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Article abstract

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Cite this article

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ABSTRACT

New collections of plant macrofossils provide a precise Middle Pennsylvanian age for the lower Wamsutta Formation red beds of the Narragansett Basin, southeastern Massachusetts, USA. The Wamsutta assemblage indicates strong correlation with the Cumberland Group of the Maritime Provinces of Canada, which was earlier considered to be of late Langsettian to early Duckmantian age. This correlation is supported by the presence in the Wamsutta Formation of the following plant-fossil species: Neuralethopteris schlehanii, Neuropteris obliqua, Senftenbergia plumosa, Calamites suckowii, Annularia asteris, Annularia cf. microphylla, Asterophyllites charaeformis, Asterophyllites grandis, Asterophyllites lindleyanus, Sphenophyllum cuneifolium, and Sphenopteris valida. Moreover, the new fossil flora resembles the Middle Pennsylvanian floras of western Europe, such as the Laveineopteris loshii Subzone. The new flora is especially similar to those of the Iberian Peninsula, where there is a complete succession of Carboniferous macrofloral zones, and this similarity confirms a late Langsettian or early Duckmantian age for the lower Wamsutta macroflora. The new collections of Wamsutta Formation plant fossils, along with a smaller existing collection, represent the oldest known macroflora in the Narragansett Basin.

INTRODUCTION

The Wamsutta Formation red beds of the Narragansett Basin contain sparse plant fossils. Shaler et al. (1899) reported Calamites and Cordaites, which demonstrate that the Wamsutta Formation is of Carboniferous (Pennsylvanian) age, like the overlying Rhode Island Formation. Further substantial progress on the paleobotany of the Wamsutta...
Formation was not made until Knox (1944) reported the following florule from the lower Wamsutta Formation: Annularia stellata; Calamocladus (= Asterophyllites longifolius); Calamites equisetiformis (= Asterophyllites equisetiformis); Calamites suckowii; Pecopteris sp.; Neuropteris cf. N. raritervis (probably a misidentification); Sphenopteris valida; Sphenopteris sp.; Oligocarpa splendens; Cordaites cf. C. robbi; Cordaites cf. C. communis; "Sigillarian leaves" (lycopsid leaves = Cyperites).

Two problems exist with Knox's (1944) collection. First, he provided no photographs or drawings of the specimens that he collected to help validate his identifications. Second, we do not know where Knox's collection is located. A recent thorough search of the paleobotanical collections at Harvard University revealed no sign of the collection ever having been housed there. Inquiries as to whether the collection perhaps had been deposited in the Tufts University collections or included in the Brown University Paleobotanical Collections (and possibly relocated to the University of Philadelphia) were also unsuccessful.

Until now, no additional paleobotanical collections have been assembled with the aim of establishing the age of the Wamsutta Formation red beds in the Narragansett Basin. However, Oleksyshyn (1976) derived an Asturian age for the Rhode Island Formation based on macroflora from an open-pit quarry; the Rhode Island Formation interfingers with and overlies the Wamsutta Formation. Also, Lyons et al. (1976) recorded a florule from the Upper Pondville Conglomerate, which underlies and interfingers with the Wamsutta Formation, in the adjacent Norfolk Basin; they assigned it a late Duckmantian age.

A summary of the new fossil-plant collections from the lower Wamsutta Formation in the Narragansett Basin, we illustrate each of the plant-fossil taxa recognized in Knox (1944), and re-evaluate his identifications. Lastly, we establish the age of the lower Wamsutta Formation macroflora on the basis of correlations with similar, poorly understood Middle Pennsylvanian macrofloral zones in the Maritime Provinces and western Europe.

### GEOLOGICAL CONTEXT

The Narragansett Basin of Massachusetts and Rhode Island (Fig. 1) is a composite, intermontane rift basin formed from the coalescence of small grabens as Laurentia and the Avalon microcontinent collided during the Paleozoic (Shaler et al. 1899; Mosher 1983). The Basin is primarily infilled with terrestrial sedimentary rocks, and depositional environments include alluvial fans, braided and meandering streams, floodplains, swamps, and freshwater ponds (Shaler

<table>
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<th>Table 1.</th>
<th>Wamsutta taxa systematically arranged, with semi-quantitative (not census sampled) occurrence information. VC = Very Common, C = Common, R = Rare, VR = Very Rare.</th>
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<tr>
<td><strong>Order Medulales</strong></td>
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<tr>
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<tr>
<td>Neuropteris schlepianii</td>
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<td>Family** Cordaitales**</td>
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<tr>
<td>Laveineopteris cf. loshii</td>
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<td>Family** Cyclopteridaceae**</td>
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<td>Cyclopteris sp.</td>
<td><strong>VR</strong></td>
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<td>Family Neurodendopteridaceae</td>
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<td>Neuropteris cf. obliqua</td>
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<td><strong>Order Lyginopteridales</strong></td>
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<td>Family** Lyginopteridaceae**</td>
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<td>Mariopteris sp.</td>
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<td>Family** Botryopteridales**</td>
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<td>Senftenbergia plumosa</td>
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<td>Family** Filicophyta insertae sedis**</td>
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<td>Sphenopteris valida</td>
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<td><strong>Order Equisetales</strong></td>
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<td>Family** Calamitaceae**</td>
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<td>Calamites suckowii</td>
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<td>Annullaria asteris</td>
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<td>Annullaria cf. microphylla</td>
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<td>Annullaria radiata</td>
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<td>Asterophyllites charaeformis</td>
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<td>Asterophyllites grandis</td>
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<td>Asterophyllites lindleyanus</td>
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<td>Asterophyllites longifolius</td>
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<td>Calamostachys sp.</td>
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<td>Sphenophyllum cf. cuneifolium</td>
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<td>Family** Sigillariaceae**</td>
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<td>Sigillaria sp.</td>
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<td>Cyperites sp.</td>
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et al. 1899; Mosher 1983, and references therein; Knecht et al. 2011; Getty et al. 2013). Recent radiometric dating of rhyolite near the base of the Wamsutta Formation (one of the lowermost units of the Basin) suggests that crustal extension began in the Late Devonian, about 373 Ma (Maria and Hermes 2001; Thompson and Hermes 2003). Previously, most authors (e.g., Shaler et al. 1899; Towe 1959; Quinn and Oliver 1962; Skehan et al. 1979; Mosher 1983) had considered basin formation to have begun with the deposition of the Pondville Conglomerate in the Early Pennsylvanian. The Pondville Conglomerate, the basal unit in the Narragansett Basin, is 150 m thick and consists of grey to greenish conglomerate containing quartzite, granite and schist clasts up to 60 cm thick interbedded with red and green slate, arkose, and siltstone (Shaler et al. 1899; Skehan et al. 1979). The conglomerate sometimes grades upward into grey sandstone that may grade into red Wamsutta Formation beds (Skehan et al. 1979) (see below).

Most strata in the Narragansett Basin comprise the Rhode Island Formation, which overlies and interfingers with the Wamsutta Formation. The Rhode Island Formation is almost 3000 m thick and consists of grey polymictic conglomerate, grey quartz-rich sandstone and siltstone, grey to black shale, and coal seams up to 10 m thick (Lyons 1977; Skehan et al. 1979; Murray et al. 2004). A rich variety of fossil plants and, to a lesser extent, faunal ichnofossils and insect wings, occur in the formation (Scudder 1893; Oleksyshyn 1976; Sproule and Getty 2012; Getty et al. 2013, 2017). Biostratigraphic analyses of the fossil flora indicate that the age of the Rhode Island Formation ranges from early Bolsovian to Stephanian B or C (Oleksyshyn 1976; Skehan et al. 1979; PCL and RGS unpublished data).

The uppermost lithostratigraphic unit in the Narragansett Basin is the Dighton Conglomerate, which is 300–450 m thick and correlative to the Purgatory Conglomerate in the southern Narragansett Bay (Skehan et al. 1979; Lynn and Kroll 2006). The Dighton Conglomerate, which extends in age almost up to the Permian, is the highest unit in the Basin (Mosher 1983; Mosher and Rast 1984; Wagner and Lyons 1997; Rich 2006) and is composed of coarse-grained sandstone with quartzite cobbles, some of which display molds of Cambrian brachiopods (Shaler et al. 1899; Skehan et al. 1979).

The Wamsutta Formation red beds overlie the Pondville Conglomerate and are composed of conglomeratic to arkosic sandstone, cross-bedded medium- to fine-grained sandstone, siltstone, and red shales that are approximately 300 m thick and rare freshwater limestone (Shaler et al. 1899; Mutch 1968). In the Narragansett Basin, the Wamsutta Formation includes a significant number of volcanic
deposits, and felsic clasts are common in conglomeratic layers (Skehan et al. 1979; see Fig. 2a). Maria and Hermes (2001) and Thompson and Hermes (2003) considered the volcanic rocks to be of Devonian age based on U–Pb dating, which suggests that an unconformity is present within the formation. One of us (PCL) has seen that the contact between the volcanic rocks and the overlying portions of the Wamsutta Formation is gradational, and thus we consider the volcanic rocks to be of Pennsylvanian age, like the rest of the formation. The red beds within the Wamsutta Formation, which originated as a series of well-drained alluvial fans, include several horizons with plant fossils. However, deposits with compressed, carbonized (coalified) flora representing anoxic, swampy environments are absent (Shaler et al. 1899; Knox 1944; Lyons and Chase 1976).

The Wamsutta study area, from which the new fossil-plant collections were obtained, comprises several small, closely spaced outcrops on the southern flank of Red Rock Hill in North Attleboro, Massachusetts (Fig. 2b), the type locality of the Wamsutta Formation. The outcrops lie about in the middle of that part of the lower Wamsutta succession exposed locally, and are stratigraphically 100 m or more above

Figure 2. (a) Exposure of the Wamsutta Formation in study area. Note granite (light-colored) and volcanic (dark) clasts in conglomerate with coarse sandstone at upper right. Geologic hammer gives scale. (b) Exposure of Wamsutta Formation in study area. Note steeply dipping, laminated sandstone (lower left centre) overlain by thick conglomerate. Height of hand sledge is 40 cm. (c) Amphibian footprints with biserial Diplichnites trackway (arrowed; lower left). Note fossil seed fern (arrowed; Neuropteris?) just above footprint (arrowed) at lower right. Collection of RGS. Scale equals 3 cm. (d) Tonganoxichnus sp., an archaegnathan insect ventral impression. Collection of RGS. Scale in millimetres. (e) Neopteran insect wing (Order Blattaria?). Collection of RGS. Scale in millimetres.
outcrops of the basal felsite flows (Knox 1944). The north-easterly to northwesterly trending strata are capped by laterally continuous conglomerate with coarse-grained, poorly sorted, cross-bedded sandstone 3 m or more thick. Beds dip east-northeast at about 60 degrees in the northwestern part of the study area, and northwest at 35 degrees in the south-eastern margin.

Underlying the study area’s coarse strata are thin sequences of ripple-laminated, medium- to fine-grained sandstone beds, graded in cross-section and draped with red shale, that indicate recurrent flooding (and drainage) by shallow streams on a low-relief floodplain. Plant detritus, especially cordaitalean leaves, sphenophyte stems, and pteridosperm pinnae, are sometimes preserved on bedding-plane surfaces, as are sedimentary structures including raindrop imprints, asymmetric current ripples, wrinkled mud, tool marks, pressure casts ("mud bubbles"), gas-escape structures, and occasional desiccation cracks. Entombed stem casts with whorled leaves and/or plant stem cross-sections with pin-wheeled eddy patterns frequently poke through these substrates, indicating the presence of upright sphenophytes and other plants in growth position. Red shale predominates in a few areas and forms distinct plant-fossil facies with thicknesses of up to approximately 1 m. Accompanying the fossil plants are well-preserved ichnofossils, such as numerous footprints of reptiles and basal amphibians (Fig. 2c), plentiful biserial Diplanchites trackways (referred to myriapods) (Fig. 2c), other invertebrate traces including four ephemeropteran and 11 archaeognathan insect impressions (Fig. 2d), and a rare neopteran (order Blattaria?) insect wing (Fig. 2e). Thin lithified paleosols lacking in fossil macroflora may underlie and/or overlie the fossiliferous fine-grained sandstone and shale horizons. All told, the Wamsutta Formation study area displays many of the characteristics of a moist, sub-humid (seasonally dry to seasonally wet), clastic wetland habitat on an alluvial fan landscape lush with tropical forestation (see Bashforth et al. 2014).

MATERIAL AND METHODS

Simple hand sledges and pry bars were used to loosen and extricate specimens from two low-lying, unexploited, well-indurated ledges exposed in the study area on Red Rock Hill (41.937° N, 71.353° W; 41.939° N, 71.354° W); the latter ledge produced all of the insect fossils. Knox’s (1944) fossil-flora locality, a road cut along the west side of U.S. Route 1 near the Attleboro–North Attleboro town line, is about 250 m to the southeast of the study area. Approximately 100 m north of the study area, fossil tetrapod footprints have been reported (Willard and Cleaves 1930; Knecht and Benner 2008) along with an ephemeropteran insect impression, several archaeognathan insect impressions, and at least two insect wings (Knecht et al. 2011; Benner et al. 2015).

The new collection of the lower Wamsutta Formation macroflora is reposited in the Harvard University Paleobotanical Collections (HUPC) in Cambridge, Massachusetts. HUPC accession numbers have been assigned that refer to the slabs on which the specimens are preserved. Images of key and/or uncertain plant species were sent to paleobotanists for their opinions on identification. Specimens were photographed in natural sunlight with an Olympus VR 350 16-megapixel camera. Measurements were made directly from the specimens and are given in standard international (SI) metric system units.

SYSTEMATIC PALEONTOLOGY

Order Medullosales Corsin 1960
Family Alethopteridaceae (Lesquereux 1884) Corsin 1960 emend. Anderson et al. 2007

Genus Neuralethopteris Cremer 1893

Neuralethopteris schlehanii (Stur 1877) Cremer 1893 (Figs. 3a–b)

Remarks. Pinnules are usually tongue-shaped with a rounded apex, and have a cordate base in some parts, or are stalked or attached to the base in other areas, but are not recurrent or confluent as in alethopterids. A strong midvein nearly reaches the pinnule apex, and lateral veins extend at an acute angle,forking two or three times to meet the margin almost perpendicularly (Goubet et al. 2000), as opposed to meeting the margin obliquely as in neuropterids such as Neuropteris obliqua and Laveineopteris loshii. The rachis is striated longitudinally (Bell 1944).

Neuralethopteris schlehanii is a common component of Namurian and Langsettian tropical rainforest florules in Europe, and is a key arborescent species, perhaps 3 m tall, in similar Langsettian-age florules from the Cumberland Group of the Maritime Provinces (Bell 1944; Falcon-Lang et al. 2006). Neuralethopteris schlehanii has been seldom recorded in the United States (Goubet et al. 2000), and is recorded herein for the first time in the Narragansett Basin. Sizable frond fragments of Neuralethopteris schlehanii were found in the lower Wamsutta littering bedding planes of shallow, low-energy, sand-bed channels and low-relief fluvial embankments, in association with Cordaites cf. principalis, Laveineopteris cf. loshii, Calamites suckowii, and Asterothylites lindleyanus. Neuralethopteris schlehanii became abruptly extinct in western Europe at the end of the Langsettian and is, therefore, very useful biostratigraphically (Goubet et al. 2000).

Family Cyclopteridaceae Corsin ex Wagner 1964

Genus Laveineopteris Cleal et al. 1990

Laveineopteris cf. loshii (Brongniart 1831) Cleal et al. 1990 (Figs. 3c–d)

Remarks. J.-P. Laveine (personal communication) was
uncertain of the species designation; consequently, we only tentatively assign the Wamsutta specimens to *Laveineopteris loshii*. The median vein is strong; the lateral veins arise acutely, are dense, bifurcate acutely several times, and meet the margin at oblique angles. The terminal pinnule is deltoid, and subjacent pinnules may be broadly attached (Cleal and Shute 2003). Large fronds grew on arborescent plants that may have approached the local forest canopy in height.

Associated cyclopterid shade leaves may be preserved nearby (Cleal and Shute 2003).

Although well known throughout the Duckmantian substage, *Laveineopteris loshii* first appears and becomes common in the wetland deposits of middle Langsettian tropical rain forests. As a result, Cleal (1991) proposed a biostratigraphically useful floral subzone, the "*Laveineopteris loshii* Subzone", in the Langsettian substage, based in part on this

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**Figure 3.** (a) HUPC No. 64,894, *Neuralethopteris schlehanii*. Specimen identified by Jean-Pierre Laveine, Lille, France. (b) Close-up of HUPC 64894, *Neuralethopteris schlehanii*. Note alethopteroid venation and non-decurrent, non-confluent pinnules attached to rachis. (c) HUPC No. 64,846A, *Laveineopteris cf. loshii*. Note well-preserved venation. Some pinnules appear to taper near the base. Specimen identified by Jean-Pierre Laveine, Lille, France. (d) HUPC No. 64,882, *Laveineopteris cf. loshii*. Note large deltoidal terminal pinnule and subjacent pinnules. Specimen identified by Jean-Pierre Laveine, Lille, France. (e) HUPC No. 64,870, *Cyclopteris* sp., incomplete orbicular shade leaf. All scales in millimetres.
initial appearance (see also Cleal and Thomas 1994).

In the Wamsutta Formation, sizable frond fragments of *Laveineopteris cf. loshii* (up to at least 20 cm long) are occasionally encountered as well-preserved parautochthonous detritus on the bedding planes of shallow-channel sandstones and adjacent floodplain shales. Large, broad pinnules with full-bodied, ovate pinnules distinguish *Laveineopteris loshii* from the smaller pinnules and lanceolate/lingulate with detritus occasionally *Laveineopteris cf.*. *loshii*. The absence of lobed pinnules distinguishes *Laveineopteris loshii* from species of *Mariopteris*.

Genus *Cyclopteris* Bronn 1828a

*Cyclopteris* spp.  
(Fig. 3e)

Remarks. Isolated, orbicular and kidney-shaped leaves that display radiating venation are characteristic of the fossil-genus *Cyclopteris* (Janssen 1965; Cleal and Shute 2003). *Cyclopteris* pinnules were attached to those parts of adult trees below the canopy, which were probably shade leaves. Sun leaves, such as *Laveineopteris loshii* fronds, may have grown high above on the same tree (Cleal and Shute 2003). The isolated and detached nature of *Cyclopteris* pinnules, combined with characteristic radiating venation, easily distinguishes them in the field.

Only one example of *Cyclopteris* has been found in the Wamsutta Formation red beds. By contrast, the associated sun leaves, *Laveineopteris cf. loshii* pinnules (Figs. 3c–d), have been found as preserved litter on bedding planes in several fluvial wetland horizons.

Family *Neurodopteopteridaceae* Laveine 1966

Genus *Neuropteris* (Bronn 1822) Sternberg 1825 emend. Cleal et al. 1990

*Neuropteris obliqua* (Bronn 1833–1834) Zeiller 1886  
(Figs. 4a–b)

Remarks. This species of arborescent seed fern is characterized by mixoneuroid venation, with the lateral pinnules near the apices of the pinnae united to the rachis, and a small, lanceolate terminal pinnule (Bell 1944; Darrah 1969; Josten 1991). Pinnules may be highly variable in morphology (Darrah 1969).

*Neuropteris obliqua* is distinguished from species of *Mariopteris* in not having lobed pinnules, from *Neuropteris schlehanii* by its shorter midvein and acutely rising lateral veins, and from *Laveineopteris loshii* by its smaller, more lanceolate terminal pinnule and less robust subjacent pinnules.

*Neuropteris obliqua* is most frequently encountered in late Langsettian and early Duckmantian rainforest florules worldwide, such as those found in the Iberian Peninsula of western Europe (Cleal 2005; Wagner and Álvarez-Vázquez 2010), and in the Cumberland Group in the Maritime Provinces (Bell 1944; Falcon-Lang et al. 2006; Falcon-Lang et al. 2010; Bashforth et al. 2014). *Neuropteris obliqua* is the dominant seed fern found in the Wamsutta Formation in the study area. It was approximately 3–5 m tall and usually preserved as parautochthonous (full or partial) lone pinnae, isolated pinnules, or rarely as multiple attached pinnae compressed/adpressed upon bedding planes of mudstone and shale representing low-energy channel, proximal flood-plain, and soggy non-swamp settings.

Order *Lyginopteridales* (Corson 1960) Havlena in Dvořák and Růžička 1961

Family *Lyginopteridaceae* Potonié 1900 emend. Anderson et al. 2007

Genus *Mariopteris* Zeiller 1879

*Mariatorpseris* sp.  
(Fig. 4c)

Remarks. Pinnae from this genus of arborescent seed fern are characterized by pinnules, entire or lobed, with broad bases. Pinnules may exhibit a thick, leathery or coriaceous appearance (Janssen 1965; Darrah 1969). Lobed pinnules that lack a conspicuous midvein distinguish *Mariopteris sp.* from *Neuropteris obliqua* and *Neuralathopteris schlehanii*. Generally, *Mariopteris* sp. pinnules are less segmented than those of *Sphenopteris*, and their venation may be more coarse and conspicuous (Janssen 1965).

*Mariopteris* sp. ranges through the Westphalian, and is represented in the new Wamsutta collections by a single specimen, collected in a sandstone reflecting a shallow-channel setting along with specimens of *Laveineopteris cf. loshii*, *Cordaites principalis* and *Neuropteris obliqua*.

*Mariopteris* sp., sensu lato Zeiller 1879  
(Fig. 4d)

Remarks. This form has similar characteristics to the above form described as *Mariopteris* sp. Pecopteroid or (less commonly) sphenopteroid sub-triangular pinnules are attached for the full width of the base, are more or less united, and are often decurrent (Darrah 1969). The largest pinnules may display a cordate, sessile base (Bell 1944).

Only two specimens have been collected by the authors to date (Figs. 4c–d), both retrieved on bedding planes of shale-draped sandstone deposited in shallow, low-energy channels.
small, narrow, crenulated pinnules distinguish it from other fern-like taxa in the Wamsutta Formation. *Senftenbergia plumosa* was an abundant Late Carboniferous fern that historically had long been assigned to family Pecopterideae, as *Pecopteris plumosa*. Only recently has *Senftenbergia plumosa* been recognized as a representative of the Late Carboniferous fern family Tedeleaceae (Pšenička and Bek 2001).

According to some authors (e.g., Brousmiche Delcambre et al. 1998), *Senftenbergia plumosa* ranged throughout the Carboniferous, but Cleal (2005) gave it an early Langsettian to middle Bolsovian age. *Senftenbergia plumosa* is abundant and widespread in the Cumberland Group of the Maritime Provinces, and is also a common representative of the

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**Order Botryopteridales** Meyen 1987  
**Family Tedeleaceae** Eggert and Taylor 1966

**Genus Senftenbergia** Corda 1845

*Senftenbergia plumosa* (Artis 1825) Stur 1885  
emend. Bek and Pšenička 2001

(Fig. 4e)

**Remarks.** Pinnae are small, and so venation may be difficult to discern, but the margins of pinnules are crenulated. The midvein is slightly flexuous, a branch entering a lobe divides once, with the upper arm perhaps dividing again (Bell 1944). The small size of the lanceolate pinnae and the very small, narrow, crenulated pinnules distinguish it from other fern-like taxa in the Wamsutta Formation.

*Senftenbergia plumosa* was an abundant Late Carboniferous fern that historically had long been assigned to family Pecopterideae, as *Pecopteris plumosa*. Only recently has *Senftenbergia plumosa* been recognized as a representative of the Late Carboniferous fern family Tedeleaceae (Pšenička and Bek 2001).

According to some authors (e.g., Brousmiche Delcambre et al. 1998), *Senftenbergia plumosa* ranged throughout the Carboniferous, but Cleal (2005) gave it an early Langsettian to middle Bolsovian age. *Senftenbergia plumosa* is abundant and widespread in the Cumberland Group of the Maritime Provinces, and is also a common representative of the
arborescent wetland macroflora in the lower Wamsutta Formation in the study area. Although one frond fragment has been collected, most Wamsutta specimens of this taxon are found as detached parautochthonous pinnae in fine-grained sandstones representing shallow sand-bed channels and their proximal low-relief embankments and floodplains.

Order Filicales Engler & Prantl 1902
Family Filicophyta incertae sedis

Genus Sphenopteris (Brongniart 1822) Sternberg 1825

Sphenopteris valida Dawson 1868

Remarks. Lacy lobate pinnules branch off on each side and bifurcate at acute angles, are alternate, petiolate and obtuse, and may taper gradually to a point (Janssen 1965; Josten 1991). A single vein enters the pinnule from the rachis, forming several times such that one or two veins enter each lobe of the distal margin (Janssen 1965). Sphenopteris valida differs from Neuropteris obliqua, Laveineopteris loshii, and Neuraletthopteris schlehanii in having lobate pinnules without a prominent midvein, and from Senftenberga plumosa in having lobate, non-crenulated pinnules. Marioperis sp. does not have petiolate pinnules.

Biostratigraphic ranges given by authors for this species vary, but Wagner and Álvarez-Vázquez (2010) list Sphenopteris valida (= Sphenopteris artemisiacifoliiodes; see Stopes 1914; Bell 1944) as ranging primarily from Langsettian to Duckmantian. Reported previously by Knox (1944) and found typically in the Wamsutta Formation as well as lying horizontally upon bedding plane surfaces. Occasionally, these casts are transversely imprinted with needle-like leaves of Asterophyllites longifolius.

Annullaria asteris Bell 1944

Remarks. This seldom accurately reported species is characterized by very small, rather stiff, straight-sided leaflets 2 to 4.5 mm long with acuminate tips. Up to ten isophyllous, narrow leaflets only slightly constricted at the base may comprise a verticil (Bell 1944; Álvarez-Vázquez and Wagner 2017). Ultimate branches are very slender (Bell 1944); two sizes of leaflets may be displayed on branches of the same plant (Abbott 1958). Annullaria asteris is distinguished from other whorled taxa by being the most diminutive; its stiff, straight leaves separate it from the slightly longer, sickle-shaped leaves of Annullaria microphylla.

Annullaria asteris is very common in the Wamsutta Formation in the study area, where it probably grew as luxuriant groves of short groundcover in abandoned channels, on flanks of channels, and/or as rapid recovery growth on proximal floodplains of low-energy streams. Annullaria asteris is sometimes associated with Asterophyllites charaeformis (Álvarez-Vázquez and Wagner 2017) and, interestingly, a similar association is noted in two new Wamsutta specimens. Reportedly ranging from Kinderscoutian to Bolsovian in western Europe (Wagner and Álvarez-Vázquez 2010; Álvarez-Vázquez and Wagner 2017), Annullaria asteris is reported herein for the first time in the Narragansett Basin.

Annullaria cf. microphylla Sauveur 1848

Remarks. Annullaria microphylla is marked by longitudinally striate ultimate axes, with anisophyllous verticils displaying 12–16 small, sickle-shaped leaflets 5–6 mm in length. The taxon is only slightly larger in size than Annullaria asteris, but Annullaria microphylla may also be distinguished from Annullaria asteris by its sickle-shaped leaflets with characteristic thick, vaunted laminae (Álvarez-Vázquez and Wagner 2017). The Wamsutta specimens are only tentatively assigned to this species due to their poor preservation.

The species has not been recorded commonly in Europe or the Maritime Provinces, but Wagner and Álvarez-Vázquez (2010) and Álvarez-Vázquez and Wagner (2017) reported it from Langsettian to lower Duckmantian strata. Annullaria cf. microphylla is also an uncommon species in the
Wamsutta Formation in the study area, but like *Annularia asteris* probably grew as low shrubs along shallow stream embankments and proximal low-relief floodplains. This is the first record of specimens associated with *Annularia microphylla* in the Narragansett Basin.

*Annularia radiata* (Brongniart 1822) Sternberg 1825  
(Fig. 5e)

**Remarks.** Linear lanceolate leaves 11–24 mm in length, 12–16 per verticill with an acute apex characterize this species (Álvarez-Vázquez and Wagner 2017). Isophyllous whorls are spread out on the substrate but do not overlap. *Annularia radiata* is sometimes confused with *Annularia stellata*, but is easily distinguished, since the latter species displays distinctly larger anisophyllous whorls of leaves (Abbott 1958; Álvarez-Vázquez and Wagner 2017).

*Annularia radiata* occurs most often in upper Langsettian and Duckmantian strata (Álvarez-Vázquez and Wagner 2017). In the Wamsutta Formation, sizable paraautochton-

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**Figure 5.** (a) HUPC No. 64,893A, *Sphenopteris valida*. Elongate-ovate pinnules are characteristic. Scale equals 3 cm. (b) HUPC No. 64,872, *Calamites suckowii*. Internodes shorter than long and coarse ribs alternate at nodes. Scale equals 3 cm. (c) HUPC No. 64,863B, *Annularia asteris* showing two sizes of very small, rigid leaflet. Scale equals 3 cm. (d) HUPC No. 64,839, *Annularia cf. microphylla*. Whorled leaflets are 5–6 mm in length. Scale equals 8 cm. (e) HUPC No. 64856A, *Annularia radiata*. Leaflets are linear lanceolate with acute apices. Scale in millimetres.
ous groups of whorled leaves sometimes occur on clay-draped channel-sandstone substrates, though two- or three-whorled branch fragments are more common. Workers in the Maritime Provinces and in the Narragansett Basin area (mis)applied the synonym *Annularia acicularis* to specimens of *Annularia radiata*, or to specimens too fragmentary (e.g., Oleksyshyn 1976, p. 156, figs. 7B-7D) for precise identification (Álvarez-Vázquez and Wagner 2017). Perhaps this explains why *Annularia radiata* has not been reported previously in the Narragansett Basin.

**Genus Asterophyllites** Brongniart 1822

*Asterophyllites charaeformis* (Sternberg 1825)
Göppert 1844
(Fig. 6a)

**Remarks.** Verticils with up to 10 hook-shaped, pointed leaves 2–3 mm in length (as long as, or slightly longer than, the internodes), and the compact nature of the shoot combine to make this smallest *Asterophyllites* species distinctive. Axes are slender and can be striate vertically (Bell 1944; Álvarez-Vázquez and Wagner 2017). The sickle-shaped leaves of *Asterophyllites grandis* are somewhat larger than those of *Asterophyllites charaeformis*, and distinguish the two species.

In the macrofloras of western Europe, the Iberian Peninsula, and the Cumberland Group of the Maritime Provinces, *Asterophyllites charaeformis* is recorded as ranging from late Namurian to Bolsovian (Wagner and Álvarez-Vázquez 2010; Álvarez-Vázquez and Wagner 2017). *Asterophyllites charaeformis* is a fairly common taxon in the Wamsutta Formation, and likely grew as diminutive groundcover in abandoned low-energy channels, and on low-relief floodplains and interfluves. The close resemblance of two of this study’s specimens to Matthew’s (1906) specimen, with *Annularia asteris* nearby, is notable (see Álvarez-Vázquez and Wagner 2017, fig. 13).

*Asterophyllites grandis* (Sternberg 1825) Geinitz 1855
(Fig. 6b)

**Remarks.** Longitudinally striate axes with internodes 2.5–4 mm long characterize this species. Curved, sickle-shaped leaves 4–6 mm in length are distinctive and overlap slightly at successive whorls (Álvarez-Vázquez and Wagner 2017). Commonly, only two opposite, uninnerved leaflets in a verticil are preserved (Bell 1944). Here recorded for the first time in the Narragansett Basin, *Asterophyllites grandis* is noticeably larger and more robust than *Asterophyllites charaeformis* (Abbott 1958), with which it may sometimes be confused. *Asterophyllites grandis* occurs in upper Namurian to upper Bolsovian strata in western Europe and the Maritime Provinces (Álvarez-Vázquez and Wagner 2017). Numerous specimens of *Asterophyllites grandis* have been collected by the authors in the Wamsutta study area, and apparently represent the foliage of a shrub- or small tree-like sphenophyte inhabiting shallow stream embankments and/or their proximal floodplains.

*Asterophyllites lindleyanus* Göppert 1848
(Fig. 6c)

**Remarks.** Up to 16 linear-lanceolate, whorled leaves 10–14 mm long per verticil are characteristic of this taxon. *Asterophyllites lindleyanus* closely resembles *Asterophyllites equisetiformis*, and the two species can be confused, but the latter’s leaves are often more delicate and thread-like (Álvarez-Vázquez and Wagner 2017). The leaves of *Asterophyllites grandis* are shorter than those of *Asterophyllites lindleyanus*, with a smaller length to width ratio, whereas the leaves of *Asterophyllites longifolius* are twice as long as those of *Asterophyllites lindleyanus*.

*Asterophyllites lindleyanus* is common throughout the Wamsutta Formation in the study area, as well as in the Cumberland Group in the Maritime Provinces (Álvarez-Vázquez and Wagner 2017). *Asterophyllites lindleyanus* seems to have flourished in or near narrow channel margins and levees, as many (parautochthonous) specimens have been recovered by the authors in sandy streambed horizons. This is the first report of *Asterophyllites lindleyanus* in the Narragansett Basin.

*Asterophyllites longifolius* (Sternberg 1825)
Brongniart 1828a
(Fig. 6d)

**Remarks.** This long-ranging late Namurian to Stephanian species is readily recognized by its 25–40 mm long leaves, which may overlap several verticils. Each verticil may comprise up to 20 leaves. Usually, leaves of *Asterophyllites longifolius* are thin and filiform (Álvarez-Vázquez and Wagner 2017), but may be occasionally sickle-shaped.

Isolated leaves of *Asterophyllites longifolius* are fairly common in the Wamsutta Formation in the study area, but are difficult to notice among the stems, tendrils, and vegetative litter on fossiliferous substrates. Elongate, needle-like leaves of *Asterophyllites longifolius* are sometimes imprinted into limb casts of Calamites suckowii that are rarely encountered lying flat on stream channel beds and flanks.

*Calamostachys Schimper* 1869

**Genus Calamostachys sp.**
(Fig. 6e)

**Remarks.** *Calamostachys* is the cone-shaped fructification of calamiteans (Josten 1991), which range widely throughout Pennsylvanian time. In contrast to *Lepidostrobus* sp., horizontally arranged, short and stiff bracts may present a distinctive saw-toothed appearance across the cone of *Calamostachys* (Bell 1944).

*Calamostachys* is fairly common in the study area, but is
Order Bowmanitales Meyen 1978
Family Bowmanitaceae Meyen 1978

Genus Sphenophyllum Brongniart 1822

*Sphenophyllum cuneifolium* (Sternberg 1821)
Zeiller 1878–1879
(Fig. 7a)

Remarks. This form features cuneate leaves 5–15 mm long.

generally too poorly preserved to observe details that would determine assignment to a species. Like the fragments of the parent calamitean plants, *Calamostachys* cones are encountered in the Wamsutta Formation as parautochthonous litter on fine-grained bedding planes of strata that represent abandoned channels and adjacent floodplains.

Figure 6. (a) HUPC No. 64,898, *Asterophyllites charaeformis*. Very small cupped leaflets are typical of this species. Scale equals 3 cm. (b) HUPC No. 64,859, *Asterophyllites grandis*. Four specimens on one slab. Scale equals 3 cm. (c) HUPC No. 64,897, *Asterophyllites lindleyanus*. 10–14 mm leaflets are characteristic. Scale equals 3 cm. (d) HUPC No. 64,849, *Asterophyllites longifolius*. Elongate leaves 25–40 mm in length distinguish species. Scale equals 3 cm. (e) HUPC No. 64,899A, *Calamostachys* sp. Note short, rigid bracts arranged horizontally. Scale in millimetres.
six in a verticil, with slightly arching, dentate to deeply laciniate distal margins and straight to slightly convex lateral margins (Abbott 1958). Sphenophyllum cuneifolium has pronounced polymorphy (Darrah 1969), and may have been lianescent. Farther down the axis, leaves are progressively more deeply cleft both centrally to laterally, forming finger-like lobes (Bell 1944; Abbott 1958). A single leaf enters each leaf base and dichotomizes 2–4 times asymmetrically, with only one vein entering a distal tooth or lobe (Abbott 1958; Darrah 1969). Sphenophyllum emarginatum is larger, but the distal margins of its non-dissected leaves are obtusely triangular and appear rounded or crenulated to the naked eye. Sphenophyllum majus is considerably larger than Sphenophyllum cuneifolium, and Sphenophyllum oblongifolium has whorls of four equidimensional leaves with a pair of distinctly shorter leaves.

Sphenophyllum cuneifolium is known to occur from Langsettian to Bolsovian time (Cleal 2005). The single specimen of Sphenophyllum cuneifolium found in the new Wamsutta collections demonstrates that this genus was a rare member of the study area macroflora.

Order Cordaitales Campbell 1902
Family Cordaitaceae Grand'Eury 1877

Genus Cordaites Unger 1850

Cordaites cf. principalis (Germar 1848) Geinitz 1855
(Fig. 7b)

Remarks. This long-ranging species is distinguished by long strap-like leaves clustered near the top. The leaves have rounded and obtuse apices with fine veins and one to five “secondary veins”. The trees are tall (up to about 30 m, with trunks 1 m in diameter), and have primitive gymnosperm foliage (Janssen 1965; Darrah 1969). Complete, coriaceous, sword-like leaves may be 50 cm or more in length (Darrah 1969). Cordaites principalis may be confused with stems of Calamites spp., but horizontal nodes are lacking in the former.

Cordaites principalis is ubiquitous throughout the Upper Pennsylvania strata of western Europe (Brousmiche Delcambre et al. 1998) and Atlantic Canada (Bashforth et al. 2014). Similar specimens are abundantly preserved as abandoned streamed detritus throughout the Pennsylvanian strata of the Narragansett Basin, although poor preservation prevents unequivocal identification at the species level. Nonetheless, these fossils indicate that arborescent cordaitaleans were a prominent component of the basin’s tropical rainforests. In the Wamsutta Formation red beds, including those within the study area, leaves of Cordaites cf. principalis often overlie one another in a crisscrossed “jackstraw” fashion on interfluve channel substrates and in the “washes” of associated floodplains. Undulating veins, wrinkles and possible invertebrate traces are commonly observed on leaf surfaces.

Genus Artisia Presl in Sternberg 1838

Artisia sp.
(Fig. 7c)

Remarks. This taxon comprises the distinctly ribbed pith casts of cordaitalean stems (Josten 1991). Faster growing stems caused the pith to rupture, exposing transverse pith diaphragms at the points of septation (Janssen 1965; Darrah 1969).

Artisia is widely distributed in western Europe and North America. Despite the abundant occurrence of Cordaites cf. principalis leaves throughout the Wamsutta Formation red beds, Artisia is rarely found in the formation. Until now, Artisia has never been recorded in the Narragansett Basin.

Order Lepidodendrales Prantl 1883
Family Sigillariaceae Unger in Endlicher 1842

Genus Sigillaria Brongniart 1822

Sigillaria sp.
(Fig. 7d)

Remarks. The genus Sigillaria is distinguished from Lepido-
dendron by its possession of vertically-aligned, usually hexagonal leaf scars, separated by straight or undulating furrows extending the full length of the trunk (Janssen 1965; Josten 1991). Sigillaria was an arborescent lycopod related to modern clubmosses and quillworts. Up to 30 m in height, it had a straight, gently tapering, singly or sparsely branched trunk surmounted by a cluster of grass-like leaves. Petrified casts of Sigillaria trunks range up to almost 2 m in diameter (Janssen 1965).

Sigillaria dominated many Late Pennsylvanian tropical rainforest wetlands in western Europe and North America. Only one specimen of Sigillaria, however, was collected in the Wamsutta Formation study area (Fig. 7d). Its presence in a matrix of planar fluvial-channel sandstone suggests that the plant was tolerant of disturbances due to flooding. Despite the rarity of Sigillaria specimens, numerous leaves (Cy-
perites) of arborescent lycopsids, probably associated with Sigillaria, were found by the authors (see below), frequently intermingled with various stems and other plant litter on shallow fluvial or wet and soggy floodplain substrates.

Genus Cyperites Lindley and Hutton 1832

Cyperites sp.
(Fig. 7e)

Remarks. The genus Cyperites comprises the long thin leaves of arborescent lycopsids (Janssen 1965), including Sigillaria (Bashforth 2005). They may have a “grass blade-like” appearance, with a prominent central vein, striae on both sides, and subparallel lateral margins (Josten 1991).
Cyperites may be distinguished from Calamites twigs and stems by the absence of horizontal nodes, and from Cordaites by its narrower width and the presence of a single distinct medial vein.

Cyperites is long-ranging, from Namurian to Stephanian, and is ubiquitous throughout tropical rainforest wetland strata of that interval worldwide. In the Wamsutta Formation of the study area, Cyperites is encountered as isolated leaves, or intermingled in an overlapping “jackstraw” fashion with Cordaites leaves, Calamites stalks and other parautochthonous stems on fluvial-channel and floodplain substrates.

AGE OF THE WAMSUTTA FORMATION

Stratigraphic ranges for 15 of the lower Wamsutta Formation taxa are given in Figure 8. In common with Knox
(1944), who first attempted to determine the age of the Wamsutta Formation through its fossil flora, we can confirm the presence of *Calamocladus* (=*Asterophyllites lindleyanus*), *Cyperites* (=lycopsid leaves) and *Sphenopteris valida* (=*Sphenopteris artemisiaefoliioides* as indicated by Bell 1944). Of the florule reported by Knox, we did not find *Annularia stellata* or *Neuropteris rarinervis* Bunbury (= *Laveineopteris rarinervis*). With regard to the latter, Knox (1944) likely observed not *Laveineopteris rarinervis* but *Neuropteris obliqua*, one of the most common and morphologically variable taxa in the Wamsutta Formation flora. Consequently, Knox erroneously concluded that his florule was of Asturian age, but Darrah (1969) determined it to be Bolsovian.

The macroflora reported in this paper (Table 1; Figs 3–7) correlates most closely with that of the Lower Pennsylvanian Cumberland Group of the Maritime Provinces. The Cumberland Group was assigned a late Langsettian to early Duckmantian age by Bell (1944); later Dolby (1997) assigned the Cumberland Group a late Yeodanian to Langsettian age based on palynological assemblages (Fig. 8). Bashforth et al. (2014) considered the Cumberland Group redbeds in New Brunswick to be mainly late Langsettian in age. Correlation to the Cumberland Group is indicated by the following plant-fossil species in the Wamsutta Formation: *Neurallethopteris schlehanii*, *Neuropteris obliqua*, *Senftenbergia plumosa*, *Calamites suckowii*, *Annularia asteris*, *Annularia cf. microphylla*, *Asterophyllites charaeformis*, *Asterophyllites grandis*, *Asterophyllites lindleyanus*, *Sphenopteris cuneifolium*, and *Cordaites principalis*.

Figure 8. Chart depicting biostratigraphical ranges of 15 fossil-plant species collected in the Wamsutta Formation, inferred from known ranges in western Europe and the Maritime Provinces of Canada. Ranges following Brousmiche Delcambre et al. (1998), Cleal (2005), and Álvarez-Vázquez and Wagner (2017). Shaded-gold column shows the resultant age determination of the lower Wamsutta fossil macroflora, i.e., late Langsettian to early Duckmantian.

Two aspects of the Wamsutta Formation macroflora are notable. First, the domination of cordaitaleans and sphenophytes (Table 1). The abundance of sphenophytes enables a detailed comparison with the assemblages recorded by Álvarez-Vázquez and Wagner’s (2017) in the Maritime Provinces. We record many of the species that they listed, including three species of *Annularia* and four of *Asterophyllites*. Second, the next most common taxa in the Wamsutta Formation are the medullosan seed ferns (see Table 1), represented most abundantly by *Neuropteris obliqua*. This taxon is abundant in the Langsettian and Duckmantian strata of western Europe (Laveine 1967; Josten 1991). Bell (1944) also reports the presence of *Neuropteris obliqua* in the Cumberland Group.

The Wamsutta Formation macroflora reported here also resembles late Langsettian/early Duckmantian florules of the Iberian Peninsula, where there is a complete succession of Carboniferous macrofloral zones (Wagner and Álvarez-Vázquez 2010). At least thirteen of the twenty-two Wamsutta taxa that we report (Table 1) are listed by the latter authors in their Langsettian and Duckmantian macrofloral tables (their table 1, p. 257 and table 2, p. 266), including *Neuropteris obliqua*.

Finally, the presence in the Wamsutta macroflora of *Laveineopteris cf. loshii* and *Neurallethopteris schlehanii* further suggests a late Langsettian to early Duckmantian age for the lower Wamsutta Formation red beds. Regarding
Laveineopteris loshii, Cleal and Thomas (1994) identified a Laveineopteris loshii Subzone, denoting the first appearance in Europe of Laveineopteris loshii in the middle Langsettian (and, by inference, its associated shade leaf Cyclopteris sp.; see Cleal and Shute 2003). Thus, the initial appearance of Laveineopteris cf. loshii suggests a biostratigraphic lower limit for the age of the lower Wamsutta Formation macro flora—i.e., middle Langsettian. On the other hand, Neuralethopteris schlehanii, a key marker species with a well-known range in western Europe (including the Iberian Peninsula) and North America (Goubet et al. 2000; Wagner and Álvarez-Vázquez 2010) suggests an upper limit for the age of the macroflora of the Wamsutta Formation in the study area—i.e., late Langsettian to early Duckmantian. This results from a general consensus that Neuralethopteris schlehanii became extinct abruptly at the close of the Langsettian substage (Goubet et al. 2000; Wagner and Álvarez-Vázquez 2010). However, there are a few reports of its occurrence in the early Duckmantian (e.g., Bell 1944).

CONCLUSIONS

The Early Pennsylvanian age of the macroflora in the lower Wamsutta Formation of the Narragansett Basin is more precisely defined on the basis of new plant-fossil collections from Red Rock Hill in North Attleboro, Massachusetts. Biostratigraphic comparisons of the new fossil flora with Middle Pennsylvanian tropical rainforest macrofloras in western Europe, especially those of the Iberian Peninsula where there is a complete succession of Late Carboniferous macrofloral zones, and with correlative tropical rainforest macrofloras of the Cumberland Group of the Maritime Provinces, reveal a late Langsettian to early Duckmantian age for the study area’s Wamsutta fossil plants. Since the horizons in which the fossil plants were found are stratigraphically 100 m or more above the base of the Wamsutta Formation exposed in the area, the lowermost horizons were likely deposited wholly within Langsettian time.

Comprising a series of well-drained alluvial fans, the sedimentary rocks of the Wamsutta Formation red beds contain sparse plant fossils, which are usually fragmentary and poorly preserved. Nonetheless, a total of 22 plant macrofossil taxa are identified herein. Fossil-plant species reported for the first time in the Narragansett Basin, and occurring exclusively within the lower Wamsutta Formation, are the following: Neuralethopteris schlehanii, Annularia asteris, Annularia cf. microphylla, Annularia radiata, Asterophyllites grandis, Asterophyllites lindleyanus, and Artisia sp. The lower Wamsutta represents a seasonally dry (moist subhumid) upland fluvial environment with clastic sediment input that experienced repeated flooding. It supported a tropical rainforest, but was devoid of persistent swamps and peat deposits. The new collection of Wamsutta Formation macroflora, supplemented by a smaller collection assembled by Knox (1944), represent the oldest known macroflora within the Narragansett Basin.

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