).

**Emended specific diagnosis.** Leaf compound, trifoliolate or paripinnate, consisting of three to six pinnately alternate to opposite leaflets; leaf stipulate, petiole thin; lateral leaflets petiolulate, terminal leaflets petiolulate or sessile, narrow oblong to narrow elliptic; leaflet margin entire. Primary venation of leaflets pinnate; primary vein multi-stranded, stout to massive, course straight. Secondary venation eucamptodromous or slightly brochidodromous; intersecondary veins common, composite. Tertiary veins thick relative to secondary veins, predominantly percurrent, slightly convex; arrange-
FIGURE 13. *Sapindopsis powelliana* (Lesquereux) comb. nov. 1, UF15706-14830, leaf with four leaflets. Note two ultimate leaflets appearing to be bilobed (indicated by arrow on the left), two oppositely arranged lateral leaflets, and stipule at the base. Scale bar equals 1 cm. 2, Enlargement of Figure 13.1 (indicated by arrow on the right) to show secondary and tertiary venation. Scale bar equals 2 mm. 3, Enlargement to of Figure 13.1 to show venation of the stipule. Scale bar equals 2 mm. 4, UF15706-14814, an elliptic leaflet with acuminate apex. Scale bar equals 1 cm. 5, Enlargement of Figure 13.4 to show secondary and tertiary venation. Scale bar equals 3 mm.
ment on secondary vein predominately alternate. Quaternary veins orthogonal, forming quadrangular or pentagonal imperfect areoles; veinlets simple, curved or straight.

Description. Leaf compound, trifoliolate (Figure 15) or paripinnate (Figure 16.1); leaflets alternate (Figure 16.1) to opposite (Figures 13.1, 14); leaf stipulate (Figures 13.1, 15); petiole thin, stipules up to 1.5 cm long and 0.6 cm wide, with venation parallel to the long axis; leaflets narrow oblong (Figures 13.1, 14.1-2, 15) to narrow elliptic (Figure 13.4); leaflet 6.5 cm to 13.5 cm long and 1.5 cm to 2.5 cm wide (L/W 4.3 to 5.4); leaflet margins entire, structurally reinforced, apex acute to acuminate, base acute and asymmetrical, terminal leaflet tissue decurrent on rachis; two ultimate leaflets oppositely arranged, with outer sides lamina decurrent on rachis (Figure 13.1), giving a bilobed appearance; the apex of the ultimate leaflets sometimes lobed once, with deep sinus extending about 30%
distance of lamina length, bracing of the sinus accomplished by forking of primary veins and then running along the margin within sinus (Figure 14.2). Lateral leaflets petiolulate and terminal leaflets petiolulate or sessile (Figure 14.2); petiolule thin, about 1 mm wide and up to 3.5 cm long (Figure 15). Primary venation of leaflets pinnate; primary vein multi-stranded, stout to massive (the ratio of vein width to lamina width is ca. 4%), course straight. Secondary venation eucamptodromous (Figure 9.2, 9.5; secondary veins upturned and gradually diminishing apically inside the margin, connected to the superadjacent secondary veins by a series of cross veins without forming prominent marginal loops) or slightly brochidodromous; secondary veins moderate in thickness, ca. 15 pairs per leaflet lamina; secondary veins uniformly originate from primary vein at moderate to narrow acute (less than 65°) angle, up to 15 pairs per leaflet, subopposite, uniformly curved and diminish near lamina margin; intersecondary veins common, composite (made up of coalesced tertiary vein segments for over 50% of its length). Tertiary veins thick relative to secondary veins, angle of origin acute-obtuse (AO, lower side of the secondary vein and upper side of the secondary veins), predominately percurrent, slightly convex; arrangement on secondary vein predominately alternate.
Quaternary veins orthogonal (arising at right angles), forming quadrangular or pentagonal imperfect areoles; veinlets simple, curved or straight.

**Number of specimens examined.** 110.

**Neotype (designated here).** UF15706-14830 (Figure 13.1-3).

**Other specimens illustrated.** UF15706-14814 (Figure 13.4-5); 24670 (Figure 14.1); 4812 (Figure 14.2).
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14.2); 24719 (Figure 15); 24711 (Figure 16.1); 24675 (Figure 16.2).

Remarks. Lesquereux (1892, p. 155, plate 56, figures 4, 5) described two specimens from the Dakota flora and assigned them to an extant genus and established a new species, Rhus powelliana. He did not designate a holotype specimen for the new species and both specimens he described are currently missing. Based on Article 9.7 of the International Code of Botanical Nomenclature (McNeill et al., 2012), we here designate one specimen from Hoisington III locality, Kansas (UF15706-14830; Figure 13.1-3) as the neotype. We transfer this species to the genus Sapindopsis based upon the following diagnostic characters: 1) pinnately compound leaf with terminal lobes or leaflets commonly more or less united at their base (Figure 13.1); 2) distal leaflets with lamina decurrent to the leaf rachis and continuing as a narrow marginal lamina on either side of the rachis (Figure 14.1); 3) the decurrent lamina on the rachis missing entirely toward the leaf base (Figure 14.2); 4) leaflets elliptical to lanceolate-shaped, stout midrib extending to the apex of each leaflet (Figures 13-15); and 5) eucamptodromous secondary venation.

The genus Sapindopsis was established by Fontaine (1889) and the generic diagnosis was emended by Berry (1911d). A total of eight species were originally proposed. Dilcher and Basson (1990) emended the generic diagnosis and proposed that the simple leaf forms (e.g., Sapindopsis cordata Fontaine, 1889, p. 296, plate 147, figure 1), in synonymy with Ficophyllum crassinerve Knowlton (1919) and Sapindopsis elliptica Fontaine (1889, p. 297, plate 147, figure 3), and in synonymy with Rogersia longifolia Berry (1911d) should be excluded from the genus Sapindopsis. The large distance between adjacent secondary veins and the well-defined intercostal region by secondary veins in Rogersia are distinct from those of the Sapindopsis powelliana leaves. The other six species, including Sapindopsis magnifolia Fontaine (1889, p. 297, plate 151, figures 2, 3, plate 152, figures 2, 3, plate 153, figures 2, 3, plate 154, figures 1, 5, plate 155, figure 6; Berry, 1911d, p. 471, plate 86, figures 1-3, plate 87, figure 1, plate 88, figure 1; Berry, 1922, p. 214, plate 55, figure 5, plate 56, plate 57, figure 2, plate 59, figure 3; Doyle and Hickey, 1976, p. 166-167, figures 17-19; Hickey and Doyle, 1977, p. 35, figures 33-38), Sapindopsis variabilis Fontaine (1889, plate 151, figure 1, plate 152, figures 1, 4, plate 153, figure 3, plate 154, figures 2-4, plate 155, figures 2-5; Berry, 1911d, p. 469, plate 83, figures 109, plate 84, figures 1-2, plate 85, figure 1; Berry, 1922, p. 213, plate 55, figures 2-4), Sapindopsis parvifolia Fontaine (Fontaine, 1889, p. 300, plate 154, figure 6), Sapindopsis brevifolia Fontaine (Fontaine, 1889, p. 300, plate 153, figure 4, plate 155, figures 1, 7, plate 163, figure 3; Berry, 1911d, p. 473, plate 87, figures 2-5; Berry, 1922, p. 216, plate 55, figure 1, plate 59, figure 1), Sapindopsis tenuinervis Fontaine (Fontaine, 1889, p. 301, plate 153, figure 1), and Sapindopsis obtusifolia Fontaine (Fontaine, 1889, p. 301, plate 156, figure 13, plate 159, figures 3-6), can be distinguished from Sapindopsis powelliana by their sessile leaflets. Sapindopsis belviderensis Berry (Berry, 1922a, p. 216-217, plates 49-54; Hickey and Doyle, 1977, p. 35-36, figures 39-40) has toothed leaflets which is different from Sapindopsis powelliana.

Sapindopsis powelliana is similar to a specimen (Sapindopsis sp.) illustrated as the frontispiece by Hickey and Doyle (1977). This specimen is from Red Point, Cecil County, Maryland (upper Subzone II-B; late Albian), and it may belong to the same species. They are similar in that both species have lateral leaflets with long petiolules and lobed terminal leaflets.

Dilcher and Basson (1990) described one species (also see Krassilov and Bacchia, 2000), Sapindopsis anhouryi from the Nammoura locality, Lebanon. The stipules of this species are similar to those of S. powelliana leaves but they differ in that the leaflets of Sapindopsis powelliana are petiolulate and the secondary venation is eucamptodromous. Well-defined intercostal regions by secondary veins and intramarginal veins are also absent from the leaflets of Sapindopsis powelliana.

Huang and Dilcher (1994) described four types of pinnately lobed leaves from the Cheyenne Sandstone. Of these leaves, Sapindopsis sp. B is similar to the trifoliate leaf (Figure 15) from the Hoisington III locality. Unfortunately, it is difficult to compare them in detail because no venation higher than the second order can be observed for the Cheyenne Sandstone specimens.

Sapindopsis powelliana leaves can be distinguished from other Cretaceous species of Sapindopsis reported around the world by its entire leaflet margin, petiolulate leaflets, eucamptodromous secondary venation, and predominately persistent tertiary venation. These species include, Sapindopsis cf. Sapindopsis magnifolia/variabilis Fontaine (Upchurch et al., 1994, p. 40, figures 54-57), Sapindopsis minitifolia Upchurch et al. (1994, p. 41, figures 58, 59), Sapindopsis lebanensis Krassilov and Bacchia (Krassilov and Bacchia, 2000), Sapindopsis anhouryi from the Nammoura locality, Lebanon.
always curved abaxially; apex attenuate; base nor-
proximally on rachis; rachis thin. Leaflet lorate,
sionally with one leaflet attached at some distance
arranged on the distal portion of the rachis, occa-
four leaflets; most commonl y three leaflets closely

Doyle and Hickey (1976) and Dilcher and Basson (1990) recognized two distinct types of Sapindopsis leaves. One type has fused terminal leaflets and decurrent lamina tissue on the rachis. The other type rarely has fused terminal leaflets and frequently the leaflets are petiolulate and lack any decurrent lamina tissue on the rachis. Most leaves of Sapindopsis powelliana belong to the second type but with occasional exceptions (Figure 14.1). As suggested by Dilcher and Basson (1990), a detailed study of the leaf types of Sapindopsis is needed in order to separate these forms into dis-

Compared with all other species assigned to Sapindopsis, S. powelliana seems to have the highest leaf venation rank. Most other species of Sapindopsis from the Potomac Group have the second leaf rank. This is in agreement with Doyle and Hickey (1976) and Hickey and Doyle’s (1977) summary of the Potomac leaf sequence because the Dakota Formation is younger than the Potomac Group.

Although the genus Sapindopsis was origi-
ally proposed by Fontaine (1889) as a member of the Sapindaceae, there is no evidence that Sapin-
dopsis is more closely related to extant Sapinda-
cae than to other groups (Crane, 1989; Crane et al., 1993; Upchurch et al., 1994). Many features of leaf architecture and cuticular anatomy in fossil Sapindopsis, and its co-occurrence with platanoid reproductive structures show that they are proba-
bly more closely related to Platanaceae (Hickey and Wolfe, 1975; Hickey and Doyle, 1977; Upchurch, 1984; Upchurch and Dilcher, 1990; Wang et al., 2011).

Sapindopsis retallackii sp. nov.

Specific diagnosis. Leaf compound, with three or four leaflets; most commonly three leaflets closely arranged on the distal portion of the rachis, occasion-
ally with one leaflet attached at some distance proximally on rachis; rachis thin. Leaflet lorate, always curved abaxially; apex attenuate; base nor-
mal acute; lateral leaflets sessile and terminal leaf-
lets petiolulate; margin entire, usually revolute. Primary venation pinnate; primary vein stout; course straight (medial leaflet) or recurved (lateral leaflets). Secondary venation brochidodromous; secondary veins thin relative to primary vein; opposite to subopposite, decurrent; angle of divergence acute, joining superadjacent secondary veins to form a loop enclosing an intercostal area, with two series of loops (tertiary and quaternary in order) in excostal region; exmedial branches of secondary veins common, forming tertiary veins. Intersecond-
ary veins common. Tertiary vein moderate; orthog-
onal reticulate but tending to be percurrent, retroflexed. Quaternary veins random reticulate, forming incompletely closed meshes. Veinlets sim-
ple, linear.

Description. Leaflet lamina base symmetrical. Form lanceolate, L/W >5, 10 cm long (estimated length) and 1.8 cm wide. Apex missing. Base nor-
mal, acute; petiolule 1 cm long and 1 mm wide. Margin entire. Petiole normal, short, 0.8 cm long and 0.5 mm wide. Primary venation pinnate; pri-
mary vein stout, slightly curved, up to 0.5 mm wide at the widest portion of lamina. Secondary venation festooned brochidodromous; secondary vein fine relative to primary vein; secondary veins diverging from primary vein at narrow acute angles, with lowest one to two pairs more acute than pairs above; secondary veins slightly recurved after diverging from primary vein, then extending a distance to about one fourth to one half of half lamina before joining superadjacent secondary or intersecondary veins to form two series of loops in the excostal region. Intersecondary veins common, one to three pairs per intercostal region; intersecondary veins simple or occasionally forking at variable distance after diverging from primary vein; intersecondary veins almost the same width as secondary veins. Tertiary veins hair-like; diverging from primary veins, exmedial (lower) side of secondary veins or intersecondary veins at moderate acute to narrow acute angles, predominately exmedially ramified, orientation parallel to secondary or intersecondary veins, connected by cross veins of the same order.

Number of specimens examined. 30.

Holotype. UF15706-3153 (Figure 17).

Species epithet. In recognition of Greg Retallack and his contribution to the Dakota Formation geol-
y and paleobotany.

Occurrences. Hoisington III locality, Kansas.

Remarks. Only one specimen, representing a ter-

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Springfield locality and Pleasant Dale locality, Nebraska (Wang and Dilcher, in preparation).

_Sapindopsis retallackii_ is similar to _Sapindopsis_ sp. A (Huang and Dilcher, 1994) in having three leaflets but they differ in that _S. retallackii_ occasionally has four lorate leaflets, and brochidodromous secondary venation. _Sapindopsis retallackii_ can be distinguished from other _Sapindopsis_ species (see previous section) by the following suite of characters: compound leaf with petiolulate terminal leaflets, sessile lateral leaflets, lorate leaflet shape, brochidodromous secondary venation, the presence of exmedial branches of secondary veins that branch into tertiary veins, and common intersecondary veins.

**Clade ROSIDS Angiosperm Phylogeny Group, 2009**

**Genus ANISODROMUM Upchurch and Dilcher, 1990**

**Type species.** _Anisodromum wolfei_ Upchurch and Dilcher, 1990

_Anisodromum wolfei_ Upchurch and Dilcher 1990

**Figure 18**

**Description.** Leaflet lamina and base symmetrical. Shape narrow ovate or elliptic, L/W 2 to 2.6, lamina 3 to 4.7 cm wide and 8 to 9 cm long. Apex missing. Base obtuse. Margin entire. Observed petiolule with a 0.2 to 0.3 mm wide deciduous lamina tissue; petiole 0.5 mm wide and 2 mm long. Primary vena-
tion of leaflets pinnate; primary vein stout, multi-stranded, course straight. Secondary venation predominately eucamptodromous; secondary veins moderate relative to primary vein, multi-stranded, ca. nine pairs per lamina, opposite or slightly sub-opposite, decurrent; angle of divergence moderate acute (45° to 65°), with basal pairs at slightly more obtuse angles; basal secondary veins joining superadjacent secondary veins or their exmedial branches to form loops very close to margin; other secondary veins uniformly curved and diminishing near margin; adjacent secondary veins connected

FIGURE 18. Anisodromum wolfei Upchurch and Dilcher, 1990. 1, UF15706-14818, general leaflet shape. Note long petiolule. Scale bar equals 1 cm. 2, Enlargement of Figure 18.1 to show strong secondary veins and percurrent tertiary veins oriented perpendicular to the primary vein. Scale bar equals 2 mm. 3, UF15706-24566, general leaflet shape. Scale bar equals 5 mm. 4, Enlargement of Figure 18.3 to show percurrent tertiary veins oriented almost perpendicular to the primary vein. Scale bar equals 2 mm.
by percurrent tertiary veins. Tertiary veins percurrent, moderate relative to secondary veins; angle of origin AO (acute on exmedial side of secondary vein and obtuse on admedial side of secondary veins) or RO (right on primary vein and obtuse on admedial side of secondary vein); course predominately straight, oriented almost at right angle with primary vein; arrangement close (interval between veins less than 0.5 cm). Quaternary vein orthogonal reticulate. Veins of higher order not observed.

Number of specimens examined: 3.

Specimens illustrated. UF15706-14818 (Figure 18.1-2); 24566 (Figure 18.3-4).

Remarks. Upchurch and Dilcher (1990) described six specimens from the Rose Creek locality, Nebraska and they proposed that the compound leaves most closely resemble *Sapindopsis* but they differ in that *Anisodromum wolfei* has secondary venation that shows asymmetric behavior and much less brochidodromous looping, more percurrent and closely spaced tertiary venation, and a structurally reinforced margin. These characters were interpreted as significant at generic level by analogy to extant pinnately compound Rosidae, where genera are distinguished on the basis of secondary, tertiary, and marginal venation (Upchurch and Dilcher, 1990).

*Anisodromum upchurchii* sp. nov. Figure 19.1-2

Specific diagnosis. Lamina base acute, decurrent. Primary venation pinnate; primary vein stout, multi-stranded, straight. Secondary venation brochidodromous, secondary veins moderate relative to primary vein, opposite, subopposite or alternate; angle of divergence wide acute, uniformly curved apically; secondary vein spacing distant. Intersecondary veins common, usually opposite to a secondary vein on the other side of lamina. Tertiary veins moderate relative to secondary veins, percurrent, course straight; angle of origin AO (acute on exmedial side and obtuse on admedial side of secondary veins or intersecondary vein) or RO (right on primary vein and obtuse on admedial side of secondary or intersecondary veins), primary vein-tertiary vein angle almost right (90º); tertiary vein interval close (interval between veins less than 0.5 cm). Veins of higher order not observed.

Number of specimens examined: 4.

Holotype. UF15706-24576 (Figure 19.1-2).

Derivation of epithet. In honor of Garland R. Upchurch in recognition of his contributions to Dakota Formation angiosperm paleobotany.

Remarks. *Anisodromum upchurchii* differs from *Anisodromum wolfei* in that it has large angles of divergence for secondary veins, large distance between secondary veins, fewer secondary veins, and intersecondary veins arranged opposite to secondary vein on the opposite side of the lamina. *Anisodromum upchurchii* and *A. schimperi* are similar in that they both have irregularly spaced secondary veins. They differ in that the secondary venation of *A. schimperi* is festooned brochidodromous and its tertiary veins are orthogonally reticulate near primary vein.

The wide divergence angle of the secondary veins and the primary-tertiary vein angle of *Anisodromum upchurchii* are similar to Morphotype CM05 (*Magnoliaephyllum* sp.) from the mid-Cretaceous Cedar Mountain Formation of Utah, USA (Harris and Arens, 2016, p. 646-648, figure 4.1-3). They differ in that the divergence angle of secondary veins is larger and the percurrent tertiary veins is less dense in *Anisodromum upchurchii* compared with those of Morphotype CM05.

It is unknown if the specimens from the Hoisington III locality represent simple or compound leaves. Therefore, we tentatively assign these specimens to the genus *Anisodromum*. Future collection with better preserved specimens may confirm this assignment or help separate them from this genus.

*Anisodromum schimperi* (Lesquereux) comb. nov. Figure 19.3-4

Basionym. “*Persea*” *schimperi* Lesquereux (1892, p. 103, plate 16, figure 5).

Emended specific diagnosis. Lamina margin entire. Primary venation pinnate; primary vein stout, multi-stranded, course straight. Secondary venation festooned brochidodromous; secondary veins moderate relative to primary veins, subopposite to alternate, decurrent, spacing irregular, with
one side more closely spaced than the other side; secondary vein curved apically, joining tertiary veins from superadjacent secondary veins to form two series of loops. Tertiary veins moderate relative to secondary veins, predominately percurrent, orthogonal reticulate near primary vein; course irregular, straight or sinuous; primary-tertiary vein angle oblique.

**Description.** Whole lamina and base symmetrical, lamina elliptic, L/W 1.5, lamina 2.9 cm to 3.1 cm wide and 4.5 cm to 5.5 cm long. Apex obtuse. Base acute normal. Margin entire. Petiole not observed. Primary venation pinnate; primary vein stout, multi-

**FIGURE 19.** 1-2, *Anisodromum upchurchii* sp. nov., UF15706-24576, middle portion of lamina showing strong primary and secondary veins (scale bar equals 1 cm) (1) and enlargement of an area in Figure 19.1 (indicated by arrow) to show percurrent tertiary veins (scale bar equals 2 mm) (2). 3-4, *Anisodromum schimperi* (Lesquereux) comb. nov., UF15706-24635, leaf showing uneven spacing of secondary veins and reticulate tertiary veins (scale bar equals 5 mm) (3) and; UF15706-24633, leaf showing secondary venation and reticulate tertiary veins (scale bar equals 5 mm) (4).
secondary vein curved apically, joining tertiary veins (AO); course irregular, straight or sinuous; primaryRedirected to internal server error

tertiary vein angle oblique. Veins of higher order (AO); course irregular, with one side more close than the other; angle of divergence more acute than those of the other side; secondary veins, predominately percurrent, orthogonal reticulate near primary vein; angle of origin (AO); course irregular, straight or sinuous; primary-tertiary vein angle oblique. Veins of higher order not observed.

Number of specimens examined. 2.

Neotype (designated here). UF15706-24633 (Figure 19.4).

Other specimens illustrated. UF15706-24635 (Figure 19.3).

Remarks. Characters such as entire margin, multi-stranded primary vein, and festooned brochidodromous secondary venation indicate the affinity of this species with Magnoliales, but the uneven secondary vein spacing may indicate these specimens may represent leaflets of a compound leaf.

Lesquereux (1893, p. 103, plate 16, figure 5) described one specimen from a locality in Minnesota, USA and assigned it to the extant genus Persea. In this report, we retain Lesquereux's species epithet and transfer his specimen to Anisodromum. Because his original specimen is lost and he did not designate a type, we here emend his species diagnosis and designate a specimen from the Hosiington III locality as the neotype.

Genus CITROPHYLLUM Berry, 1911a

Type species. Citrophyllum aligera Lesquereux, 1892 (designated by Upchurch and Dilcher 1990)

Citrophyllum aligera Lesquereux, 1892

Figure 20

*1892 Ficus aligera Lesquereux, p. 84, pl. 10, figs. 3-6.

*1909 Citrophyllum aligerum (Lesquereux) Berry: Berry, p. 258, pl. 18a, figs. 1-8.

*1911a Citrophyllum aligera (Lesquereux) Berry: Berry, p. 169, pl. 21, figs. 1-8.

Emended specific diagnosis. Leaf simple, whole lamina and base symmetrical. Apex obtuse. Margin entire. Petiole aliform, base inflated; demarcation between lamina wing and lamina distinct. Primary venation pinnate; primary vein stout, course straight. Secondary venation festooned brochidodromous; secondary veins thin relative to primary vein, subopposite to alternate; angle of divergence moderate acute; secondary veins uniformly curved, joining exmedial branches of superadjacent secondary vein at right or obtuse angles to form two series of loops. Intersecondary veins common, simple. Tertiary veins random reticulate, but tending to be percurrent near leaf margin. Quaternary and quinternary veins random reticulate.

Description. Leaf simple, whole lamina and base symmetrical. Form ovate to narrow elliptic, L/W 1.3 to 2.6, lamina 3 to 6 cm wide and 8 cm long. Apex obtuse. Base normal obtuse or cuneate. Margin entire. Petiole aliform, base inflated; petiole 0.5 to 1.8 mm wide and 2.5 cm to 4 cm long; lamina wing 0.2 mm to 1.8 mm wide on each side; demarcation between lamina wing and lamina distinct. Primary venation pinnate; primary vein stout, 0.5 mm to 1 mm wide, course straight. Secondary venation festooned brochidodromous; secondary veins thin relative to primary vein, ca. nine pairs per leaf lamina, subopposite to alternate; angle of divergence moderate acute (ca. 50º), with lowest pair more obtuse than pairs above; secondary vein uniformly curved, joining exmedial branches of superadjacent secondary vein at right or obtuse angles to form two series of loops. Intersecondary veins common, one to two between adjacent secondary veins, simple, extending a distance of one third to one half of half lamina. Tertiary veins random reticulate, but tending to be percurrent near leaf margin. Quaternary and quinternary veins random reticulate.

Number of specimens examined. 6.

Neotype (designated here). UF15706-24646 (Figure 20.1).

Other specimens illustrated. UF15706-24645 (Figure 20.2); 24332 (Figure 20.3-4).

Remarks. Citrophyllum aligera differs from Citrophyllum doylei, a species established by Upchurch and Dilcher (1990, p.43-45, plate 27, figures 1, 3, 4; text-figure 21), in having a longer petiole and festooned brochidodromous secondary venation. Some specimens from the Hosiington III locality occasionally don’t have alate petioles. These specimens may represent the further reduction of lamina tissue on the petiole.

The genus Citrophyllum was established by Berry (1909) to accommodate fossil leaves with elliptical to ovate or ovate-lanceolate, coriaceous leaves, with entire or slightly undulate margins and stout conspicuously alate petioles. These characters are generally similar to those of the modern genus Citrus. Berry (1909, p. 258) transferred Les-
Lesquereux’s (1892) Dakota specimens, originally assigned to the extant genus *Ficus*, to *Citrophyl-

lum*. As discussed by Upchurch and Dilcher (1990, p. 44), the modern affinities of the genus *Citrophyl-

lum* is with Sapindales instead of Rutaceae because it doesn’t possess features that characterize Rutaceae. Leaves of Rutaceae have been reported only from the Cenozoic of Asia (Guo, 2011; Xie et al., 2013) and Africa (Pan, 2010), but no Rutaceae fossils have been found from the Cretaceous.

**Clade Unknown**

**Genus WINGIA gen. nov.**

**FIGURE 20.** *Citrophylum aligera* Lesquereux, 1892. 1. UF15706-24646, base of leaf showing lamina wing on petiole. Scale bar equals 5 mm. 2. UF15706-24645, base of leaf showing a long petiole with lamina wing and inflated petiolar base. Scale bar equals 5 mm. 3. UF15706-24332. leaf with a long petiole with thin lamina wing and inflated petiolar base. Scale bar equals 1 cm. 4. Enlargement of an area in Figure 20.3 (indicated by arrow) to show looping marginal veins. Scale bar equals 1 mm.
**Generic diagnosis.** Leaf simple, irregularly five-lobed; lamina margin with glandular teeth at or near apex of lobes, strongly reinforced; sinus rounded, bracing accomplished by two secondary veins originated from adjacent primary veins; serrations minute, simple, typically three veins entering a tooth, medial vein originating from exmedial side of secondary vein, basal vein tertiary in order and running very close to the margin before entering the tooth, vein of apical side not well developed. Petiole long and thin, enlarged both distally and proximally (perhaps these are pulvini). Primary venation basal actinodromous; primary vein stout to massive, multi-stranded; medial primary vein straight in course; inner lateral primary veins apically curved; outer primary veins apically curved or recurved. Secondary venation predominately brochidodromous or semicraspedodromous when margin is toothed near apex; secondary veins thin relative to primary veins; originating from primary veins at wide acute to angle, straight or slightly curved and then abruptly curved very close to the margin to join superadjacent secondary veins to form rectangular or rhomboidal intercostal regions; Intersecondary veins common, composite. Tertiary veins orthogonal reticulate, forming predominately quadrangular meshes.

**Type species.** *Wingia expansolobum* (Upchurch and Dilcher) Wang and Dilcher (designated here).

**Derivation of generic name.** In honor of Scott Wing in recognition of his contribution to angiosperm paleobotany.

**Remarks.** Upchurch and Dilcher (1990, p. 48-49, plate 31, text figure 25) established a new species based upon one specimen from the Rose Creek locality, Nebraska and assigned this species to *Dicotylophyllum*. This species differs from all other lobed leaves from the Dakota Formation of Kansas, including "*Aralia* quinquepartita" Lesquereux (1871, p. 302, 1874, plate 15, figure 6) and "*Sterculia* lugubris" Lesquereux (1883, plate 6, figures 1-3), in having small teeth and five lobes. The genus *Dicotylophyllum*, first established by Saporta (1894), is heterogeneous and it may only include probable nymphaealan leaves (see Halamski, 2013, p. 429-430 for detailed discussion). Therefore, we establish a new genus for this species from the Dakota Formation.

*Wingia expansolobum* (Upchurch and Dilcher) comb. nov.

Figures 21-23

**Basionym.** *Dicotylophyllum expansolobum* Upchurch and Dilcher (1990, p. 48-49, plate 31, text-figure 25).

**Specific diagnosis.** Leaf simple, irregularly five-lobed, margin toothed near apex of lobes, serrations minute, simple. Petiole long and thin, enlarged both distally and proximally. Primary venation basal actinodromous. Secondary venation predominately brochidodromous or semicraspedodromous when margin is toothed near apex; typically, one series of loops present in the excostal region if margin is entire; secondary veins below sinus and between adjacent primary veins forming inverted ‘V’ pattern. Intersecondary veins common, composite. Tertiary and quaternary veins orthogonal reticulate, forming predominately quadrangular meshes.

**Description.** Leaf simple, five-lobed, lamina varies from 3.2 cm long and 6 cm wide to 15 cm long and 15 cm wide; sinus deep, extending 75% to 80% of distance from apex to lamina base; medial lobe narrow obovate, narrow ovate or narrow oblong, symmetrical; lateral lobes asymmetrical, shape irregular, curved apically; outer lobes occasionally underdeveloped (leaf tending to be trilobate); apex of lobes obtuse; base of lamina obtusely cuneate or strongly rounded to give a truncate appearance; margin toothed near apex of lobes, strongly reinforced; sinus rounded, bracing accomplished by two secondary veins originated from adjacent primary veins, these two veins join and then fork to run along the margin; serrations minute, simple; serration type ranging from concave to straight on apical side and straight to convex on basal side, spacing irregular; typically three veins entering a tooth, medial vein originating form exmedial side of secondary vein, basal vein tertiary in order and running very close to the margin before entering the tooth, vein of apical side not well developed. Petiole long, thin, 8 cm long and 1 mm wide; pulvini (lower and upper) present; upper pulvinus extending up to 1 cm proximally along the petiole. Primary venation basal actinodromous; primary veins stout to massive, multi-stranded; medial primary vein straight in course; inner lateral primary veins apically curved; outer primary veins apically curved or recurved. Secondary venation predominately brochidodromous or occasionally semicraspedodromous near apex; secondary veins thin relative to primary veins; originating from primary veins at wide acute angles, straight or slightly curved and then abruptly curved very close to the margin to join superadjacent secondary veins to form rectangular or rhomboidal intercostal regions; typically one series of loops present in the excostal region if margin is entire; secondary veins below sinus and between adjacent primary veins forming inverted
"V" pattern. Intersecondary veins common, composite, one to three per intercostal region. Tertiary veins orthogonal reticulate, forming predominately quadrangular meshes, tending to be percurrent and arranged at very oblique angles (<20°) with primary veins. Quaternary veins orthogonal reticulate. Veins of higher order not observed.

**Number of specimens examined.** 52.

**Holotype.** UF15713-8304 (designated by Upchurch and Dilcher, 1990).

**Specimens illustrated.** UF15706-30158 (Figure 21.1-3); 24788 (Figure 21.4); 14825 (Figure 22.1-2); 24461 (Figure 22.3-4); 14828 (Figure 23.1-3).

**Remarks.** This species was established by Upchurch and Dilcher (1990) based on one specimen from the Rose Creek I locality, Nebraska. We emend the specific diagnosis because specimens

**FIGURE 21.** *Wingia expansolobum* (Upchurch and Dilcher) comb. nov. 1, UF15706-30158a, specimen showing a complete leaf. Note the long petiole with a swollen base and the extension of lamina tissue on the petiole. Scale bar equals 1 cm. 2, Enlargement of the petiole in Figure 21.1 to show the swollen base. Scale bar equals 2 mm. 3, UF15706-30158b, enlargement of a leaf on the back of the same specimen to show high order venation and glandular teeth. Scale bar equals 3 mm. 4, UF15706-24788, specimen showing a deeply lobed leaf. Note apically curved outer lateral lobes. Scale bar equals 1 cm.
from the Hoisington III locality yield more information on the leaf features, especially the variations of leaf morphology. For example, secondary venation of *Wingia expansolobum* can vary from predominately brochidodromous to semicraspedodromous. High order venation (tertiary and quaternary veins) is well preserved on the specimens from the Hoisington III locality, Kansas.

The proximal and distal enlarged areas on the petioles of *Wingia expansolobum* leaves may be pulvini. In extant plants, the pulvinus, a motor organ for leaf movement (regulating leaf position),

**FIGURE 22.** *Wingia expansolobum* (Upchurch and Dilcher) comb. nov. 1, UF15706-14825, specimen showing a large leaf with irregular shape of lateral lobes. Note the structurally reinforced margin on sinus, pluvinus lamina extension on the thin and long petiole. Scale bar equals 1 cm. 2, Enlargement of Figure 22.1 to show the pluvinus lamina extension on the petiole. Scale bar equals 2 mm. 3, UF15706-24461, enlargement of Figure 22.4 (area indicated by arrow) to show looping of secondary veins near the lamina margin. Scale bar equals 1 mm. 4, UF15706-24461, specimen showing a small leaf. Scale bar equals 5 mm.
FIGURE 23. Wingia expansolobum (Upchurch and Dilcher) comb. nov. 1, UF15706-14828, specimen showing a multi-lobed leaf. Scale bar equals 1 cm. 2, Enlargement of Figure 23.1 (area indicated by the upper arrow) to show secondary and tertiary venation. Scale bar equals 1 mm. 3, Enlargement of Figure 23.1 (area indicated by the lower arrow) to show venation near the lamina margin. Scale bar equals 2 mm.
is often found in the Fabaceae (Satter and Morse, 1990; Rodrigues and Machado, 2008). A search using a subset of characters (vegetative morphology and anatomy: leaves (form)) in Intkey (Dallwitz et al., 1995) and using two characters (pulvinate and well-developed leaves) of Wingia expansolobum resulted in 11 families (Averrhoaceae, Barbulaeaceae, Empectraceae, Lardizabalaceae, Leguminosae, Marantaceae, Santalaceae, Sapindaceae, Staphyleaceae, Umbelliferae, and Zygophyllaceae) that have pulvinate leaves. When more characters (including leaf petiolate, simple, palmately-lobed, and primary vein palmate and in this order) are used, only two families, Leguminosae and Umbelliferae, remain. Both families do not seem to have similar leaf morphology as that of Wingia expansolobum. As indicated by Upchurch and Dilcher (1990), no extant species of Magnoliidae has five-lobed organization with basally palmate (actinodromous) primary venation. The modern affinities of Wingia expansolobum remain indeterminate but future collection of specimens with cuticular material may help clear this uncertainty.

Many recent discoveries of the Early Cretaceous fossil leaves with similar tooth pattern have been proposed to be related to the Eudicots. For example, Iterophyllum lobatum Barral et al. (Ranucucales; Barral et al., 2013) is a fossil species based on leaves from the late Barremian of Las Hoyas, Spain. The simple leaf is petiolate and palmately-lobed with chloranthoid-like glands at the lobe apices and the sinuses. Fairlingtonia thyrsopteroides (Fontaine) Jud (Jud, 2015) and Vernifolium tenuiloba Jud and Sohn (Jud and Sohn, 2016) from the Early Cretaceous Potomac Group in Maryland and Virginia, USA were also placed within the Eudicot clade, although their familial and ordinal affinities are unknown.

Wingia cf. expansolobum (Upchurch and Dilcher) comb. nov.

Figure 24

Description. Leaf simple, five-lobed, leaf 4 cm long and 8 cm wide (estimated maximum width); sinus deep, extending 75% to 80% of distance from apex to lamina base; medial lobe narrow ovate, symmetrical; lateral lobes asymmetrical, curved apically; apex of lobes obtuse or slightly acute; base of lamina obtusely cuneate or truncate; margin toothed; sinus rounded; serrations minute, simple; serration type ranging from concave to straight on apical side and straight to convex on basal side, spacing irregular; typically three veins entering a tooth, medial vein originating from exmedial side of secondary vein, basal vein tertiary in order and running very close to the margin before entering the tooth, vein of apical side not well developed. Petiole not observed. Primary venation basal actinodromous; primary veins thin, multi-stranded; medial primary vein straight in course; inner lateral primary veins apically curved; outer primary veins apically curved. Secondary venation semicraspedodromous; secondary veins thick relative to primary veins; originating from primary veins at moderate acute angle, slightly curved. One pair of secondary veins in each lobe originating from the extreme base of the leaf lamina and running almost parallel to the primary veins for almost half of the lobe length before joining the secondary veins originated from the primary veins within the lobes. Tertiary veins orthogonal reticulate. Veins of higher order not observed.

Number of species examined. 2.

Specimens illustrated. UF15705-14824 (Figure 24.1, 24.3); 24460 (Figure 24.2).

Remarks. These two specimens are placed in Wingia cf. expansolobum. They differ from Wingia expansolobum in having smaller leaves, relatively thin secondary veins, a pair of distinctive secondary veins originating from the extreme base of the leaf lamina and running parallel to the primary veins for almost half of the lobe length, semicraspedodromous secondary venation, and a predominately toothed margin. Even leaves of the same size (Figure 21.4) of Wingia expansolobum have different venation patterns. We currently separate these two specimens from Wingia expansolobum because of these differences.

Genus DICOTYLOPHYLLUM Saporta, 1894

Dicotylophyllum skogii sp. nov.

Figure 25

Specific diagnosis. Lamina thin. Base cordate. Margin entire. Petiole thin, multi-stranded. Primary venation pinnate; primary vein multi-stranded. Secondary vein thin, decurrent on primary vein, subopposite; angle of divergence moderate acute, uniform. Intersecondary veins common, simple. Tertiary veins thin, originating from secondary or intersecondary veins at right angles, forming rectangular or square meshes that are oriented with long axis parallel to secondary veins.

Description. Observed lamina 11 cm long and 5 cm wide; lamina very thin. Base cordate. Observed margin entire. Petiole thin, multi-stranded, 6 cm long and 1.5 mm wide. Primary venation pinnate; observed primary vein 1.5 mm wide, multi-stranded, slightly curved. Secondary venation not observed; secondary vein thin relative to primary
vein, 8 pairs observed, decurrent on primary vein, subopposite; angle of divergence moderate acute (ca. 45°), uniform. Intersecondary veins common, simple, 1 to 2 pairs between two adjacent secondary veins. Tertiary veins thin, originating from secondary or intersecondary veins at right angles, forming meshes rectangular or square in shape and oriented with long axis parallel to secondary veins. Veins of higher order not observed.

**Number of specimens examined.** 1.

**Holotype.** UF15706-24573 (Figure 25).

**Derivation of epithet.** In honor of Judith Skog in recognition of her contribution to Dakota paleobotany.
Remarks. The observed lamina is 11 cm long and estimated leaf length may be up to at least 15 cm. Although secondary venation pattern is not observed because of the fragmented nature of the specimen, the combined features of this leaf type, including thin lamina structure, thin primary and higher order veins, large spacing of secondary veins (eight pairs observed on 11 cm lamina), the presence of intersecondary veins, and tertiary venation patterns is distinctive from any other Dakota Formation angiosperm leaves. The multi-stranded primary vein and the presence of intersecondary veins indicate its possible affinity within the Magnoliidae. The thin lamina texture, tertiary and quaternary veins forming rectangular meshes with long axes parallel to secondary veins are observed in some extant herbaceous angio-

**Figure 25.** *Dicotylophyllum skogii* sp. nov. 1, UF15706-24573, fragment of leaf to show the petiole and strong primary vein. Scale bar equals 1 cm. 2, Enlargement of Figure 25.1 (area indicated by the upper arrow) to show secondary and intersecondary veins. Scale bar equals 2 mm. 3, Enlargement of the base of the leaf in Figure 25.1 (area indicated by the lower arrow) to show venation. Scale bar equals 2 mm.
sperms, indicating that *Dicotylophyllum skogii* may be an herbaceous plant which grows near water.

*Dicotylophyllum leptovenum* Wang and Dilcher, 2009

**Figure 26**

**Description.** Leaf base asymmetrical. Apex missing. Base acute, decurrent. Margin entire. Petiole short, stout, 1 cm long and 1.5 mm wide. Primary venation pinnate; primary vein massive, multi-stranded, course straight, observed primary vein about 1.2 mm wide. Secondary venation brochi-dodromous; secondary veins thin relative to primary vein, 6 pairs observed in a 6 cm long lamina; secondary veins originating from primary vein at wide acute angle (ca. 70º), joining superadjacent secondary veins at a distance of about one fifth of half lamina to enclose an intercostal region; secondary veins uniformly curved, but becoming sinuous when they intersect with intersecondary veins. Intersecondary veins always present, two or three per intercostal region, simple. Tertiary veins course irregular, intersecting with intersecondary veins or randomly reticulate, anastomosing to form meshes irregular in shape and size. Quaternary veins irregular in course, exmedially ramified or forming incompletely closed meshes irregular in shape and size. Veinlets simple, curved. Vein pattern in excosta region not observed.

**Number of specimens examined.** 1.

**Specimens illustrated.** UF15706-24734 (Figure 26).

**Remarks.** The specimen from Hoisington III locality has well-preserved high order veins compared with the specimen from the Courtland I locality, Minnesota.

**DISCUSSION**

Vegetation along the East Margin of the Cretaceous Western Interior Seaway (WIS) during the Mid-Cretaceous

During the latest Albian, angiosperms were widespread and replaced gymnosperms as the
WANG & DILCHER: EARLY CRETACEOUS LEAVES

dominant plants in many environments, as shown at the Hoisington III and other localities (Retallack and Dilcher, 1986). The Hoisington plant fossil assemblage is the best representative of a mixed deposit of several distinct, local plant communities (Retallack and Dilcher, 1981a, 1981b; Retallack and Dilcher, 2012). A swamp woodland vegetation probably included Archaeaanthus linnenbergeri (or Liriodendron kensense; Crane and Dilcher, 1984; Dilcher and Crane, 1984), Lesquerella elocata (Crane and Dilcher, 1984); Calodax delevoryana (Dilcher and Kovach, 1986); Sapindopsis bagleyae, Sapindopsis reallackii, and other members of the Platanaceae; Prisca reynoldsii and members of the Lauraceae, and conifers. Prominent understory plants include ferns, Anemia dicksoniana (Heer) Krassilov, Anemia dactotensis (Rushforth) Skog and Dilcher, Gleichenia camptophylla (Debenham and Ettinghausen) Heer, Gleichenia deliculata Heer, Matonidium brownii Rushforth (Skog and Dilcher, 1994; Hu et al., 2006) and some herbaceous angiosperms (e.g., Skogia leptoselis). Free floating aquatic plants in the open lake water include Marsilea johnhallii (Skog and Dilcher, 1992), Aquatofila fluitans, and Brasenites kensense (Wang and Dilcher, 2006a). Submerged aquatic plants, e.g., Donlesia dactotensis (Dilcher and Wang, 2009), an extinct member of the Ceratophyllaceae from the coeval Dakota Formation localities, probably also occupied some of the habitat in the fresh water lakes. The semi-aquatic plant, Isoetites phyllophila (Skog and Dilcher, 1994), lived in shallow water or on wet margin that may have experienced short-term drying.

Among all localities from the Dakota Formation, angiosperm leaves are the most species-rich at the Hoisington III locality based on raw counts of species from a collection of 1,500 specimens stored in the Paleobotany collection at the Florida Museum of Natural History. Six species of ferns, one species of quillworts, and six species of conifers have been reported from this locality (Table 3). This further confirms the observation that flowering plants were the dominant group in the local floras along the Cretaceous Western Interior Seaway. The abundant conifer pollen in the microfossil record represents the inland and more elevated floras to the east (Retallack and Dilcher, 1986).

Comparisons of Angiosperm Species Richness among Dakota Localities

Compared with the Braun Ranch locality in Kansas (Wang and Dilcher, 2006a), the Rose Creek locality in Nebraska (Upchurch and Dilcher, 1990), and the Courtland I locality in Minnesota (Wang and Dilcher, 2009), the Hoisington III locality is the most species-rich in angiosperms with 24 species (Table 1). The Hoisington III locality and the Rose Creek I locality share four species, i.e., Crassidenticum decurrens, Pabiania variloba, Anisodomum wolfei, and Wingia expansolobum. The Hoisington III and Courtland I also have four species in common, i.e., Jarzenia kanbrasota, Rogersia dactotensis Wolfiophyllum paffianum, Dicorytophyllum leptovenum, and Credneria cyclophylla. Of the 70 species recognized from the four localities, only one species, i.e., Crassidenticum decurrens is shared by three localities (Hoisington III, Rose Creek and Braun Ranch). In summary, there is little species overlap between any two localities of the Dakota Formation leaf assemblages. These leaf assemblages are all Dakota time (late Albian to early Cenomanian) equivalent. However, they do not represent floras living at exactly the same time because the deposition of the Dakota Formation extended over at least five million years (Koch and Brenner, 2009). At present, there is no reliable dating of these megafossil localities. Therefore, their relative ages are uncertain. It is not known how the time difference may have affected the diversity pattern of the leaf assemblages among these localities.

Effects of Environments on the Diversity of the Dakota Flora

Paleoenvironmental interpretation of the four localities by various authors indicates that angiosperm leaf megafossils represent floras that lived under different environments. For example, Farley and Dilcher (1986) interpreted the Braun’s Ranch locality as marsh lakeside and the Rose Creek I locality as tidally influenced distributary margin. The Hoisington III locality was interpreted as a fresh water lake or lagoon environment with river influence (Retallack and Dilcher, 1981a, 1981b; Skog and Dilcher, 1992). It has not been possible for us to resolve the relative age of these localities with sufficient precision in order to place them in chronological sequence in the stratigraphic column. This makes it difficult to assess the effects of floristic changes through time on the evolution/diversity of the Dakota flora. At present, the small number of overlapping species might be explained by the environmental differences at the various localities (Table 1). Similar environments, even though separated by a great geographical distance (or with a great time difference with similar paleoclimate condition, see discussion below), tend to
have more similar taxa than two different environments with a distance of only several kilometers (Kvacek and Dilcher, 2000).

**Comparison with the Palynological Record**

The high diversity of angiosperm leaf megafossils at different localities is in conflict with the microfossil record (Farley and Dilcher, 1986; Dilcher and Farley, 1988), which indicates a high percentage of non-angiosperm palynomorph abundance. This was interpreted as resulting from the dominance of ferns and gymnosperms in the upland and coastal regional floras and the dominance of angiosperms in local coastal environments (Skog and Dilcher, 1994; Kvacek and Dilcher, 2000). It has also been suggested that the Dakota Formation was formed at a time just prior to the development of wind dispersed angiosperm pollen (Kvacek and Dilcher, 2000). All these suggest that the high diversity of angiosperms from different localities is controlled by environment, rather than being a sampling effect (Farley and Dilcher, 1986), which further suggests successful competition of angiosperms in coastal environments with gymnosperms and ferns during that time (Retallack and Dilcher, 1986; Kvacek and Dilcher, 2000).

**CONCLUSIONS**

In this report, we recognize 24 fossil leaf species from the Hoisington III locality, Kansas, USA. This leaf assemblage represents the highest species richness of the four Dakota Formation localities (Braun Ranch and Hoisington III, Kansas, Courtland I, Minnesota, and Rose Creek, Nebraska). So far, a total of 70 fossil leaf species have been recognized from these four localities (Table 1). There is little overlap of angiosperm leaf species between any two Dakota Formation localities. At present, the high diversity of angiosperms that is locality specific and the little species overlap between any two Dakota Formation localities can best be interpreted as the evolution of angiosperm diversity, each being unique to the different environments in which they grew (Retallack and Dilcher, 1986).

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