# A PHYTOGEOGRAPHICAL ANALYSIS OF TAXUS（TAXACEAE） BASED ON LEAF ANATOMICAL CHARACTERS 

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A phytogeographic analysis of 845 Taxus specimens is presented based on leaf anatomical characters for the number of stomata rows in a stomata band and the number of epidermal cells that lack papillae between the leaf margin and stomata band．The specimens are arranged by continent，country，state or province，and species，and represented geographically on three maps：（1）North America，（2） the Euro－Mediterranean，and（3）Asia．Taxus is least diverse and most distinct taxonomically in North America，and most diverse and least distinct taxonomically in southwest China．Stomata data show several clines in North America，an obvious south to north decrease for the Mesoamerican yew（T．globosa）and Pacific NW yew（T．brevifolia）populations combined，and a less obvious reverse cline for the Canada yew（T．canadensis）．The results are discussed in review of other paleobotanical data．It is suggested that Taxus immigrated to North America from Asia across a Pacific land connection during the Cretaceous，and from Europe to North America across North Atlantic land bridges during the Tertiary．The low diversity of Taxus in North America is suggested to be the result of the K／T extinction event． In the Euro－Mediterranean，evolution of Taxus is suggested to have been impacted more by extinction as a result of climatic changes during the Tertiary，and by hybridization during the Pleistocene．The greater diversity in SW China is indicated to be the result of less extinction there and more frequent hybridization during the Pleistocene，not only among authochtonous species，but also allochtonous species as a result of the Himalayan uplift．

## CHINESE ABSTRACT

基于每一条气孔带内气孔列数目以及叶边缘和气孔带之间缺乏乳突的表皮细胞数目的解剖特征，本文对 845 份红豆杉属 Taxus标本进行了植物地理分析。标本按洲，国家，州或省以及种来排列，其地理分布显示在 3 幅地图上：（1）北美，（2）欧洲一地中海，（3）亚洲。红豆杉属在北美的多样性最低且分类上区别最明显，而在中国西南部的多样性最高且分类上区别最不明显。气孔数据显示红豆杉属在北美有几个渐变群，中美洲红豆杉 T．globosa 和太平洋西北红豆杉 T．brevifolia 的居群由南向北明显减少，加拿大红豆杉 T．canadensis 由北向南减少得不太明显。综合考虑了古植物学资料，这些结果支持以下一些假说。红豆杉属在白垩纪和第三纪分别从亚洲和欧洲穿过太平洋陆地连接和北大西洋陆桥迁移到北美。白垩纪一第三纪界线绝灭事件造成了红豆杉属在北美低的多样性。第三纪气候变化造成的绝灭和更新世发生的杂交对欧洲一地中海地区红豆杉属的演化影响更大。由于喜马拉雅山的抬升，红豆杉属的土著种和外来种在更新世的杂交比较频繁，而且在中国西南部绝灭较少，所以该属在中国西南部的多样性较高。

## INTRODUCTION

The genus Taxus has included eight geographically defined species：（1）T．baccata L．—Europe，N Africa and SW Asia（Franco 1964），（2）T．cuspidata Siebold \＆Zucc．－temperate E Asia（Krüssmann 1985；Ohwi 1965）， （3）T．wallichiana Zucc．－Himalayas（Krüssmann 1985），（4）T．sumatrana（Miq．）de Laub．—S China，Philip－ pines，Taiwan，Sulawesi，and Sumatera（de Laubenfels 1988），（5）T．globosa Schltdl．－N Central America to Mexico（Ferguson 1978），（6）T．brevifolia Nutt．－NW North America（Ferguson 1978；Hils 1993），（7）T． floridana Nutt．ex Chapm．－W Florida（Ferguson 1978；Hils 1993；Price 1990），and（8）T．canadensis Mar－ shall—NE North America（Ferguson 1978；Hils 1993；Price 1990）．Except for T．sumatrana，these were also recognized by Pilger $(1903,1916,1926)$ as subspecies of T．baccata．

These eight geographically defined taxa－generally accepted as species（Bailey 1933；Farjon 1998； Hils 1993；Krüssmann 1985；Rehder 1940；Silba 1984）—have not been clearly distinguished（Ferguson 1978；Price 1990），including several other species and varieties that have been recently recognized（Farjon 1998，2001；Fu et al．1999）；however，I will show that this traditional geographical classification is distorted to recognizing more species where diversity in the genus is least（North America）and less species where
diversity is greatest (SW China). This will be based on quantitative leaf character attributes for the number of stomata rows (SR) in a stomata band, and the number of marginal cells (MC) across an abaxial marginal zone without papillae. These data will be summarized on geographical maps of North America, EuroMediterranean, and Asia, and analyzed in the results section of this study. This is followed by a discussion of phytogeographical relationships. Data for all herbarium specimens studied are provided in an appendix according to continental and political regions and taxonomy.

## MATERIALS, STANDARDS AND METHODS

MATERIALS.-The materials of Taxus include 845 specimens of fresh and dried branchlets with leaves from throughout the natural range of the genus (Appendix), and an undetermined number of specimens from cultivated plants in Australia (1), England ( $\sim 50$ ), France ( $\sim 20$ ), and the United States ( $\sim 300$ ), and from miscellaneous other sources, the main one was Phyton, Inc ( $\sim 65$ ), now Phyton Biotech, a commercial company specializing in producing taxol from tissue culture of Taxus.
STANDARDS.-The genus Taxus Linnaeus (Taxaceae Gray) is defined by cone and leaf morphology (Florin 1931, 1948c, 1951) in relationship to other "taxad" genera, characterized by producing an arillocarpium (Spjut 1994)—a type of cone in which the seed is subtended by a fleshy arillate bract (Airy Shaw 1973; Cheng \& Fu 1978; Florin 1948a; de Laubenfels 1988).

Taxads include both extant and extinct taxa; the extant genera, in addition to Taxus, are Amentotaxus (5-6 spp., China, Vietnam), Torreya (6 spp., E Asia, N America), Austrotaxus (1 sp., New Caledonia), Cephalotaxus (8-11 spp., E Asia), and Pseudotaxus (1-2 sp., China) (Fu et al. 1999). Molecular studies employing ribosomal RNA (Chaw et al. 1993, 1995), chloroplast DNA (Tsumura et al. 1995), or RAPD (T. Wang et al. 2000), suggested Amentotaxus and Torreya to be more closely related to each other than to Cephalotaxus or Taxus, and that Cephalotaxus is basal to two clades, (1) Torreya/Amentotaxus and (2) Taxus/Pseudotaxus/Austrotaxus (Cheng et al. 2000); however, whether these clades should be treated in separate orders, families, subfamilies, or tribes, is controversial (Hill 1998).

Cones.-Pseudotaxus and Taxus produce a terminal seed on a lateral (secondary) short shoot (André 1956; Dupler 1920; Miller 1988) that is only partly surrounded by a loose cupular bract, whereas in other genera the seed is more fully and tightly covered by the aril (Florin 1948b; Sahni 1920). Cephalotaxus is distinct for its biovulate cone scales from which usually only one ovule matures (Singh 1961). Amentotaxus differs for its terminal, "racemose" male shoots (Cope 1998; Fu et al. 1999). The Austrotaxus cone was regarded as isolated from other taxads based on anatomy of the seed coat (Bobrov et al. 2004).

The closely related Pseudotaxus (1-2 spp., China, Fu et al. 1999) differs from Taxus by a white arillocarpium (Cheng 1934), and additional sterile scales in male cones (Florin 1948c).

Leaves.-Taxus leaves are differentiated from those in other taxad genera by papillose cells that define the "stomatic apparatus" (Dilcher 1969; Florin 1931, 1948c, 1951, 1958). This apparatus includes 4-8 small subsidiary papillose cells that encircle each stoma (Florin ring) and adjacent (accessory) papillose cells (Figs. 1A, 1B, 1C). Stomata develop in longitudinal rows (periclinal) in a stomatal region divided into two bands by a midrib (e.g., Fig. 1A). The midrib and marginal cells vary in size, shape, and development of papillae. Further details—with photomicrographs—can be found in Ferguson (1978), Florin (1931, 1951), Jinxing and Yuxi (2000), Kvaček (1984), and Kwei and Hu (1974).

Mammillae, not to be confused with papillae (Bertrand 1874), develop singly over most of the cell's surface as large lens like bumps. Under a dissecting scope (30x) they appear most conspicuous along leaf margins, less so on the epidermal surfaces (adaxial or abaxial). Papillae, by contrast, are smaller and numerous on a cell-like pimples. They develop in 1-3 distinct or concrescent rows, generally discernible only under a microscope-at least $100 \times$, and only on the abaxial surface. Papillae are always present in stomata bands, gradually diminishing in prominence outside the bands towards the leaf margins, and may develop entirely or partially or not at all on midrib cells.

The development of stomata in rows and their differentiation by papillose accessory cells, which together make up the stomata band, are the most distinguishing features of Taxus relevant to data in this study, compared
to other extant genera of taxads (Florin 1931, 1951). For example, Pseudotaxus has glaucous stomata bands (Cheng 1934; Florin 1931, 1948b, 1948c, 1948d) devoid of papillae, except on subsidiary cells, and more stomata rows per band-23-28 rows (Florin 1948c), in contrast to 4-21 rows in Taxus (Appendix). Amentotaxus and Torreya (Amentotaxaceae) have papillose glaucous bands largely of subsidiary cells (periclinally arranged), rather than accessory cell types (Florin 1951, 1958). Austrotaxus (Austrotaxaceae; Nakai 1938; Florin 1958), which also differs conspicuously by its long-linear leaves-comparable to some Podocarpus spp.-has stomata evenly scattered across the entire abaxial surface without clear differentiation of rows and bands in which the epidermal cells are similar to those on the adaxial surface-irregularly quadrate (or pentagonal) as in Taxus. These differences, and the presence of other features such as sclereids and resin canals in leaves of Torreya (Bertrand 1874) and Cephalotaxus, would seem to support classification of the taxads in different families (Amentotaxaceae, Austrotaxaceae, Cephalotaxaceae, Taxaceae).

The features of the stomata band that distinguish Taxus from other extant taxads do not apply to extinct taxads, however (Florin 1951, 1958; Harris 1976a, 1976b; Kvaček 1984; Miller 1977). Photomicrographs of many taxad fossils from Jurassic deposits presented by Florin (1958) show remarkable detail that are strikingly similar to extant Taxus in leaf epidermis (Kvaček 1984; Meyen 1984), except for narrower stomata bands with fewer stomata rows (Kvaček 1984). Indeed, some leaves, which included twigs and arillocarpia, were assigned to Taxus; these are T. bornholmiensis Florin with 4-5 stomata rows, T. harrisii Florin with 5 stomata rows, and T. jurassica Florin with 3-5 stomata rows; however, none of these appear to belong to the genus Taxus. Harris (1976a, 1976b), for example, transferred T. jurassica to Marskea, an extinct genus characterized by op-posite-decussate leaves (Florin 1958; Harris 1976b), which are clearly evident in Florin's (1958) photograph of Marskea jurassica. Jurassic taxads include many other extinct genera (Florin 1958) that may have existed since the Triassic (Florin 1951; Meyen 1984).

Epidermal cells adjacent to leaf stomata bands, the midrib and marginal areas, are usually papillose in part. Epidermal cells on the abaxial surface nearest the leaf margin appear to have evolved in some species by extension (folding) of the upper (adaxial) surface to the lower (abaxial) surface; leaves of many specimens are revolute along their margins in which the abaxial epidermal cells are often more similar to those above than to the adjacent cells below (Nicolosi 1982).

The comparative morphological relationships of Taxus to other extant taxads (Florin 1931, 1948c; Appendix) indicate that the ancestral Taxus leaf had a partially differentiated abaxial epidermis in which stomata developed in definite rows but not in distinct bands. Evidence for this can be seen in $T$. wallichiana and allied species in the E Himalayas and SW China. Their leaf stomata occur not only in stomata bands, but also on the abaxial midrib; essentially, stomata develop across the entire abaxial leaf surface to within several cells of the margin-in up to 21 rows. The abaxial marginal and accessory epidermal cells are all nearly rectangular and papillose-in sharp contrast to the epidermal cells on the adaxial surface that are much shorter, $\pm$ trapezoidalpentagonal, and without papillae.

## METHODS

More than 1,000 herbarium specimens (A, BH, BM, BOLO, E, GH, K, M, NA, NY, P, PE, PH, S, U, US; Holmgren et al. 1990) were studied of Taxus throughout the natural range of the genus to assess morphological variation in characters that involve branches, bud-scales, leaves, and cones; 845 are cited in the appendix, and additional specimens are mentioned in this paper. Each specimen was photographed with a Nikon camera using 35 mm color film with 35 mm and 60 mm lenses. Stafleu and Cowan (1976-1988) were consulted for location of types, other specimens of historical relevance to this study, and references.

From each herbarium specimen of Taxus, one mature leaf was selected for microscopic study of anatomical features. The Taxus leaf was soaked in water for $8-16 \mathrm{hrs}$. The leaf was then transversely sectioned in the mid region as bryologists routinely section leaves of mosses for taxonomic identifications. With a single-edge razor blade and dissecting needle as a guide, 5-10 transverse sections were generally made. Then an abaxial epidermal layer was removed from both remaining leaf portions, generally $0.5-2.0 \mathrm{~mm}$ in length, by scraping mesophyll parenchyma from the epidermal layer with a razor blade. Occasionally, the entire abaxial leaf


FIG. 1A. Mid leaf sections of Taxus caespitosa var. Iatifolia, from a cultivated plant in Maryland, U.S.A, Spjut 10485 (wba), representative of the Baccata Group, Cuspidata Alliance. Top: transverse section (T-sect.), $\sim 100 \times$, showing elliptical shaped epidermal cells, mesophyll layers of anticlinal palisade layer of parenchyma and spherical parenchyma cells, drawn by Karen Parker. Bottom: abaxial epidermal layer from margin (left) to midrib (right), $\sim 250 \times$, showing a marginal border of 8 smooth (non-papillose) cells wide followed by a stomata band with 13 rows of stomata, and a midrib of mostly smooth cells, drawn by R. Spjut.


Fig. 1B. Mid leaf sections of Taxus brevifolia var. reptaneta from Siskiyou Co., California, U.S.A., representative of the Wallichiana Group, R. Spjut \& T. Spjut 11835 (wba, type). Top: T -sect. $\sim 100 \times$ shows tall angular epidermal cells, drawn by Karen Parker. Bottom: abaxial epidermal layer from margin (left) to midrib (right), $\sim 250 \times$, shows marginal region of 10 smooth cells across of which 6 rows are inflated, followed by 8 rows of papillose cells, 5 stomata rows, and a papillose midrib, drawn by R. Spjut.


FIG. 1C. Mid leaf sections representative of the Taxus Sumatrana Group, drawn by R. Spjut. From top to bottom: Top: T. mairei var. maire from Guangdong (China), Tsang 20694 (US), showing elevated and truncated midrib along abaxial surface with enlarged epidermal cells in comparison to smaller elliptical epidermal cells on adaxial surface, the lower T-section, T. mairei var. mairei from Yunnan, Maire s.n., isotype, showing truncated and channeled midrib with larger spherical parenchyma cells lying against smaller epidermal cells-most conspicuous along midrib and marginal zones. Lower two sections: abaxial epidermis from margin to across the midrib; upper most from Guizhou, isotype (A) of $T$. speciosa, showing marginal region of 23 smooth cells in width, a stomata band with 16 rows of stomata, and a smooth midrib; the lowest section from holotype of T. sumatrana, showing long rectangular cells and 12-14 stomata rows.
surface and a medial portion of the adaxial epidermis were removed. All sections were examined under magnifications of $100 \times, 250 \times$, and $400 \times$ (Nikon binocular microscope) for cell shape, number of stomata rows, number of cells marginal to stomata bands, and for papillae position and distribution across the abaxial leaf surface. The results were sketched and described on small packets $3 \times 5$ inches. A temporary slide of the sections and photographs of the herbarium specimen were retained for each packet. Leaves from fresh specimens were also similarly studied throughout the range.

Figs. 1A-1C show diagrammatic leaf sections of the mid region that is representative of three species


FIG. 2A. Number of stomata rows per band (SR) in leaves of Taxus plotted from locality data on representative herbarium specimens from North America; see Appendix for specimen data. Specimens from northern Mexico indicated in yellow numbers are not easily distinguished from those in Florida; therefore, these are considered T. globosa var. floridana.


FIg. 2B. Number of stomata rows per band (SR) in leaves of Taxus plotted from locality data on representative specimens from the Euro-Mediterranean; see Appendix for specimen data. Most specimens were found to have 8-10 stomata rows per band as indicated in yellow numbers. Occasional plants with higher counts, 11-15 stomata rows per band, are shown in red. Both yellow and red numbers belong to the Baccata Alliance. Numbers in white belong to $T$. canadensis. Most T. canadensis have less than 8 stomata rows per band, but a few with higher counts—up to 10 stomata rows per band—are shown.
groups of Taxus. These include (1) a complete transverse section and (2) an epidermal portion of the abaxial surface from one margin to across the midrib. As previously indicated, similar sketches were made on 3.5 $\times 5$ inch $(8 \times 12.5 \mathrm{~cm})$ packets for most herbarium specimens studied except only portions of the stomata and marginal areas were drawn, while number of stomata in a band and number of marginal cells adjacent


FIG. 2C. Number of stomata rows per band (SR) in leaves of Taxus for representative specimens in EAsia of the Baccata and Wallichiana Groups; see Appendix for specimen data. The Sumatrana Group, which is common in SE China, is not shown to contrast the difference between the Wallichiana and Baccata Groups. The Wallichiana Subgroup includes T. suffnessii in white, T. florinii in blue, and T. wallichiana var. yunnanensis in black. The Chinensis Subgroup within the Wallichiana Group includes T. chinensis shown in purple in mainland China and related species in the Chinensis Subgroup in the Philippines, Sumatera, Sulawesi, and Taiwan, and also two species in mainland China, in yellow. The Cuspidata Alliance includes four species; T. cuspidata, T. biternata, and T. caespitosa, in maroon, and T. umbraculifera, in black. Note higher counts for T. contorta (8-11) and lower counts for T. wallichiana (11-14) where the ranges of these taxa overlap in the central Himalayas. The widest range in number of stomata rows was found in N Myanmar to SW China where T. florinii is recognized to have $7-12$ rows per band, and T. suffnessii from 13-20 rows per band.
to stomata bands were recorded. The leaf margin can be difficult to pinpoint when leaves are rounded along margins. This is determined by the smallest cell that is usually mammillose. Both types of leaf sections were examined to determine where papillae develop on cells between the margin and stomata band.

Variation due to mechanical preparation, error in counting, and environmental factors (Deryugina \& Nesterovich 1981), were only generally assessed—for practical reasons—from duplicates that were unintentionally included in this study, occasional field collections that were collected at various heights from one or several trees of a population (top, middle, and lower branches of T. brevifolia from trees in California and Oregon), and from test cases of selected leaves at various developmental stages from shrubs in cultivation. Practical reasons include damage to herbarium specimens caused by removing a leaf, and the time required to prepare leaf sections and record data, approximately one hour for each specimen.

Herbarium specimens studied are listed in an appendix with data on numbers of stomata rows per band (SR) and numbers of marginal cells (MC) without papillae. Specimen data are arranged by continent, then by country within continents, and finally by taxa, generally from south to north in North America, and from west to east in Eurasia, and then east and south from the Himalayas to Indonesia. Leaf anatomical data are further arranged by decreasing order in number of stomata rows (SR), and by increasing order in number of marginal cells (MC) except for T. canadensis, the Sumatrana Group, and for duplicate specimens belonging to the same species, or duplicate specimens from the same locality or collection number. Only minimal collection data are cited, although for many specimens data were minimal. If the stomata count varied on each side of the midrib of a single leaf, this is indicated by a slash; for example, T. canadensis frequently had 5 stomata rows in one band and 6 in the other (5/6). A dash between numbers indicates a variable range, especially when more than one leaf from the same specimen was studied, or a dash alone indicates absence of data. It should
be remembered that these data are a byproduct of an overall taxonomic study of the genus Taxus (see Spjut 2007); i.e., they were not compiled with this paper in mind. Additional character features that appeared to correlate with the findings are also noted (e.g., length of epidermal cells/width of epidermal cells or $1 / \mathrm{w}$ ).

RESULTS
Numbers of Stomata Rows (SR).-Figures 2A, 2B and 2C show numbers of stomata rows per band on maps of North America, Europe, and Asia, respectively, for representative specimens listed in the Appendix. Each number represents a count from a single leaf of a herbarium or fresh specimen; the datum is plotted at the general location where it was reported to have been collected. Where the count varied on each side of the leaf midrib, or among duplicate specimens, the highest number was scored.
Stomata data for duplicate specimens, or among specimens from different plants at the same locality, are summarized under five cases as follows:
(1) For 62 duplicates of herbarium specimens included in this study, the same number of stomata rows per band was found in 19 duplicate specimens. Among the remaining 43 duplicates, 17 differed only by 1 row, another 17 differed by 2 rows, 6 varied by 3 rows, and 3 varied by as much as 4 rows. Duplicates that varied by 4 stomata rows appear to have been collected from different habitats within the same general locality. Examples are Farges 128 from Sichuan, Tsiang Ying 1425 from Guangdong, and several cases where the same collection number was reported from multiple locations, Wilson 1265, from two sites in Sichuan and one in Hubei, and Wilson 4053 from two locations in Sichuan (Rehder \& Wilson in Sargent 1914).
(2) From a single shrub of T. caespitosa Nakai var. latifolia (Pilg.) Spjut in Maryland, $\sim 35$ yrs of age and 3 m in height, 16 leaves were selected from branchlets with various exposures to light and age ( $1^{\text {st- }} 2^{\text {nd }} y r$ ). With one exception, 15 were found with 13 stomata rows per band; one leaf—plucked from a well-shaded branchlet near the main trunk—had 16 rows.
(3) From trees of T. brevifolia, leaves from top, middle, and lower branches had the same count at two sites-one in California, and another in Oregon-but varied by l-2 rows in leaves from different trees at the same sites.
(4) In Taiwan, individuals of a population collected on five occasions (Appendix) had the same number of stomata rows at one site (Tongshi 7), but varied from 11-14 rows per band at two other sites (Tongshi 5, Tongshi 6).
(5) The age of the leaf was apparently not a factor in 20 leaves studied of one cultivated individual of T. mairei (Lemée \& Lév.) S.Y. Hu ex T.S. Liu var. speciosa (Florin) Spjut; 16-18 stomata rows per band were found in all leaves from buds to $3^{\text {rd }}$ yr branchlets.

Generally, the number of stomata rows per band varied by a count of 3 for about half of the individuals or species. In NW North America (T. brevifolia), this occurred frequently among different individuals within a population, whereas in Europe (Baccata Alliance) leaves of Taxus exhibited the same range in variation whether obtained from the same plant or from different plants at the same site. A wider range, $13-18$ stomata rows, in the E Himalayas (T. wallichiana), and a narrower range, 5-6 or 6-7 stomata rows, in NE North America (T. canadensis), were also evident.

Despite this range in variation, the numbers of stomata rows per band in North American Taxus (Fig. 2A) show a distinct cline from south to north for the Wallichiana Group (Central America to Florida, California). Specimens from Honduras to southern Mexico had $7-11$ rows per band (Bertrand 1874); those from N Mexico and Florida had (5-) $7(-8)$ rows per band in further contrast to $4-7(-9)$ stomata rows per band in the Pacific NW. This northward decline in stomata rows per band is also apparent within the Pacific NW by the highest count of 9 rows found in specimens from California, compared to the lowest count of 4 rows from specimens at more northern locations-in the Rocky Mountains. This cline was also noted to be associated with an increase in length of epidermal cells relative to width ( $1 / \mathrm{w}$ ), ranging from an average of ca. $3 \times 1 / \mathrm{w}$ to $8 \times 1 / \mathrm{w}$.

A reverse cline, however, is evident in the NE North America by 9 rows per band occasionally found in specimens from Newfoundland, compared to the more common 6 rows per band as reported by Bertrand (1874), which includes specimens from Quebec to Ontario, south to Kentucky (Spjut 1998a, 2000a, 2000b).

In the Euro-Mediterranean, Taxus leaves most often had 8-10 stomata rows per band (Fig. 2B); however, notably higher counts and lower counts were found at widely scattered locations (Fig. 2B, red and white numbers; appendix in bold type). Among the higher counts are specimens from the Caucasus Mountains, one with 15 rows (Princeps Kascelsky, ex. Herb. Hort. Imper. Petro), another with 13 rows (Woronowa s.n.). Occasional specimens with 12 rows are from Bosnia (Biol. Inst. Dubrovnik 37), Romania (Topa, Bot. Mus. Exsic), Finland (Florstöm s.n.), Portugal mainland (Yoller 61), Portugal Azores (Goncalves 4625), and Algeria (Swingle s.n.). Those that were noted to differ in other morphological features are indicated in bold type, including also
specimens with 11 stomata rows per band, such as one specimen from England (Bowden \& Hillman 433) that had globose shaped epidermal cells, instead of the usual elliptical shape; others such as Hauti 28894 from the British Isles and Busch s.n. from the Caucasus Mountains had a broader marginal zone of bare cells, as indicated later under results for marginal cells. A higher number of stomata rows with a wider leaf margin indicate a higher density of stomata as seen in the Cuspidata Alliance. The range of variation for the Baccata Alliance was greatest in Transcaucasia.

Leaves with fewer than 8 stomata rows per band include seven specimens with 7 rows-from Germany, Austria, Switzerland, Bosnia, and Turkey; these belong to the T. baccata Alliance. Those with fewer than 7 rows were relatively rare ( $1-2 \%$ of 196 specimens cited for the Euro-Mediterranean Region), two specimens with 5 rows from Norway and Sweden, and one with 4 rows from Slovenia. These are considered T. canadensis as shown later. A cline is not apparent in the Euro-Mediterranean as it is in North America; however, the relative frequent occurrence of stomata in $8-10$ rows per band appears significant when compared to a greater range of variation in E Asia (7-21 rows per band, Fig. 2C).

In E Asia, the number of stomata rows per band ranged from 7-16 in the temperate region, and from (5-) 7-19 (-21) in the tropical region with two patterns converging in the Himalayas, one from the west with 5-8 ( -11 ) rows of stomata, and another from the east with $7-21$ rows per band.

Number of Marginal Cells (MC).—The absence of papillae nearest the leaf margin, as measured by the number of marginal cells (MC) across between the margin and stomata band, is depicted geographically in Figs. 3A and 3B, and detailed in the Appendix.

Data on marginal cells lacking papillae along the abaxial surface (MC) were recorded less often for North American species because they were distinguishable early in the study (Hils 1993; Spjut 1992, 1993). It was recognized that marginal cells of T. canadensis always lacked papillae (Hils 1993, Spjut 1992, 1993, 1998a, 2000b), which has since been determined to vary from 11-19 cells across in North American plants (Appendix); the absence of papillae on the abaxial midrib is in sharp contrast to the papillose midribs of other North American species (T. brevifolia, T. globosa).

In the Euro-Mediterranean, the abaxial surface of leaves of most Taxus specimens ( $\sim 75 \%$ ) lacked-papillae along a relatively narrow marginal zone of $4-7$ cells across (Fig. 3A). This included the lectotype for $T$. baccata and two specimens from the Caucasus Mountains that, unlike the lectotype, were found to have a relatively high stomata count as noted earlier-one with 13 stomata rows per band, and one with 15 stomata rows per band-and also one specimen from Bosnia with 12 stomata rows per band. A specimen from the Caucasus Mountains-that had abaxial marginal papillae to within one cell from the margin—lacked papillae on nearly half of the cells across the midrib in the median region (Woronowa s.n.).

Leaves of Euro-Mediterranean specimens with a relatively broad zone of bare cells between the margin and stomata band-from $8-24$ cells across-were found less frequently ( $\sim 25 \%$ ). These are from widely scattered places. Many are indistinguishable from T. canadensis in North America—based on additional characters of branching, phyllotaxy, and color (Figs. 4-5); therefore, are referred to T. canadensis (Spjut 2000b). In Europe, leaves of T. canadensis may include a transitional zone of papillose cells between the stomata band and margin (Fig. 4, specimen from Morocco). These plants may be hybrids between T. canadensis and $T$. baccata.

The Cuspidata Alliance showed an intermediate range of values for abaxial marginal cells without papillae, (6-)8-18(-24) cells across (Figs. 3B, 6), compared to the Baccata Alliance, (1-)4-7(-11) cells across (Fig. 3A) and the Sumatrana Group, 8-36 cells across (Fig. 3B, 6). The higher stomata counts in relatively narrower stomata bands for the Cuspidata Alliance (see also Dempsey \& Hook 2000) and Sumatrana Group means in effect they have a higher leaf stomata density, recognizing also that stomata density is related to width of the epidermal cells and the width of the stomata band (Nicolosi 1982), and that a transitional zone of papillose cells is always present in the Baccata Alliance but not in the Cuspidata Alliance. Data on number of papillose cells across the abaxial margin were not included in this study because this was observed to be highly variable, although the absence of papillae in T. canadensis has taxonomic significance in North American species as already indicated (Hils 1993; Spjut 1992, 1993, 1998a, 2000b).


FIG. 3A. Number of epidermal marginal cells (MC) without papillae between the margin and stomata band (abaxial surface nearest margin) for representative specimens of Taxus from the Euro-Mediterranean Region; see Appendix for locality data. Numbers in red show the more common range in variation-a leaf margin 4-7 cells across-that corresponds to T. baccata and its allies in the Euro-Mediterranean Region. Numbers in white contrasts the higher counts-a leaf margin 6-24 cells wide-that belong to T. canadensis. Data in yellow numbers indicate intermediates that are morphologically similar to T. biternata, T. canadensis, T. cuspidata, or T. contorta, appearing more frequent in specimens from the Caucasus Mountains (Appendix, data in bold type)..


Fig. 3B. Number of epidermal marginal cells $(M C)$ without papillae between the leaf margin and stomata band for three species groups of Taxus obtained from herbarium specimens cited in the Appendix. Data for the Wallichiana Subgroup and T. contorta (Baccata Group) are summarized, and data for the Chinensis Subgroup-summarized in Fig.6-are excluded here to further contrast the wide range in variation seen in the T. cuspidata Alliance (Baccata Group) and the Sumatrana Group. The Sumatrana Group includes T. celebica in maroon, T. mairei in black, T. kingstonii in white, and T. sumatrana in yellow. The Cuspidata Alliance includes T. biternata in orange, T. cuspidata in maroon, T. caespitosa and T. umbraculifera in red. Note that the widest range in variation occurs in SW China.

FIg. 4. Examples of T. canadensis var. canadensis as characterized by the regularly isodichotomous branchlets, and acute to acuminate leaves that spread from branchlets more by their petioles than by their blades, arranged in nearly two ranks, and by the abaxial surface of leaves lacking papillae on at least 8 cells across from the margin. $M C=$ Marginal Cells across the marginal zone without papillae. $S R=$ Stomata Rows in one stomata band. The specimen from Morocco has a partially papillose margin, 8-9 of the 18 cells between the margin and stomata band lack papillae. Specimens above left, above right and left: Bean et al. 19634 (PH), Lowe 570 (BM), Font Quer 1928 (BM).

|  | Nova Scotia | Madeira | Morocco |
| :--- | :---: | :---: | :---: |
| MC | 15 | 14 | $8-9(18)$ |
| SR | 7 | 11 | 8 |



The absence of papillae along the abaxial marginal surface is most variable in E Asia ( $0-36$ cells across, Figs. 3B, 6). This variation is related to taxonomic features that define the species groups and the species themselves as shown in Fig. 6. In Fig. 3B, the numerical data for the Taxus wallichiana Group, and for the NW Himalayan species, T. contorta (Baccata Group), were summarized because leaf stomata bands are consistently bordered by a relatively narrow marginal zone of epidermal cells without papillae—most often 4 cells across (Fig. 6)—as seen also in the Baccata Alliance (Fig. 3A, Appendix). Asian plants with a marginal border of fewer than 4 cells across generally belong to Taxus wallichiana var. yunnanensis (W.C. Cheng \& L.K. Fu) C.T. Kuan, or to T. suffnessii Spjut, whereas specimens with an abaxial leaf margin exceeding 7 cells in width usually belong to the Sumatrana Group and Cuspidata Alliance, but there are several notable exceptions. One is Taxus chinensis with a leaf margin of $4-12$ cells wide that is clearly intermediate between the Sumatrana Group and the Cuspidata Alliance, corresponding also to its intermediate geographical posi-


FIG. 5A. Examples of T. canadensis var. adpressa from North America and Europe. MC refers to number of cells without papillae between the margin and stomata band. SR refers to the number of stomata rows in a stomata band. Specimens from left to right: Finks.n. (US), Anderson s.n. (US: 1091452), Asplund s.n. (US), and Berglund s.n. (S: C-2066).

|  | lowa | Norway | Sweden | Slovenia |
| :--- | :---: | :---: | :---: | :---: |
| MC | $12-15$ | 18 | 22 | 24 |
| SR | 6 | 5 | $5-8$ | $4-6$ |


tion in central China. Another is the central Himalayas T. contorta var. mucronata Spjut, a variety that is recognized by shorter reflexed leaves as in T. umbraculifera (Cuspidata Alliance) but also with slightly more stomata rows per band ( $8-11$ ) and a wider leaf margin ( $8-10$ cells across) than what is usually seen in the typical variety of NW Himalayas.

Data in Figs. 3B and 6 also contrasts the wide variation in the number of marginal cells in the Sumatrana Group and Cuspidata Alliance with other Asian taxa. The Sumatrana Group and Cuspidata Alliance share the elliptical shape of epidermal cells as seen in T-section, in contrast to the angular leaf epidermal cells of the Wallichiana Subgroup, and also lack of papillae on the abaxial midrib as well as along marginal regions (e.g., T. sumatrana, Fig. 1C). This group usually occurs at elevations below 2000 m in contrast to T. wallichiana found mostly above 2300 m . One exception, T. kingstonii Spjut, in the Sumatrana Group, is ecologically and morphologically intermediate between the Wallichiana and Sumatrana Groups.

Variation in the number of abaxial marginal cells recorded from the same plant, or related plants at the same locality, was assessed similarly to data compiled on number of stomata rows. Among duplicate herbarium specimens, the count was the same in nearly half of the duplicate sets. Most variation within individuals, or within a population of individuals, occurred in the Sumatrana Group (T. celebica [Warb.] H.L. Li, T. kingstonii, T. mairei [Lemée \& H. Lév.] S.Y. Hu ex T.S. Liu, T. sumatrana; Fig. 3B, Appendix). For example, leaves from duplicate specimens of T. mairei from Guangdong often lacked papillae along the abaxial margin on either 14 or 24 cells across. Similar dimorphic differences are evident in specimens from Sichuan and Guizhou, and in T. kingstonii from Yunnan. These differences may be due in part to leaves from different plants, or from different ages of shoots, or from different heights on the plant (de Laubenfels 1988), or in T. celebica, from slightly different regions of the leaf.

In the case of Wilson 1265 (A, BM, K, S, US), which was mentioned earlier as having been collected from three different locations (Rehder \& Wilson in Sargent 1914), one site in western Sichuan at 600-650 m included duplicate specimens that was found to have either 15 (US) or 21 (A) stomata rows per band, and either 16 (US) or 25 (A) marginal cells. Wilson also collected seed (Wilson 1265) from Sichuan near Mt. Emei and/or Yachou Fu at 600 m from which leaves in three herbarium specimens obtained from a plant grown from seed (of Wilson 1265) at the Royal Botanic Gardens-Kew lacked papillae entirely along an abaxial margin zone, 18 or 28 cells across, while all three had $8-10$ stomata rows per band. This plant is not T. chinensis as indicated in the literature (Rehder \& Wilson in Sargent 1914), but Taxus mairei var. speciosa (Florin) Spjut that appears atypical by the relatively large greenish distant leaves that are more characteristic of T. celebica. Photographs of a very similar plant in cultivation at the Royal Botanic Garden-Edinburgh (probably from Wilson 1265 seed) are shown in van Gelderen and van Hoey Smith (1996) and in Krüssmann (1985). However, Wilson 1265 (b) does include one specimen I identified as T. chinensis that was reportedly obtained from western Hubei south of "Ichang," 600-1300 m.


FIG. 6. Number of Taxus specimens according to the number of bare (without papillae) cells across abaxial surface of leaves between the margin and stomata band for selected taxa from E Himalayas to China. Note that the Sumatrana Group has the widest leaf marginal zone without papillae, and that $T$. wallichiana has the narrowest zone, mostly 4 cells wide (40 specimens).

De Laubenfels (1988), commenting on his field observations of yews in Taiwan, suggested that differences in leaf shape may be seen on the same plant and further implied the same for the presence or absence of leaf papillae.

Leaves of Taxus mairei var. speciosa that showed considerable variation were also studied from plants cultivated in the United States. An examination of 20 leaves (Phyton s.n.)—from apical buds to $3^{\text {rd }}$ yr branchlets-were found to be relatively constant in the number of marginal cells without papillae-9 cells across-and also in having 16-18 stomata rows per band. They were notably variable in shape and length of epidermal cells, especially juvenile foliage.

The abaxial leaf margin in T. mairei specimens obtained by C-j. Chang from near Hualien, Taiwan varied by four cells (4-7) at four of six locations (Nos. 2, 4, 9, 10), and by only two cells ( $0-1$ cell) at the two other locations (Nos. 1, 5).

Finally, specimens of T. celebica from South Vietnam by Schmid (1974) were found to lack papillae on either (23-) 24 or 32 cells across the leaf margin. Here Schmid (1974) reported that Taxus was polymorphic, and among his specimens at the Museum of Natural History in Paris (P), is an apparent hybrid (Schmid s.n.) between T. aff. chinensis (Poilane 4150) and T. celebica.

DISCUSSION
Phytogeography of Taxus.-Data presented for leaf character attributes of Taxus (Figs. 2 and 3) show that stomata rows and marginal cell features are most diverse in SW China, while the same number of subspecies (Pilger 1903), or species (Farjon 1998; Silba 1984) have been recognized to occur in both North America and Eurasia; thus, the traditional separation of Taxus species (or subspecies) based on these geographic discontinuities is a distorted classification. The phytogeographic data in this study support the taxonomy of yew for only the geographical disjunct occurrences in North America.

In North America, leaf stomata of Taxus brevifolia show a cline in number of stomata rows ranging from 9 rows in California to 4 rows in the northern Rocky Mountains, and also in length of abaxial epidermal cells relative to width $(1 / w)$ from an average of ca. $3 \times 1 / \mathrm{w}$ to $8 \times 1 / \mathrm{w}$. Molecular differences have been reported between coastal and inland yews (El-Kassaby et al. 1994, 1995) at more northern localities. Clinal
variation in conifers has been linked to historical migrations and hybridization patterns with the advance and retreat of glaciers since the Pliocene (Wilkinson et al. 1971); however, the cline in stomata data for the North American Wallichiana Subgroup, including Mexico, seems best explained by loss of stomata in leaves of Taxus as it may have migrated northwards during the Neogene, when the climate may have become increasingly warmer and drier, while the differences within the Pacific Northwest may be a product of more recent climatic changes (Graham 1999).

Also, a geographical species concept that recognizes T. sumatrana as widely distributed in SE Asia (de Laubenfels 1988) might conclude that its distribution was achieved from long-distance dispersal by birds, whereas geographical disjunction of Taxus in North America has been correlated with paleobotanical data (Graham 1999). This seems paradoxical; i.e. the greater variation in leaf anatomical data of Taxus in Asia should also be explained by evolution and paleogeography—perhaps the result of climatic and geomorphic changes that have occurred since the Cretaceous, a period of 130 million years (my). Therefore, the discussion that follows will focus on this latter hypothesis.

Although data on leaf stomata rows in Taxus are more variable in Asia than in North America and Europe, relationships become evident when other taxonomic features are taken into consideration (Spjut 2007). For example, in the western Himalayas, the stomata counts that range from 5-8 ( -10 ) or 8-11 stomata rows per band (Fig. 2C) are a characteristic feature of T. contorta Griff. This species is also recognized by the long narrow leaves that have idioblasts in the spongy mesophyll (vesicular cells appearing dark red in herbarium specimens), a character trait not seen in the E Himalayan yews. Moreover, these features show a closer relationship to European yews than to Asian yews. In the E Himalayas, T. wallichiana-indicated to have 11-19 stomata rows-is recognized by leaves having large angular shaped epidermal cells as seen in T-section, by the persistent bud-scales at the base of branchlets, by the branchlets that show a marked color change in their $2^{\text {nd }} \mathrm{yr}$ of growth—from yellowish green to maroon or reddish orange, and by the bone-like parenchyma cells in the spongy mesophyll that connect in a reticulate pattern with rounded to angular intercellular spaces. These morphological features are considered more closely related to yews of SW China than to T. contorta of W-C Himalayas.

Data in the appendix take into account variation in $T$. chinensis and T. wallichiana on Mt. Emei. Specimens are arranged according to increasing number of marginal cells along the abaxial surface of the leaf without papillae. In T. wallichiana, the epapillose marginal cells, which are consistently 4 wide for numerous specimens in the Himalayan Region (see also Fig. 10 in Spjut 2007), appear to show greater variation on Mt. Emei where it was found that two of six specimens had a leaf margin 8 cells wide. Similarly, T. chinensis outside of Mt. Emei was usually found to have a relatively narrow leaf margin of 4-7 cells wide, 25 of 30 specimens ( $83 \%$ ); only 2 specimens ( $7 \%$ ) were found with a leaf margin greater than 8 cells wide, whereas on Mt. Emei, 11 of 30 specimens ( $37 \%$ ) had a relatively broad leaf margin ( $8-12$ cells across). The broader leaf margin in T. chinensis from Mt. Emei could be the result of recent hybridization with species of the Sumatrana Group, or possibly reflects historical introgression with T. umbraculifera of NE China. Hybridization might also account for similar variation in T. chinensis for three specimens from Guizhou, Shaanxi, and Vietnam.

The development of leaf papillae in Taxus along the abaxial marginal zone may be partially correlated with latitude as evidenced by the narrower range of marginal cells without papillae ( $7-24$ cells across, Figs. 3B, 6) for the T. cuspidata Alliance in temperate NE Asia, compared to that of the more widely distributed Sumatrana Group (8-36 cells across, Figs. 3B, 6) in SE tropical Asia. At increasingly higher latitudes, plants with more papillae on their leaves obviously receive greater protection from ultraviolet rays of the sun-during the longer summer days. The refractivity (protective) effect of papillae on Taxus leaves has indeed been mathematically demonstrated (von Frimmel 1911). Nevertheless, hybridization between the tropical and temperate species alliances in E Asia cannot be ruled out.

In the Cuspidata Alliance, I have observed that papillae are of lower stature and concrescent near cell walls in which the cell walls appear thicker, examples of which are shown in Jinxing and Yuxi (2000). This may be evidence of introgression with the Wallichiana Group from which T. chinensis allegedly evolved. As
indicated, leaves of $T$. chinensis often have elliptically shaped epidermal cells in T-section, a slightly wider marginal border, ranging from 4-12 smooth cells across (Fig. 6), and midrib papillae often more conspicuous along cell walls. Thus, the Cuspidata Alliance, which is undoubtedly related to the Baccata Alliance (Collins et al. 2003; J. Li et al. 2001), may have acquired an expanded leaf margin as a result of hybridization with species of the Sumatrana Group.

In cultivated individuals related to T. cuspidata and T. mairei, papillae sometimes were found on midribs of young leaves, but not the older leaves. However, the odd leaf mentioned earlier for one cultivar ( $T$. caespitosa) with 16 instead of 13 stomata rows/band was found with low papillae on its midrib, whereas the other 15 leaves had smooth midribs; this odd leaf may have retained juvenile characteristics due to lack of exposure to light. I have also completely "skinned" leaves to evaluate the distribution of papillae from base to apex in specimens from Europe, Taiwan and the Philippines, and have found papillae to develop more in the upper half (towards apex). The presence of midrib papillae on juvenile leaves, thus, may indicate an ancestral trait that should not be treated as a justification for lumping all variation within a geographical area under one species.

This alleged ancestral trait is also evident among specimens that are intermediate between T. chinensis and T. mairei, and the extinct T. engelhardtii (Fig. 7). The characteristics of T. mairei include larger (mamillose) epidermal cells on the abaxial midrib and marginal zones, and isodichotomous zigzag branching; those of T. chinensis are the marginal papillae on the abaxial midrib [e.g., Ching 1676 from Sichuan; Chiao \& Fan 464 (US) from Sichuan, and Tsiang Ying 1425 (P)]. A study by Kwei and Hu (1974)—that mentioned 30 of the specimens cited in the Appendix—recognized intermediates by a partially papillose midrib; however, Spjut (1992, 1993, 1998a) has since reported other correlative taxonomic characters—such as shape of leaf epidermal cells, development of papillae along the abaxial marginal zone and size of bud-scales-to help further separate these species. It should also be noted that midrib papillae can be consistently present in the W Himalayan T. contorta (Kvaček 1984), or consistently absent in the North American T. canadensis.

From Myanmar are four specimens found to have leaves almost entirely papillose within a few cells from the abaxial margin. Three of the specimens are recognized as belonging to a distinct species (T. suffnessii) by the relatively large and persistent bud-scales at base of branchlets, by the conspicuous papillae on epidermal cells, and by the relatively tall-rectangular epidermal cells as seen in T-section of leaves (Spjut 2007). One of two other specimens from NE India and Bhutan (Ludlow \& Sherriff 18762, 3719)—that was recorded to be papillose within 2 cells from the margin—differed by having elliptical instead of angular epidermal cells. It would appear, then, that the occurrence of papillae on the abaxial surface of Taxus leaves has taxonomic significance even when the numerical differences are relatively narrow as also seen in North American $T$. globosa var. globosa and T. globosa var. floridana in which intermediates are recognized to occur in northern Mexico.

Disjunct Relationships between Eastern Asia and Western North America.—Disjunct geographic distributions in Taxus and other genera have long been recognized between temperate North America and Eurasia (Axelrod 1983; Boufford \& Spongberg 1983; Good 1964; Graham 1972; Hara 1972; Kornas 1972; H. Li 1952; Tiffney 1985a; Qian 2002; Q. Wang et al. 2006); however, their rate of evolution varies. For conifers this has been considered relatively slow (Prager et al. 1976). In the genus Abies, for example, the subalpine fir in W North America [A. lasiocarpa (Hook.) Nutt.] appears more closely related to an endemic species of Taiwan [A. kawakamii (Hayata) Ito] than to any of the 10 other American species (Farjon 1990; Hunt 1993; Liu 1971). Indeed, recent molecular studies by Suyama et al. (2000) show A. mariesii Masters of Japan to be more related to species in North America than to its relatives in Japan. Additionally, species of Pseudotsuga ( $\pm 4$ spp., Farjon 1990) in Asia may have been derived from ancestors in North America (Strauss et al. 1989).

The Wallichiana Subgroup of Taxus—characterized by angularly shaped epidermal cells in T-section—occurs primarily in E Himalayas to SW China (Sichuan, Yunnan) and in North America (Spjut 1998a, 1998b, 2000a). Within this subgroup, leaves of Yunnan and Sichuan plants (T. florinii, Spjut) appear indistinguishable from those of the American T. globosa (Spjut 1998b, 2000a, 2000c). Other specimens from Myanmar


FIG. 7. Comparison of leaves of extant T. mairei (clear photos, isotype, P) with extinct T. engelhardtii (grainy photos, reproduced from Kvaček 1984), from an Oligocene deposit in Bohemia.
(T. suffnessii) are similar to $T$. brevifolia in the relatively large bud-scales and tall rectangular epidermal cells as seen in T-section (Spjut 2000c).

An analogous disjunct relationship is seen among the white pines, Pinus monticola Douglas ex D. Don of W North America and P. wallichiana A. B. Jackson of Myanmar (Axelrod 1986, Pinus griffithii [Hook. f. \& Thomson] Parl.). They are remarkably similar in cone morphology and needle chemistry. The antiquity of this relationship is supported by their turpentine chemistry of saturated straight chain hydrocarbons-undecane and heptane, the chemical structures of which are considered more archaic among the terpenoid compounds in pines (Mirov 1953). Additionally, heptane occurs in the Mexican P. ayacahuite Ehrenb. ex Schltdl. (Mirov 1953), along with a bicyclic sesquiterpene—cardenine—that has also been found in P. parviflora Siebold \& Zucc. of Japan (Mirov 1953). The close relationship among these species, which belong to sect. Quinquefoliae subsect. Strobus, is supported by molecular data (Liston et al. 1999) from which it has been suggested that the ancestors probably originated in the "Old World" (Gernandt et al. 2005).

In angiosperms, it is interesting that Phipps (1983) recognized—among ~145 species of hawthornsCrataegus mexicana Moç. \& Sessé, a widely distributed species in Mexico and Guatemala, to have its closest relative in Yunnan, C. scabrifolial (Franchet) Rehder, and that both are the "most primitive" of a taxonomically complex Laurasian genus, which has numerous species in both Mexico and in Yunnan.

For taxads and other conifers, diversity is greatest in SW China (Figs. 2, 3; Cheng \& Fu 1978; Prakash et al. 1995; Qian \& Ricklefs 1999). Ancestors related to Taxus suffnessii Spjut in Myanmar (Appendix) may have immigrated to North America across a former Aleutian (or Bering) land bridge (Hamilton 1983; Millar 1993)—as suggested for Crataegus (Phipps 1983). A logical time for this to occur would have been during the latter half of the Cretaceous (110-100 mya), after Pangaea had fragmented (Graham 1993) —when an epeiric sea (Wolfe 1975) possibly divided the North American continent into distinct west and east floras (Graham 1999; Srivastava 1994; Thorne 1972, 1978). Late Cretaceous fossils related to the Alaska cedar, Callitropsis nootkatensis (D. Don in Lambert) Florin, which includes one related sister species in North Vietnam, and is also sister to other species in North America (Little 2006), have been found on Vancouver Island (McIver 1994), and an early Cretaceous fossil, Chamaecyparis eureka Kotyk, from Eureka Sound in the Canadian Arctic, is most similar to the extant Ch. pisifera Siebold \& Zucc. in Japan (Kotyk et al. 2004). Additionally, fossil cones of Thuja smileya LePage from Late Cretaceous deposits on the North Slope of Alaska are indistinguishable from modern species (LePage 2003).

As climate temperatures declined during the Cretaceous (Axelrod 1958; Frederiksen 1994; Graham 1999; Novacek 1999; Srivastava 1994), Taxus might have retreated southwards, perhaps reaching southern Mexico by the end of the Cretaceous (65 mya); similar retreats have been suggested for other genera (Phipps 1983; Sharp 1966), but for the Tertiary Period (Phipps 1983), not the Cretaceous. A later migration and extinction of Taxus, such as in the Tertiary near the Eocene-Oligocene boundary, may seem like a more reasonable time frame for evolution of North American Taxus, but there also has to be ample time for diversification of
the Cuspidata Alliance as well as the alleged migration and extinction of the Wallichiana Group across the Sino-Japanese Region.

The end of the Cretaceous is marked by a distinct change in the geochemical and fossil records (McIver 1999; McIver and Basinger 1999; Novacek 1999)—indicating a rapid climatic warming—possibly due to a meteor impact in the Caribbean Sea that might have caused massive volcanic materials to erupt and cloud the atmosphere (O'Keefe \& Ahrens 1989) -a 'greenhouse' calamity that could explain evidence for "ecological deserts" (Tschudy et al. 1984)—and mass extinction of major taxa (e.g., dinosaurs, Novacek 1999). This could have extirpated yew north of Mexico; Cretaceous fossils of gymnosperms of the Taxodiaceae (Metasequoia, Sequoia, Sequoiadendron), and Amentotaxaceae (Amentotaxus, Torreya) are known as far south as New Mexico and North Carolina (Florin 1963) for which Taxus has had a long history in association (Florin 1951, 1963; Kvaček 1984) but whose fossils may not always be preserved or identified.

Paleontological evidence indicates that following the Cretaceous a warmer subtropical humid climate (Chaney 1947; Frederiksen 1994; Tiffney 1985a) prevailed over much of North America until the late Eocene (ca. 50 mya; Chaney 1947; Graham 1999; Novacek 1999; Srivastava 1994; Wolfe 1975). Assuming that ancestral T. globosa had survived only in Mexico, a northward migration (as the climate warmed) would account for the cline in leaf stomata data of Taxus in W North America (Fig. 2A). Other North American conifers with evidence of a southern ancestry include Douglas fir (Pseudotsuga menziesii (Mirb.) Franco), most likely derived from big cone fir (P. macrocarpa (Vasey) Mayr), endemic to S California (Strauss et al. 1989), and pines that may have drifted northwards on the San Andreas rift system (Axelrod 1986) —a system that may have included Vancouver Island originating perhaps from as far south as "lands end" off the cape (Cabo San Lucas) of Baja California (90 mya, Ward et al. 1997). A northward range extension of ancestral Taxus globosa may be further correlated with the change in a Rocky Mountain flora from paleotropical (boreotropical) to neotropical elements during the mid Eocene (Leopold \& MacGinitie 1972), and later along the Gulf as evident from biogeographical data on fishes and amphibians (Rosen 1975), maples (Acer saccharum L. Group; Humphries 1982), and other taxa (Burnham \& Graham 1999)—emphasized by Axelrod (1975, 1986).

As the climate became drier with the uplift of the W Cordillera (Chaney 1947; Wolf 1969), the range of Taxus, like other Arcto-Tertiary genera, diminished (Axelrod 1975, 1983; Graham 1993, 1999), while other taxa evolved (Axelrod 1958). Fossils of Taxus have been reported in Eocene (54-38 mya), Oligocene (38-27 mya), and Miocene (27-10 mya) strata of W North America (Gaussen 1979; Kvaček \& Rember 2000, in press; Manchester 1994; Meyer \& Manchester 1997) in association with species of Tsuga, Abies, Lithocarpus, Quercus, Acer, Alnus, Cornus, Carpinus, Castanea, Fagus, Liquidambar, Nyssa, Ostrya, Platanus, Tilia, Ulmus, and Cercidiphyllum (Graham 1999; Whittaker 1961).

Extant species of Taxus are still found with these same genera today in mixed mesophytic forests of S Appalachia (Braun 1950), China (Hou 1983), and Japan (Hayashi 1954). Taxus brevifolia allegedly evolved from an ancestral T. globosa complex as the climate became cooler and drier during the Eocene (56-34 mya; Graham 1999), while closer ties between the Mesoamerican yew and Florida yew were likely maintained until the Pleistocene as evidenced by the close similarity among many shared taxa between the two regions (Sierra Madre Oriental and S Appalachia). It is interesting that specimens of Florida yew appear indistinguishable from those occasionally collected in Veracruz and in Nuevo Leon/Tamaulipas, Mexico (e.g., Meyer E Rogers 2746, BM; Mueller 1337, BM, PH), where they reportedly occur with Carpinus caroliniana Walter, Cercis canadensis L., Frangula caroliniana (Walter) A. Gray, Hamamelis virginiana L., Liquidambar macrophylla Oerst, Magnolia schiedeana Schltdl., Prunus serotina Ehrh, and others also found in S Appalachia, including many lichens and mosses (Culberson et al. 1990; Graham 1973, 1999; Miranda \& Sharp 1950). Moreover, it has been shown that the Florida yew and Mesoamerican yew form a clade with the Pacific yew as a sister species (J. Li et al. 2001), and that the Florida and Mesoamerican yew are indeed more closely related (J. Li et al. 2001).

As previously noted, genera found with the Mesoamerican yew also occurred with Pacific yew (Graham 1999), but in the Pacific Northwest many of these genera perished—such as Carya, Disopyros, Fagus, Hamamelis, Liquidambar, Liriodendron, Magnolia, and Morus (Axelrod 1975, 1983, 1986; Graham 1999; Manchester

1999; Wood 1972). Liriodendron is one of many known from fossils in Europe and W North America with relatives now surviving only in E North America and E Asia (Axelrod 1983; Manchester 1999; Schuster 1976). They perhaps were part of a widespread Tertiary "boreotropical" forest (Wolfe 1975; Graham 1999), which may have included T. canadensis (Figs. 4, 5).

As conifer diversity declined and grassland vegetation expanded during the Miocene (Axelrod 1976; Jacobs et al. 1999), Taxus possibly had attained maximum diversity in geographic and ecological species isolation. McIver and Basinger (1989) found in Eocene deposits cones similar to western red cedar (Thuja plicata Donn ex D. Don) that may have been derived from an earlier complex related to the extinct Thuja polaris McIver et Basinger, which they described from a Middle Paleocene deposit on Ellesmere Island, whereas other cedars related to Th. occidentalis L. are not known before the Miocene (McIver and Basinger 1989). Moreover, Th. occidentalis is recognized in the fossil record from the late Pliocene (Bennike 1990). The redwood, Sequoia sempervirens (Lamb. ex D. Don) Endl., is hardly distinguishable from a former widespread S. abietina (Brongn.) Knobloch—known from the Upper Eocene to Upper Miocene (Mai 1998). A species of Taxus from a Middle Miocene deposit in N Idaho (Clarkia area Latah Formation) has nearly the same abaxial epidermal features seen in the extant T. brevifolia (Kvaček \& Rember 2000, in press). Klicka and Zink (1997) concluded from DNA evidence that North American species of song birds had already originated by early Pleistocene, and that subsequent glaciation was more of an "obstacle course" for their survival. Yew species, by comparison, are likely to evolve more slowly as a yew trunk may live 3000 years (Larson et al. 2000; Thomas \& Polwart 2003; Voliotis 1986), and still may survive by producing trunks from adventitious shoots (Hageneder 2007; Loudon 1844). The greater diversity of Taxus in SW China may also be related to less species extinction there as seen in many plant genera as a result of increasingly drier and cooler climates that had a more profound impact on the vegetation elsewhere since the Middle Miocene (Axelrod et al. 1998; Kubitzki \& Krutzsch 1998).

Relationships between Eastern North American and Eurasian Taxus.-While the Pacific floristic element of Taxus in North America is represented by three disjunct taxa within the Wallichiana Group, the Atlantic floristic element has only T. canadensis, a species that appears more related to the Baccata Group (J. Li et al. 2001; Spjut 2007) than to T. globosa by its elliptical shaped epidermal cells in T-section, and by its subcylindrical seed shape (Spjut 1998a, 2000). The lack of papillae on the abaxial leaf surface between the margin and stomata band that characterizes the North American Canada yew is also seen more frequently in yews of temperate NE Asia (Cuspidata Alliance) than in the Euro-Mediterranean (Baccata Alliance), and its leaf epidermal features are most similar to T. biternata Spjut, a species closely allied to T. cuspidata. The close relationship between T. canadensis and T. cuspidata is supported by molecular data (Collins et al. 2003). Taxus biternata differs from T. canadensis by the 2-3 angled seeds (tapered part) developing on $1^{\text {st }}$ yr branchlets, and by the tree habit (Spjut 2007). The complete lack of papillae along the abaxial leaf marginal zone is a relatively rare occurrence in European yew ( $<2 \%$ ), but this character trait may have once been common in that region; for example, three species described by Kvaček (1984) from leaves of fossil assemblages in Europe-dating from Oligocene to Pliocene-all lacked papillae entirely between the stomata bands and margins.

Furthermore, a "Taxus (sp. 1," Kvaček 1984; Fig. 8) of Lower Miocene age is, in my opinion, T. canadensis. Its leaves are more similar to American plants than to European plants, which differ by the distinctly papillose stomata bands—except perhaps for rare North American specimens (e.g., Coy \& Glen from Ithaca, New York). These extant European variants could be referred to the extinct T. grandis Kräusel or T. inopinata Givulescu (1973)—described from Tertiary deposits in Europe (Kvaček 1984). Data for numbers of stomata rows (Figs. 2A, 2B) also support my hypothesis that the North American T. canadensis came from Europe, possibly arriving late Paleocene or Eocene when migration across the Atlantic was possible by land (McKenna 1983; Tiffney 1985b), as suggested for the evolution of Cornus sessilis Torr. ex Durand (Xiang et al. 2005, 2006). During this period the Gulf Coast flora shows evidence of many immigrants from Europe (Frederiksen 1994, 1995) that included species of Fagopsiphyllum, Hydrangea, Iodes, Koelreuteria, Langtonia, Nyssa, Palaeophytocrene, Pentoperculum, Platanites, Platycarya, Pyrenacantha, Sargentodoxa, Symplocos, Tapiscia, Tetraclinus, and Tilia (Manchester

1999). It is interesting to note that 10 species (in 9 genera) of lichens recently discovered to occur in E North America—on Mt. Katahdin in Maine—were previously known only from northern and/or central Europe, except for one species that also occurs in Greenland and Siberia (Fryday 2006).

Relationships between the Euro-Mediterranean and Asian Taxus.-The European yews have all been considered a single species, T. baccata; however, additional species appear evident. The lectotype (of T. baccata) has leaves arranged mostly parallel to one other along two sides of a branchlet in a flat spray with $8-10$ stomata rows/band and a papillose undersurface-from the midrib to near the margins. The typical European yew is further characterized by isodichotomous branching, pale glaucous green leaves similar in color on both surfaces, and cones maturing on branchlets that have terminated their growth as evident in specimens from England, Germany, Austria, Switzerland, Czech Republic, Spain, Portugal, Algeria, Morocco, Italy, Albania, Bosnia, and Turkey. This is in contrast to another widespread species, T. recurvata, that I recognize by a less parallel arrangement to the leaves with a sharper contrast in color between the leaf surfaces-notably yellowish green on the abaxial surface and dark green on the adaxial surface, and by cones developing on branchlets that continue their growth. Both have many naturally occurring varieties based on differences in leaf arrangement, leaf texture and branching, and leaf anatomical differences.

Intermediates include specimens similar to T. contorta in the W Himalayas. Examples are characterized by long linear $\pm$ distichously arranged leaves that in relative thickness to width (as seen in T-section) are similar to either the W Himalayan T. contorta (1.5-2.0 mm wide, $<0.5 \mathrm{~mm}$ thick, e.g. Biol. Inst. Dubrovnik from Bosnia, Barabas from Romania, Davis 13667 from Turkey), or to the E Asian T. biternata Spjut (2.0-2.5 mm wide, 0.25-0.33 mm thick, e.g., Petrak from Czech Republic, Moniz from Madeira, Handel-Mazzetti from Greece, Anderson 42 from Bulgaria, Davis \& Hedge 32208 from Turkey). The W Himalayan T. contorta is distinguished from most Euro-Mediterranean T. baccata by the presence of dark red, or sometimes yellowish, parenchyma cells (idioblasts) in the leaf spongy mesophyll (in herbarium specimens); however, European specimens occasionally have idioblasts in the leaf mesophyll. In fresh specimens, the intermediates would probably be difficult to distinguish. Of further significance is that the abaxial leaf midrib of T. contorta is always papillose (Kvaček 1984), whereas in T. baccata, the abaxial midrib varies from smooth to papillose.

I suggest that ancestral T. contorta entered the Himalayas from the north during the Miocene uplift (Krishnan 1974), or earlier (Najman \& Garzanti 2000), before it arrived in Europe (Frederiksen 1995). Its leaves would likely have lost stomata in adapting to the rising Himalayas where environmental selection would also likely favor the development of papillae on the abaxial leaf midrib (von Frimmel 1911).

As glaciers advanced during the early Pleistocene (2.5-1 mya), the cooling temperatures may have led ancestral T. contorta to also retreat into Europe where it then allegedly hybridized with other species of Taxus that may have flourished in a subtropical evergreen laurel-conifer forest (Axelrod 1975; Klaus 1989; Kvaček 1984; Mai 1989; Palamarev 1989), but may have found refuge in ravines and coastal areas. Similar patterns of evolution have been suggested for European species of Abies (Fady et al. 1992). The significance of numerous European refugia for conifers was suggested by Fady-Welterlen (2005) to account for their "significantly higher" "within species diversity" "than that of other conifer species worldwide." In this regard, it is interesting to note that the association of Taxus with "ancient forests" on limestone cliffs in Iowa and in Europe is partly attributed to the topography of the habitat that offers protection from Homo sapiens (Larson et al. 2000), whereas a severe decline in European Taxus is generally recognized (Heinze 2004).

Taxus engelhardtii Kvaček, described from a late Oligocene deposit in "NW Bohemia," was associated with a mixed mesophytic forest with prevailing broad-leaved componets (Kvaček 1984; Kvaček \& Walther 1998) that included Laurophyllum (4 spp.), Cercidiphyllum, Liriodendron, Acer, Ostrya, Betula, Craigia and other genera. Its leaf shape and arrangement is much like T. mairei in Sichuan, Yunnan (Fig. 10) and Guangdong where similar forest types still occur today. Vegetation in these areas-characterized by a distinct dry season—includes species of Cercidiphyllum and Liriodendron (Hou 1983) that have since become extinct in Europe. In Vietnam, the closely related T. celebica occurs in a laurophyll oak forest from 1000-1600 m in elevation in association with Cinnamomum obtusifolium Roxb. ex Nees, Michelia foveolata Merr. ex Dandy, M. mediocris Dandy, Castanopsis fissa (Champion ex Benth.) Rehder \& E. H. Wilson, Quercus bambusifolia Hance
and others (Dung 1996; Schmid 1974.). Extant yews in Europe, however, bear little resemblance to T. engelhardtii; therefore, it may not have contributed to hybrid complexes that are now apparent in Europe. Rather it may have simply been extirpated from the Euro-Mediterranean Region due to changes in the climate.

However, another European complex appears to have been derived in part from the Irish yew (T. fastigiata Lindley), a possible relict of a former subtropical forest. Known initially from two trees in Ireland-that were transplanted at the time of discovery ( $\sim 1770$; Veitch et al. 1881), it has always been regarded a distinct yew even though taxonomists have included it under T. baccata (Loudon 1844; den Ouden \& Boom 1965). Its linear dark green leaves that are spirally arranged in whorls seem less evolved. Moreover, such radial phyllotaxy is infrequent among yews-occurring at disjunct locations in the Old World, particularly coastal regions-in Spain, Morocco, British Isles, Honshu, Hokkaido, and Sakhalin. Leaves of the Irish yew in cultivated specimens from widely scattered locations were found to be remarkably similar in lacking papillae along 6-15 cells across an abaxial marginal zone and on the midrib as well; these specimens, which are not included in the Appendix, are from Australia (Boorman, New South Wales, A), North America (Spjut s.n. Oregon, California, Maryland, wba), and Europe (Baker, Yorkshire, BM; Stewart Hort., Florence Court, Ireland, K; Baenitz, Lusitania, S). Perhaps European yews during the Tertiary were more like those now seen in E Asia but have since acquired more papillae on their leaves through introgression with ancestral $T$. contorta, the alleged replacement species. Many yew specimens with dark metallic green foliage from Great Britain appear intermediate between T. contorta and T. fastigiata; examples are the "Dovaston yew" (T. baccata var. dovastoniana) and the English yew (in England, T. recurvata).

Evolution within the European T. canadensis complex is also evident as seen in leaves of one specimen from the former N Yugoslavia (Slovenia, Fig. 5) by the relatively fewer (4-7) stomata rows per band and inflated epidermal cells. Related plants in Madeira, southern France, Norway, and Sweden have more stomata (5-9 rows/band) and less inflated epidermal cells. Characteristics of the Slovenian yew (obtuse leaf apex, 4 stomata rows/band, wedge-shaped epidermal cells) are evident in a fossil leaf from a Pliocene deposit in Bohemia, Czech Republic ("Taxus sp. 2," Kvaček 1984).

The increase in cell size and loss of stomata in leaves of the Slovenian yew may reflect adaptation to changes in a climate from a warm temperate humid type with uniform distribution in rainfall towards a climate with more pronounced warmer and drier seasons. The Yugoslavia region is also one of 33 sites in the Euro-Mediterranean region with "Paleomediterranean" woody taxa known from Oligocene, Miocene, and Pliocene deposits (Palamarev 1989). Thus, the Slovenian yew may be a relict of a former Mediterranean montane flora that included the conifer genera Pinus, Juniperus, Tetraclinus, Abies, Cedrus, Cupressus, and Picea (Palamarev 1989); some of these are reported with this Pliocene yew (Kvaček 1984).

I also distinguish T. mairei from T. sumatrana by the relatively short inflated epidermal cells on the abaxial midrib (Appendix; Spjut in adnot. and on illustrations of packets, A, GH, Jun 1996; Spjut 1998b, 2007). The occurrence of this species in China corresponds mostly to the "broad-leaved evergreen forests of the subtropical zone" of Hou (1983) with a climate marked by "distinct dry seasons" - "on mountains below 1100 m in the eastern humid subtropics, or on mountains between 1500 and 3000 m in the western subtropics of the Yunnan Highland" (Hou 1983). Similarly, T. brevifolia, a species confined to the North American Mediterranean climate, has wider and taller marginal epidermal cells and fewer stomata (Fig 1B), compared to its putative ancestor, T. globosa (Spjut 1998a, 1998b) that has evidently survived in the montane cloud forests of Mexico and Central America. The evolution towards larger epidermal leaf cells has also been noted between fossils and living species of Amentotaxus (Ferguson 1978).

The variation in leaf anatomical data for Taxus in SW China is also related to the convergence of different floras in that region (Bartholomew 1999; X-w. Li \& J. Li 1997; Zhengyi \& Sugong 1998). These include the "Turkmenian" in W Himalaya, Tibetan or "Indo-Chinese", Malayan, and Sino-Japanese (Mani 1974; Rao 1974; Rau 1974). In the W Himalayas Taxus is represented by the neoendemic T. contorta, usually with 7-8 stomata rows per band, and in the eastern region by the paleoendemic Wallichiana Group with 15-18 (-21) stomata rows. The relatively lower numbers for stomata rows in $T$. wallichiana ( $11-15$ ) and higher numbers for T. contorta (9-11)—where these taxa overlap in their distribution-is undoubtedly due to hybridization
and introgression (Fig. 2C). Further evidence for hybridization involving T. contorta is seen in the wider marginal region of cells along the abaxial surface of leaves (MC, Appendix) and the wider angle of leaf divergence from twigs in plants from Nepal and Bhutan (Spjut 2006).

The wide range in leaf anatomical traits for yews of SW China (Fig. 2C, 3B) is also a product of a long evolutionary history of tropical and temperate vegetation types with possibly less extinction of taxa during glacial climates (Hsü 1983), in comparison to greater glacial devastation to the floras of North America and Europe. The oscillating wet and dry periods during the Pleistocene may have led to many new combinations in Taxus between anatomical and gross morphological features that were once distinctly correlated with eco-geographic differences prior to the Pleistocene. Introgression of character traits has been correlated with data on the advance and retreat of glaciers for other conifers-such as between Picea rubens Sargent and $P$. mariana (Mill.) B.S.P. (Bobola et al., 1996), between Picea glauca (Moench) Voss and P. engelmannii Parry ex Engelm. (Wilkinson et al., 1971), among species of Pinus (Axelrod, 1986), and among species of Abies (Fady et al. 1992).

The slow evolutionary rate that I have suggested for yew is perhaps not all that surprising in view of its ability to survive almost indefinitely. Individual yew trunks can live several thousand years or more (Loudon 1844; Larson et al. 2000; Thomas \& Polwart 2003), and when they fall, the plant still survives by adventitious shoots, or by layering (Hageneder 2007; Loudon 1844); thus, it may continue to survive until perhaps a change in climate forces it to either adapt or perish. Prager et al. (1976) calculated a rate of change in the amino acid sequence for Pinaceae to occur once in every 7.5 my . In Taxus this may be longer. By extrapolation from data in Figs. 2, and from paleoclimatic changes earlier discussed, one may hypothesize that one row of stomata may become lost permanently in the Taxus leaf as it adapts to slight changes in climate over a period of 10 my during which time it may also spread a distance of some 3000 km (at the rate of $300 \mathrm{~km} / \mathrm{my}$ ).

## APPENDIX

Data for all herbarium specimens studied according to continental and political regions and taxonomy.

|  | SR | MC |  | SR | MC |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| U.S.A. Florida |  |  |  |  |  |


|  | SR | MC |  | SR | MC |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Rose 55089 (US). Trinity Co., Buckhorn |  |  | MacMillan (PH). Selkirk, 4300 ft . | 6/7 | - |
| Summit, 2600 ft | 6/7 | - | Calder \& Saville 9982 (US). SE of Nakusp | 7 | - |
| Spjut 12307 (wba). Salmon Mts., 2000 m | 6-8 | 7 | Macoun (US). Rocky Mts., Silver City | 6 | 6 |
| Spjut 12307 (wba). Salmon Mts., 1500 m | 5 | 8 | Macoun 2340 (US). Vancouver Is., Victoria | 6 | - |
| Spjut 10171 (wba). Marble Mts., 2000 m | 5 | - |  |  |  |
| Dudley (US). Salmon Mts., Foxtail Ridge | 5-6 | - | Baccata Group |  |  |
| Benson 2228 (US). Shasta Co., Hatchet Creek | 5 | 0 | Taxus canadensis |  |  |
| Grant 1281 (US). Shasta Co., Dunsmiur | 6 | 5 | U.S.A. |  |  |
| Oregon |  |  | Herb. C. W. Minott (US). Ma: Amherst | 7/8 | - |
| Fisher (US). Portland | 7 | - | Bovin \& Blain 753 (PH). Me: Cumberland |  |  |
| Collector, no.? (US). Jackson Co.: Wimer | 6/7 | 2 | Co., 425 m | 6/7 | - |
| Walpole 153 (US). Jackson Co.: Ashland | 6/7 | - | True 164 (PH). Me: Ovis Island, Long Cone | 6/7 | - |
| Coville (US). Imnaha NatI. For., Billy Meadows | 6 | 4 | Gilbert 831 (PH). Ky: Carter Co., Cascade Caverns |  | - |
| USFS (wba). Josephine Co.: above Taylor Creek, Minnow Creek Rd., 650 m |  |  | Allard 12060 (US). Wv: 900-1200 m | 6/6 | - |
| 4 specimens from nearby sites: |  |  | Women's College of Baltimore (US) | 6 | 12 |
| (1) | 5 | - | Palmer \& King 205 (US). Va | 6 | - |
| (2) | 6 | - | Taylor 424 (US). Pa: Bucks Co., Kintersville | 6 | - |
| (3) | 4-5 | - | Eames 3432 (US). Ny: Coy Glen, Ithaca | 6 | - |
| (4) | 6-7 | - | Spjut (wba) Ny: Ithaca | 6 | - |
| Lankford (wba). Clackamas Co., 1060 m | 5-6 | - | Spjut 11778 (wba) Nh:White Mts. Natl. For., |  |  |
| Lankford (wba). Clackamas Co., 930 m | 5 | - | Wildriver, 300 m | 6 | - |
| Lyall 1860 (K). Columbia River | 5 | - | Stevenson (US). Vt: Willoughby Lake | 5/6 | 12 |
| Nuttall (K: type). Columbia River | 5 | - | Weatherby 5977 (US). Ct: Boston Hollow | 6 | 11 |
| Spjut 12301 (wba). E Cascades E of Portland | 4-6 | 10 | Sheldon (US). Mn: Towers St. Laus | 6 | 12 |
| Beattie 5046 (US). Josephine Co.: 2270 ft | 4-6 | - | Fellows 5686 (US). Me: Rockport | 6 | 12 |
| Cusick 3405 (US). Eastern Oregon | 4 | 3 | Spjut 12179 (wba). Ohio: Secrest Arboretum | 5/6 | - |
| Washington |  |  | Shreeve 1971 (US). Md: Garrett Co., Bailing |  |  |
| Spjut 12302 (wba). E Cascades E of Seattle | 6-7 | 10-11 | Spring | 5/6 | - |
| Horner (US). Blue Mts. | 6 | - | Travis 119 (PH). Me: Cumberland Co. | 4-6 | - |
| Meyer 1589 (US). Thurston Co., Mud Bay | 5-6 | 6 | CANADA: Ontario |  |  |
| Grant s. n. (US). Cascade Mt. | 6 | - | McDonald 223 (US). Ontario: Sagastaweeki Is. |  | 12 |
| Cantwell (US). Orcas Island | 4/5 | - | Rouleau 2700 (US). Humber Dist., Twin Lakes | 7/7 | - |
| Fosberg (US) King. Co., Stevens Pass | 4/5 | - | Quebec |  |  |
| Idaho |  |  | Tae hé \& Lepage 332 (PH). Dartmouth River | 7/8 | - |
| Cronquist 6187 (US). 20 mi W of Riggins, |  |  | Asselin 7212. (US). St.-Charles | 7/7 | - |
| French Creek | 5/6 | - | Pennell 16734 (PH). La Belle Co. | 6/7 | - |
| Shields (wba). Idaho Co.: Allison Creek, |  |  | Bartram \& Long 649 (PH). Rimousk Co. | 6/7 | - |
| 3400 ft . | 5 | - | Fernald et al. 2404 (US). Gaspé Co., |  |  |
| Cochrane (wba). Idaho Co.: Nez Perce Natl. |  |  | Mt. St. Pierre | 6 | 15 |
| For., $5600 \mathrm{ft}$.2 plants | 6 | - | Louis-Alphonse 3547 (US). Baie Missisquo | 6 | 18 |
|  | 5 | - | Lucien 743 (PH). Laurentides, Bellerive | 6/6 | - |
| Montana |  |  | Louis-Maire 686308 (PH). Mé gantic | 5/6 | - |
| Donner (wba) Flathead Natl. For., hr. |  |  | Bovin 1268 (US). St.-Catherine | 5/6 | - |
| Columbia Falls, 3800 ft . |  |  | Chas Mohr (US). Montreal | 4/5 | - |
| middle branch | 7 | - | New Brunswick |  |  |
| Donner (wba) | 7 | - | Malte \& Watson (S: C-2153). | 6/7 | 12-13 |
| Donner (wba) | 4-6 | - | Allen 2528 (PH). St. John | 5/6 | - |
| P. C. Standley 18251 (US). Glacier Natl. Park, |  |  | Nova Scotia |  |  |
| 1400-19850 m | 5/6 | - | Gorham 45139 (US). Halifax Co.: near Halifax, |  |  |
| Thomas 11031 (US). Lake Co.: 8 mi from |  |  | St. Margaret's Bay | 7/7 | - |
| Polson, 3850 ft . | 5 | 4 | Bean et al. 19634 (PH). Yarmouth Co. | 7 | - |
| Steven Wirt 100 (MRC, wba). Flathead |  |  | 19015 (S: C-2155). | 6 | 19 |
| Co.: shrubs | 5-6 | - | (S: C-2156). Victoria Co. | 6/6 | 12 |
| Steven Wirt 100 (MRC, wba) | 5 | - | Pease \& Long 19633 (PH). Cumberland Co. | 5/6 | - |
| Steven Wirt 100 (MRC, wba) | 4-5 | - | Bissell et al. 19632 (PH). Digby Co. | 5/5 | - |

## CANADA British Columbia

|  | SR | MC |  | SR | MC |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Newfoundland |  |  | Lowe with 570 (BM, top specimen) | 8 | 10 |
| Fernald et al. 6738 (PH). Prince Edward Is. | 8/9 | - | Lowe ex Barby (US). Madeira | 7 | 12 |
| Palmer 1300 (US). Bay Is. | 8/9 | - | SPAIN |  |  |
| Palmer 1327 (US). Hermitage Bay, Balena | 7/7 | - | H. Elias 4353 (BM). Burgos: Ser. Obarenes, |  |  |
| Buochan (S: C-2130). | 7/7 | 18 | 1000 m | 11 | 5-8 |
| Rouleau 6545 (US): St. Barbe Distr., E BluePond | 7/7 | 15 | Modesto Laza Palacios (K). Ser. Tejeda \& Almijara, Malacitana Prov. | 10 | 4 |
| Banks1766 (BM). Croque | 7/7 | - | no data 1878 (US). Laguna | 10 | 4 |
| Fernald et al. 26201 (PH). NW Coast | 6/7 | - | Sennen 7087 (BM). Barcelona | 10 | 5 |
| Robinson \& Shrenk (US). St. John's | 6/7 | - | Roivainen (S: C-2075). Guipúzcoa, 900 m | 10 | 5 |
| Rouleau 5533 (US). Gander River | 5-7 | - | Sandwith 4452 (BM), Huesca: Ser. Guara | 9 | 4 |
| Fernald \& Wiegand 4414 (PH). Valley of Exploits River | 6/6 | - | Heywood \& Davis 490 (BM). Ser. Cazoria: Yedra | 9 | 4 |
| Fernald \& Long 27305 (PH). Pistolet Bay | 6/6 | - | Rodriquez (K). Serrania buenia | 9 | 8-11 |
| Wiegland \& Gilbert 27304 (PH). Highlands of St. John | 5/5 | - | Bianor-Maire (BM). Baleares, 1600 m FRANCE | 9 | 5-9 |
| EURO-MEDITERRANEAN |  |  | Fosberg 41055 (US). Jura Mts. | 10 | 4 |
|  |  |  | Endress Aug 1831 (K). Pyrenees | 10 | 6 |
|  |  |  | Endress Aug 1831 (S). Pyrenees | 8 | 5 |
| specimens ranked first by number of stomata by marginal cells for |  |  | ex Herb Comby (PH) | 9 | 4 |
| Baccata Alliance, T. canadensis noted sepa each country. |  |  | Massonnet (K). Pyrenees | 9 | 5 |
| ALGERIA |  |  | Herb. Hook., 1867 (K). Pyrenees <br> Herb. Churchillanum (K). Corsica | 7 | 4 |
| Swingle (NA). Chria near Blida | 12 | 4 | C. Lagerheim \& G. Sjogren Jul 1844 (K). |  |  |
| Reichenbach (K). Atlas, Blida | 9 | 4 | Batsmanshus Paroeciae Elfkarl by |  |  |
| Gamble (K). Atlas des Demia | 8 | 4 | Rosalagiae abundans | 8/9 | 5 |
| Olaptin (S: C-2070), Atlas, Blida. | 7/8 | 4 | Rosalagiae abundans |  |  |
| Davis 52628 (BM). Cedrus forest, 1900-1950 m | 9 | 6-7 | Taxus canadensis <br> Herb. Gombault (S). Sainte Baume | 9 | 18 |
| Univ. Algeria Apr 1912 (NA).Atlas, Blida | 9 | 6 | UNITED KINGDOM |  |  |
| MOROCCO |  |  | Gamble 19866 (K). Berkshire Dist | 11 | 4 |
| Trethewy 85 (K), pendula. Ifrane 1400 m | 10 | 6 | Bowden \& Hillman 433 (BM, globose |  |  |
| Lewalle 8670 (BM). Ifrane 1400 m | 10 | 4 | epidermal cells). Nottinghamshire | 11 | 3-5 |
| Lewalle 8670 (BM). | 8 | - | Gamble 28894 (K). Weltham woods |  |  |
| Lewalle 9670 (BM). Ifrane 1400 m | 9 | 6 | [England SW] | 11 | 5 |
| Davis 49209 (BM). Ifrane 1700 m | 9 | 5 | Albarnes 26 (K). Dorsey: churchyard |  |  |
| Lynes (BM). Mid Atlas, Azrou, 5700 ft | 9 | 6 | [England SW], 350 ft | 10-11 | 8 |
| Haout 938 (BM) | 9 | 5 | Michaelstone (K). British Isles, England | 10 | 4 |
| Davis 55121 (BM). Ifrane, Cascada, |  |  | Turrill 4903 (K). Leicestershire, Charwood |  |  |
| 1580 m | 9 | 8 | Forest | 9/10 | 4 |
|  |  |  | Ap. ys 77-390 (K). Kent | 10 | 4 |
|  | 8 |  | Turrill (K). Surrey, Box Hill [England SE] | 10 | 4 |
| Font Quer 1928 (BM).Kaloa to Tauka, 150 | 8 | -9 | Fraser (K). Surrey, Box H ill | 10 | 4 |
| PORTUGAL |  |  | Fraser (K). Surrey, Chalk Pits | 10 | 4 |
| Goncalves 4625 (BM). Azores | 12 | 4 | Bean \& Hill (K). Scotland: Neopath Castle, |  |  |
| Yoller 61 (BM). Sierra Jerez [Spain]? | 12 | 3-5 | Tweedsdale near Peebles | 10 | 4 |
| Goncalves 4491 (BM). Azores | 10 | 4 | Ball 1838 (US). Surrey, Jumper Hill | 10 | 4-5 |
| ex Herb. Moniz (K). Madeira | 10 | 4 | Ex. Herb. Bidwell (BM), dovastonianum. |  |  |
| Cyrén (S: C-2058). Ser. Estrela | 10 | 6 | Westfelton | 10 | 4 |
| Fontee et al. (S: C-2047). Ser. Estrela, $1400 \text { m }$ | 10 | 5 | Ex. Herb. Gordon (K), dovatsonianum. Westfelton | 10 | 4 |
| Fontee et al. (S: C-2047-2). Ser. Estrela, | 9 | 5 |  |  |  |
| Meaden 1865 (K). Madeira | 9 | 4 | original) | 10 | 5 |
| Moller (BM). Serra Gerez: Vidoal | 8 | 4 | Lewis 721 (BM). Monmouthshire | 10 | 5 |
| Taxus canadensis |  |  | Jarrell (K). Kent: Shorehane | 10 | 5 |
| Lowe 570 (BM, bottom specimen). Madeira | 11 | 7-14 | Bennett \& Croydon 713 (US). Riddlesdown | 10 | 5-6 |


|  | SR | MC |  | SR | MC |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Hooker (PH). Kent | 9 | 3 | HUNGARY |  |  |
| Barron (K). Kent, Buckland | 9 | 4 | Herb. Láng (PH) | 10 | 4 |
| ? (K). Kent, "var. washingtonianum" | 9 | 4 | Schönach 3084 (S: C-2061A) | 10 | 4 |
| Valpy (K). Elsing, Norfolk | 9 | 4 | Schönach 3084, Austr-Hungar. (US). 445 m | 8 | 3 |
| Boswell (BM). Shropshire, Lyth Hill | 9 | 4 | Wagriesh (US: 451917). Vorarlbergia, 445 m | 8 | 3 |
| Brubaker 1960 (PH). Druids Grove | 9 | 4 | Wagriesh (US: 481917). Dolüa | 8 | 3 |
| Roper 1525 (K). Bristol, Birdhamdown | 9 | 4 | Boros (BM). Comit. Boraod. Ohassa, 550 m | 9 | 5 |
| Carruthers (K). Ireland: Pollawaddy | 9 | 4 | Lémke (S: C-2042). Bakony: Mikl¢spalhazy | 9 | 4-5 |
| Aug. 1874 (BM). Perth Co.? [filed under Portugal] | 9 | 5 | Schönach, Aust.-Hungar. (US: 966290), epacroides 445 m | 8 | 4-5 |
| Turrill (K). Yorkshire: 3 mi. from Richmond | 8 | 4 | Ex Herb. Mus. Nat. Hungar. (S: C-2041), Bakony | 8 | 4 |
| Hubbard (K). Sussex: Bury Hill | 8 | 4 | Schönach 3084, (BM). 445 m | 9 | 6 |
| Jackson (BM). Highclere, Saddam | 8 | 4 | Taxus canadensis |  |  |
| Turrill (K). Scotland: Loch Lomond | 8 | 4-7 | Schönach 3084 (S: C-2061 R Specimen). |  |  |
| Taxus canadensis var. adpressa |  |  | $445 \mathrm{~m}$ <br> ROMANIA | 5-6 | 19 |
| Summerhayes 2581 (K). E Kent | 8 | 4-6 | ROMANIA |  |  |
| SWITZERLAND |  |  | Topa, Bot. Mus. Exsic. (US). Bucovina: |  |  |
| Kellermann (US: 518500) | 10 | 3 | $400 \text { m }$ | 12 | 16 |
| Herb. A. Gray (K) | 9 | 5 | Topa, Bot. Mus. Exsic. (S: C-2024). Bucovina, 400 m |  | 3-5 |
| Fr. Castella (US). Le Pissot sur alboue, 1000 m | 7 | 6-7 |  | 10 |  |
| GERMANY |  |  | Topa, Bot. Mus. Exsic. (US). Bucovina, 400 m 9 | 4 |  |
| Reichenbach fil. (PH). Dresden | 11 | 3 | Anderson 102 (K). Balkan Exped., Cajan Pass | 9 | 4 |
| Martius 1831 (PH). Bavaria | 9-11 | 4-6 | Mititleu \& Barabas (BM). Bucovina: |  |  |
| Martius 1831 (K). | 9/10 | 4 | Darmanesti, 500 m | 8 | 5 |
| Milchbuder (K). Bavaria | 10 | 4 | BULGARIA |  |  |
| Petzi 1444 (K).Bavaria | 8/9 | 5 | Kotschy (P) | 8 | 4 |
| Reichenbach, ex Short Herb. (PH). Dresden | 7-8 | 4 | Anderson 42 (K). Sofia: Vitorha | 9 | 4 |
| Keller (PH). Darmstadt | 7 | 4 | ITALY |  |  |
| Martius 1831 (K). Bavaria Alps | 7 | 4 | Herb. Hook. 1814 (K). Montagnes | 10 | 3/4 |
| POLAND |  |  | Levier (BM). Florentino | 10 | 4 |
| Baenitz (US). Silesia: Proskau, 180 m, "f. dovastonii" | 10 | 6 | Lenander 1933 (S: C-2008). Lago di Garda, |  |  |
| Baenitz (US) epacroides. Silesia: Breslau, |  |  | McDonald: I-37 (US). Cult. | 10 | 5 |
| 120 m, "v. recurvata" | 9 | 4 | McDonald: I-37 (PH). Cult. | 8 | 5 |
|  |  |  | Solla (US: 280040). | 9 | 3-4 |
| "f. epacroides" | 10 | 4-5 | Baroncini 16 Sep 1893 (US)8/9 | 5 |  |
| Baenitz (US), epacroides. Silesiaca: Breslau, |  |  | Fireuze (BH). Cult. | 10 | 9 |
| Scheitniger Park 120 m | 8 | 4 | Martelli (PH). "Iter Sardoum", Limabara |  |  |
| Baenitz (US). Silesia: Breslau, 120 m, "f. erecta" | 9 | 9-10 | [Sardinia] | 8/9 | 10 |
| CZECH REPUBLIC |  |  | [YUGOSLAVIA] |  |  |
| Jirasek \& Suza (K). Moravia Centr.: 4-450 m | 10 | 4-6 | Biol. Inst. Dubrovnik 37 (NA). Bosnia: |  |  |
| Jirasek \& Suza (US). Moravia Centr.: 4-450 m | 10 | 5-6 | Mt. Trebevic near Sarajevo, 1450 m Kosarim (S: C-2065), Macedonia, Petiska | 12/13 | 3-4 |
| Petrakm, Fl. Boeh. \& Morav. exsic. 99 (BM) | 8 | 4-5 |  | 10 | 5 |
| AUSTRIA |  |  | Baldacci 169 (K). Albania <br> Rohleana 1908 (BM: 17197). Montenegro | 8/9 | 5 |
| Ex Pickler Herb. 1895 (US). | 11 | 4 |  | 7/8 | 4-5 |
| Ex Herb. Pichler (US: 347988, lower |  |  | Curic 1897 (K). Bosnia |  |  |
| specimen). Tirol | 10 | 4 | Biol. Inst. Dubrovnik 136 (NA). Bosnia. |  |  |
| Ex Herb. Pichler (US: 347988).Tirol | 8 | 4 | Mt. Plasma near Jablanica, 1500 m | 6/7 | 2-4 |
| Ex Shulte Herb. 1863 (K) | 10 | 4 | Woloszczak (K), Tatra | 9 | 6-9 |
| Hayer (S: C-2034). Salzburg | 10 | 6 | Taxus canadensis |  |  |
| Gander 1869 (K). Tirol | 9 | 6 | Berglund (S: C-2066; var. adpressa). |  |  |
| Gander 1869 (US: 157025). Tirol | 9 | 6 | Slovenien: Bled, berget Straza | 4-6 | 24 |
| Keck (US) | 9 | 5 | GREECE |  |  |
| Taxus canadensis |  |  | Georginda \& Tzanoudakis 631 (BM). Artis: |  |  |
|  |  |  | Mt. Tzoumarka | 10 | 6 |
| Kulmburg | 9 | 17 | HGT884 (K). Hills N of Xant Is | 9 | 4 |


|  | SR | MC |  | SR | MC |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Heldreich (S: C-2023). Oeta, 4500-6000 ft. | 9 | 2-4 | Brzhezitzky \& Kasumov H196 (US). |  |  |
| Greola (PH). "Mts. of Tyrah" | 9 | 4 | Azerbaijan | 10 | 15 |
| Guiol 2260 (BM). Mt. Olympus | 7/8 | 3-4 |  |  |  |
| Handel Mazzetti (K). Mt. Olympus, 750-850 m | 8-9 | 8 | Kousnetzoff 89 (US: 254512). [Russian |  |  |
| TURKEY |  |  | Federation] Kuban |  |  |
| Balonsa (BM). Taurus | 9 | 5 | NORWAY |  |  |
| Balonsa (P). Taurus | 8 | 5 | Gamble 28933 (K). West Dalen | 10 | 4 |
| Davis \& Hedge 32208 (BM). Coruh, Savval Tepe above Murgul, 1400 m | 7 | 6 | Taxus canadensis Anderson (US: 1091452). Kolsås | 5 | 18 |
| Sintensis 51181892 (P) Paphalogonia: Wilajet Kastanbuli | 9 | 8 | DENMARK <br> Herb. Joh. Lange 1866 (K) | 10/11 | _ |
| Sintensis 5118 (K) | 7 | 8 | Herb. Joh. Lange 1866 (K: Right specimen) | 8 | 4 |
| Murray 936 (NA). Between Molla Veyis and Meyden, S of Ardesen, 750 m | 8 | - | SWEDEN Thedenius (PH), Göteborg | 9-10 | 4 |
| Davis 13667 (K). VA. Jenigli (Caira [Caria ?]) |  |  | Thedenius (US) | 8 | 4-6 |
| (Denizli, Boz Da, Davis 13447), |  |  | Steinvall 1872 (K). Södermanland | 10 | 6 |
| 5000-5500 ft | 8 | 8-10 | Henriksson (K). Dalsland:Gunnarans | 8/9 | 6 |
| IRAN |  |  | Holmgren (US: 1276222). Blekinge | 8 | 5 |
| Koelz 16208 (US, distinct for obconical to 4-lobate seeds). Gozlu, Mazandaran | 8-9 | 7-10 | Lindberg 419 (K). Ekerö | 11 | 8 |
| SYRIA |  |  | "washingtonianum" | 10 | 8 |
| Haradjían (K). Dúldúl: Mt. Amanos, 5000-7000 ft | 10 | 4 | Bjornstrorn, ex Mus. Stockholm (US). <br> Podermanlane? [Södermanland] | 10/11 | 10 |
| Haradjían 2341 (S). Dúldúl: Mt. Amanos, 5000-7000 ft | 8 | 4 | Taxus canadensis |  |  |
| Gesbeldagh (BM) | 9 | 4 | Asplund (US: long-needled specimen). |  |  |
| Gesbeldagh (US) | 10 | 4 | Södermanland | 9 | 20 |
| Haradjían 3865 (S). Dúldúl: Mt. Amanos, 1500-2000 m | 10 | 8-10 | Asplund (Us: short-needled specimen). <br> FINLAND | 8 | 22 |
| Delbés (P). 1000 m | 9 | 8 | Florström (BM). Alandia: Lemland | 11/12 | 4 |
| RUSSIAN REGION |  |  | Florström 1909 (K) | 10 | 3-6 |
| Estonia |  |  | Vidlund, Helsinki Exsic. (K). Lemland | 8 | 4 |
| Taxus canadensis |  |  | EAST TEMPERATE ASIA |  |  |
| Lundström 742 (S). Eosl near Karriland | 9 | 16 |  |  |  |
| Lundström 579 (S). Ösel, Sworbe | 9 | 22 | Cuspidata Alliance |  |  |
| Lundström 562 (S). Ösel, Sworbe | 7 | 15 | RUSSIAN FEDERATION |  |  |
| Transcaucasia (Caucasus Mts.) |  |  | Taxus biternata |  |  |
| Elias et al. 5615 (NA) Ukraine Crimea, 1150 m | 9 | 9 | B. Cerereu (A). Far East Region: Pryanyk For Div. | 11 | 12 |
| Princeps Kascelsky, ex. Herb. Hort. Imper. Petro. (S). Caucayas Hosharia | 15 | 6-7 | Taxus umbraculifera Comp |  |  |
| Busch (K). Caucasus, Terek | 11 | 12-14 | Kypehisnova (A). Primorye Prov. | 14 | - |
| Woronowa (S: C-2027). W Transcaucasia: Suchum, Petkir (FI. Madshara) | 13 | 1 | Kypehinova (A) Primorye Prov., Bay of Peter, First Sea Reserve, Is. Stenin | 10 | 5 |
| Dmitrieva (NA). SW Georgia, Black Sea near |  |  | Dvorakovskia \& Bokina (A). Sakhalin Is. | 11 | 8 |
| NE Turkey, Adzharia, 900 m | 11 | 4 | Palczevsky 3601 (K). Primorye Prov.: vicinity |  |  |
| Inst. Bot. Acad. Sci. Armenia (US). 22 Mar 1946 | 10 | 4 | of Vladivostok | 11 | 8 |
| Szovich 610 (S: C-2072). Armenia: |  |  | Palczevsky 3601 (US) | 9 | - |
| Tschunakuchi | 10 | 4 | Palczevsky 3601 (A) | 8 | - |
| Szovich 610 (P). Armenia: Tschunakuchi | 9 | 4 | Lyubarsky 2 (A). South, Sikhote-Alin, foothills, |  |  |
| Ex. Herb. Inst.. Bot. Ac. Sc. URSS (US: 2560106). |  |  | Mt. Hezalaza, River Beryozovoy | 8 | 7 |
| [Transcaucasia] | 9 | 4 | Esus 203 (K). Sakhalin Is. | 8 | 12-15 |
| In Russian \#75 (P). Azerbaijan [Kura Mts.] | 9 | 8 | MANCHURIA |  |  |
| Herb. Komaróv (US: 1862552). Armenia | 7/8 | 6 |  |  |  |
| Prilipko (K). Transcaucasia | 9 | 10 | Taxus biternata |  |  |
| Goghika (NA). Caucasas: Azerbaijan, Chanlar, 1800 m |  | 8 | Ex herb. hort. bot. Petro. 1860, Maximowicz Mandshuria SE | 10 | 6 |


|  | SR | MC |
| :---: | :---: | :---: |
| Skvortzov 20 Sep 1931 (A). N Manchuria, |  |  |
| Sochintzest, forest, tree | 10-11 | 8 |
| In Russian No. 75 (P). Jilin (Kirin) | 9 | 8 |
| C. H. Chen 539 (A). Jilin (Kirin) | 8 | 6 |
| Maack 1855. (GH: top specimen) | 7 | 7 |
| Purdom (GH). N China [Shaanxi: Tai-pei-shan |  |  |
| T. umbraculifera Complex |  |  |
| Palczevski [Komaróv] 88 (K). Manchuria: |  |  |
| Palczevski [Komaróv] 88 (BM) | 12 | 9 |
| Maack 1855. (P) | 11 | 8 |
| Ex herb. hort. bot Petro. 1860 (Bunge), |  |  |
| Maximowicz (P). Mandshuria SE | 12 | 10 |
| Ex herb. hort. bot. Petro. 1860, Maximowicz (S) |  |  |
| Mandshuria SE | 12 | 8 |
| Ex herb. hort. I.c. (P) | 10 | 8 |
| Ex herb. I.c.(GH) | 7 | - |
| G. Fenzel (A). Schenhsi merid., Taipei-schan | 7 | 13 |
| KOREA |  |  |

## Taxus biternata

Wilson 10519 (US). Kyongsan, Nemon-rei 1315
Wilson 10519 (A) 1014
Wilson 9484 (A), Hallai-san down to
Mushroom House, bush

Wilson 9097 (A), Shinkabachin Heizanchien
to Ehoshin, Kankyo-N Heian divide, tree 1116
Wilson 8685 (A). N. Heian Prov.: O.G.M. Co.
Wilson 8685 (US) - 10-11 12

Wilson 8685 (K) 1010
Komaróv 1897 (GH). Pen-nian Prov. 10 4-6
Wilson 10688 (US). Kyongsan, Nemon-rei
Wilson 10688 (A)
9-10 6

Wilson $10688(\mathrm{~K}) \quad 7 \quad 6$

## Taxus umbraculifera Complex

Wilson 8538 (US). Oagelet Island, 0-900 m,
bush $12 \quad 12$

Wilson 8538 (A) $12 \quad 12$
$\begin{array}{lll}\text { Wilson } 9332 \text { (A). Kyongsan Prov.: Nemon-rei, } & & \\ \text { tree } & 12 & 10 \\ \text { Faurie } 1512 \text { (BM) } & 9 / 10 & 9\end{array}$
JAPAN

| Taxus cuspidata |  |  |
| :--- | :--- | :--- | :--- |
| Ex Herb. Zuccarini (M: type, T. cuspidata) |  |  |
| $\quad$ Japan |  |  |
| Ex Herb. Zuccarini (K: type, T. cuspidata) | $11-12$ | 24 |
| $\quad$ Japan | $11-12$ | 18 |
| Jack (A). Hokkaido: Sapporo | 12 | 20 |
| Jack (GH). Hokkaido: Sapporo | $10-11$ | 16 |

## Taxus biternata

Makino 43775 (S), Tokyo Pref.: Oizuni,

Nepymawku
Mujabe1884 (A). Hokkaido: Ishikasi
$16 \quad 12$
$14 \quad 14$

|  | SR | MC |
| :---: | :---: | :---: |
| Suzuki 499003 (A), Honshu: Mt. Ooyhama, Kanagawa-Pr, cult. | 14 | 13 |
| Faurie Dec 1904 (A), [Hokkaido], cult. and in forest | 13 | 13 |
| Muroi 1969 (A). Honshu: Mt. Fujiwara | 12-13 | 14 |
| Wilson, ex. Sakurai (A), Honshu: Kyaraboken, cult. "nana" | 10 | 13 |
| Muroi 5933 (A). Honshu: Mt. Himekami | 10-12 | 10 |
| Sapporo Agric. College (PH). Hokkaido: Kitami |  |  |
| Prov, Rishiri | 11 | 11 |
| Sapporo Agric. College1885 (A). Hokkaido: Niarenai? | 9-10 | 13 |
| Sapporo Agric. College1878 (A). Hokkaido | 11-13 | 11 |
| Naito (A), ex. Herb. Kagoshima Univ. Shimane Pref., Mt. Sentsu-zan. | 11 | 8 |
| Hatusima 13858 (A). Kagoshima Pref., Mt. Takahuma, tree | 11 | 8 |
| Shiota 4441 (A). Hondo, Mino Prov., hort. | 11 | 8 |
| Muroi 30 (A), Honshu: Hyogo Pref., Mt. Hyonosen | 10 | 8 |

## Taxus umbraculifera var. hicksii

Muroi 5603 (A). Hyogo Pref.: Kumatugi, Mikata-gun 1314
Muroi 5424 (A). Mt. Hatibuse 128
Muroi 5648 (A). Wakasugi 11-12 10
Muroi 3593 (A). Iwate-Pref.: Asagishi 11-12 10
Muroi 3698 (A). Gifu-Pref:: Takayama 1110
Muroi 3715 (A). Nagano Pref.: Kamikochi 118

## ASIA: HIMALAYAS <br> Baccata Group <br> AFGHANISTAN-INDIA

## Taxus contorta var. contorta

| Sprague 730 (K). Murree | 6 | 4-5 |
| :---: | :---: | :---: |
| Aitchinson (K). Kurrum Valley, 7500-9000 ft | 6 | 4 |
| Sinnott et al. 146 (K). Between Gotchbok and Kubkot Valley, 2750 m | 7 |  |
| Stewart 15343 (US). Murree, 7000 ft | 7 | - |
| ex Herb. Schlagintweit (PH). NW of Srinagar | 5 | - |
| Stewart 7374 (PH). Sonamarg, 10,000 ft | 6/7 | 4 |
| Mukinji (K). Lada Valley | 6 | 4 |
| Stewart 8414 (US). Kashmir: Pahlgam | 5 | 4 |
| Stewart 8414 (A). Kashmir: Pahlgam | 7 | 3 |
| Stewart 8414 (PH). Kashmir: Pahlgam | 7 | 3 |
| Stewart 12001B (A). Kashmir: Pahlgam 2600 m |  | 6 |
| Schlagintweit (P). Kashmir: Báltal to Númner | 7 | - |
| Kenyoer \& Dugeon (PH). Bureah, 11,000 ft | 7 | 5 |
| Rau 31770 (A). Garhwal to Lake Hemkund, 3200 m | 7 | - |
| ex Herb. Falconer 1000 (S: C-1994). Kumaon, Dwali? 8500 ft | 7 | 4 |
| ex Herb. Falconer 1000 (P). Kumaon | 7 | 5 |
| Koelz 10285 (A). Punjab: Kulu, above Bandrole, 8000 ft | 7-8 | - |
| Schlagintweit 8941 (GH). Kashmir: Sukhi across Bamsuru and Chaia Pass to Khdrsali, 9000-15400 ft | 7-8 | 3 |


|  | SR | MC |
| :---: | :---: | :---: |
| Rodin 5313 (US). Punjab Province: Rosenhiem, |  |  |
| Murree | 8 | 5 |
| Stewart 5931 (A). Kashmir: Pahlgam, |  |  |
| Heybrook 29 (K): Kashmir: Pahlgam 2600 m | 8 | 4 |
| Lace 301 (A). Bashahr, Uri Forest | 8 | - |
| Stewart (PH: 829196). Dharmkat, Dharmsala, $6000 \text { (ft?) }$ | 8 | 8 |
| Gamble 23507 (K). Jaunsar Dist., 10,000 ft | 8 | 1-3 |
| Stewart 10663A (PH). Gulwarg, $7000-10,000 \mathrm{ft}$ | 9 | 12 |
| Laig Raus (P). Siwalik and Jaunsar Div., 10,000 ft | 9 | 5 |
| Pengelly (K). Chumba | 10 | 4 |
| Bertoloni (BOLO: type, T. orientalis). Western Sikkim | 10-11 | 3 |
| NEPAL |  |  |
| Baccata Group |  |  |
| Taxus contorta var. contorta |  |  |
| Stainton et al. 7832 (BM). Chingnon, 10,000 ft |  | 4 |
| Polunin et al. 1353 (BM). Dhotar, 9600 ft . | 9 | 4 |
| Polunin et al. 432 (BM). Chankeli Range, $8000 \mathrm{ft}$ | 11 | 5 |
| Polunin et al. 5050 (BM). W of Jumla, Belas |  |  |
| Gaejigeth, 10000 ft | 9 | 6 |
| Polunin et al. 1873 (BM). Chatlwe, 9000 ft . | 10 | 5 |
| Gardner 557 (BM). Shios Khola, 8500 ft | 9/10 | 5 |
| Stainton et al. 734 (BM). Lete, S of Tukucha, 8000 ft | 10 | 5 |
| Ottba et al. 8311066 (BM). Marayandi Khola | 11 | 0 |
| Mikage et al. 9550282 (BM). Dhaulagiri Zone, 2405 m | 10/11 | 4 |
| Stainton et al. 5616 (BM). Chingnon, $N$ of Tukucha, Gadaki Valley, 10,000 ft | 11 | 7 |
| Taxus contorta var. mucronata |  |  |
| Dobremez 2106 (BM). | 8/9 | 10 |
| Wraber 514 (BM). Hanangi: Karayundi Valley, |  |  |
| 3100 m | 8/9 | 10 |
| Sumatrana Group |  |  |
| Taxus sumatrana |  |  |
| Herb. Banerji, 1953, in adnot. T. bounoniana Carr. (A). E Nepal: Khanigaon to Kalanti, 6,000 ft. | 12 | 16 |
| Wallichiana Group |  |  |
| Taxus wallichiana |  |  |
| Wallich 6054A (M: Original Material). [Nepal] 15 |  |  |
| Wallich 6054A (K: Duplicate of Original |  |  |
| Material). [Nepal] | 14 | 5 |
| Wallich 6054A (K: Duplicate of Original |  |  |
| Wallich 6054A (S: Duplicate of Original |  |  |
| Material). [Nepal] | 13-15 | 2 |
| [Wallich] (GH: Duplicate of Original |  |  |
| Material). Napalia. | 15 | 4 |


Baccata Group
Taxus contorta var. mucronata
tal. 16035 (A). Eastern, Ha:
89.18, , 9,000 ft
tal. 16035 (BM)

## Sumatrana Group <br> Taxus celebica

Cooper \& Bulley 2833 (BM). Rinchu Timakha, 6000 ft
$13 \quad 13$

## Wallichiana Group Taxus wallichiana

Ludlow et al. 18672 (BM). Tunle La. near Kinga Rasdah, 11,000 ft
$12 \quad 2$
Ludlow et al. 18672 (A). Tunle La near Kinga Rapden, 11,000 ft
$12+5$
Grierson \& Long 4417 (A). Thimphu Dist.: summit of Dochong La, 3110 m
126

Cooper \& Bulley 2600 (BM). 7,500 ft 164
Bartholomew \& Boufford 3917 (A). Above Motithang, W of Thimphu

10-11 6

## NE INDIA \& TIBET

| Sumatrana Group |  |  |
| :---: | :---: | :---: |
| Taxus celebica |  |  |
| Kingdon Ward 19324 (BM). "Assam" [Tibet] Rima, 7000 ft | 11 | 32 |
| Clarke 38308 (K). Khasia: Maophlang | 9 | 32 |
| Taxus kingstonii |  |  |
| Mann 1885 (K). Khasia Hills, I.c. | 13 | 20 |
| Mann 1885 (BM), I.c. | 15 | 15 |
| Mann 1885 (P). Khasia Hills, I.c. | 13 | 19 |
| Kingdon Ward 18751 (A). Khasi Hills, Mawphlang, 6000 ft | 12 | 15 |
| Hooker 1337 (K). Khasia: 5000 ft | 13 | 16 |


|  | SR | MC |
| :--- | :--- | :--- |
| Hooker \& Thomson 1855 (P) | 11 | 12 |
| Hooker \& Thomson 1855 (P) | $10 / 11$ | $8-10$ |
| Hooker \& Thomson 1855 (P with seed) | 13 | 10 |
| Simmons 484 (P). Assam: Khasia | 13 | 12 |

## Taxus sumatrana

Mann 1885 (A). Khasia Hills: Nunghuai, $5000 \mathrm{ft} \quad 24 \quad 12$

| Wallichiana Group |  |  |
| :---: | :---: | :---: |
| Taxus wallichiana |  |  |
| Wallich (M: Lectotype). Eastern | 13 | 4 |
| Biswas 439 (A). E Himalaya | 13 | 5 |
| Biswas 439 (A). E Himalaya | 10-11 | 5 |
| Kurz (A). Sikkim: Tongloo | 13-15 | - |
| Raijada 18919 (A). Cult., Dehra Dun, Bot. Gard. Darjeeling | 14 | 4 |
| Griffith 5002, ex Herb. Griffith. E Himal. (P) | 12 | 0 |
| Griffith 5002, ex Herb. EIndia Co (P) | 13 | 4 |
| Griffith 5002, ex Herb. Bunge E Himal. (P) | 13 | 4 |
| Hooker 77 (P). Khasia, 5000-6000 ft | 16 | 4 |
| Hooker 77 (P). Khasia, 5000-6000 ft | 16 | 4 |
| Griffith 2(7)606 Assam (P) | 15 | 4 |
| Kingdon Ward 17271 (A). Sirhoi: 8000 ft | 15 | 4 |
| Kingdon Ward 17271 (BM). Sirhoi: 8000 ft | 15 | 4 |
| Vos et al. 148 (NA). West Bengal: Singalila |  |  |
| Range, 8400 ft | 15 | 5 |
| C. B. Clarke 436743 (BM). Khasia: 4500 ft , Vale of rocks | 14 | 4 |
| G. Watt 5955 (A). Manipur: Seriphari, $10,000 \mathrm{ft}$ | 15-17 | - |
| G. Watt 5955 (P). Manipur: Seriphari, $10,000 \mathrm{ft}$ | 17 | 4 |
| G. Watt 6493 (P). Manipur: Sirohifarar, 7000 ft | 16 | 4 |
| G. Watt 6208 (P). Manipur: Jakpho, 11,000 ft | 18 | 4 |

## Taxus wallichiana var. yunnanensis

$\left.\begin{array}{llll}\text { Hooker (K). Sikkim: 7000-10,000 ft } & 14 & 3 \\ \text { Hooker (K). Sikkim: 7000-10,000 ft } & 14 & 3 \\ \text { Kingdon Ward } 18990 \text { (BM). Jakpho Range }\end{array}\right)$

## Sumatrana Group (Taxus kingstonii)

Oliver 4 Sep 1894 (K). Bernardmyo, Ruby
Mines 158

Oliver (K) 14 May 1892,5600 ft 12 5-6

|  | SR | MC |
| :--- | :--- | :--- | :--- |
| Wallichiana Group |  |  |
| Taxus obscura (Chinensis Subgroup) |  |  |

## Taxus wallichiana var. wallichiana

Kingdon Ward 9214 (BM) Northern, Adung Valley, $97^{\circ} 30-98^{\circ} 30^{\prime}, 27^{\circ} 30-28^{\circ} 30^{\prime}, 6000$ ft. 174
Kingdon Ward 9214 (A) 164

Kingdon Ward 9375 (A). N Adung Valley,


12-13 4
Taxus wallichiana var. yunnanensis
Kingdon Ward 22819 (BM). Mt. Viatoria, 9000-10,000 ft 164
Kernode 17205 (K). Myintkyina: LaikanFenshuiling Rd, 8000 ft

123
THAILAND

| Chinensis Subgroup |
| :--- |
| Taxus obscura |
| Lobb 461 (BM) Malaya |

## Sumatrana Group (T. sumatrana)

| Kerr 20146 (K). Kao Kuading, 1200 m | 15 | 14 |
| :--- | :--- | :--- |
| Kerr 20146 (BM) | 12 | 14 |

VIETNAM

| Sumatrana Group |  |  |
| :---: | :---: | :---: |
| Evrard 305 (P). Dalat: ravin buisé an chalet |  |  |
| Rimaud | 9/10 | 24 |
| Evrard 1438 (P). Lâm Dông | 12 | 24 |
| Schmind 1960 (P). Dak Tria- Manline, 1400 m | 8 | 32 |
| Schmind 1960 (P). Dalat: Dau Lamghi | 9 | 24 |
| Schmind (P). Dalat: Dak Tria, 1610 m | 11 | 32 |
| Van Cuong 12891960 (P). Dalat: Manline, 1610 m | 12 | 23 |
| Wallichiana Group |  |  |
| Taxus chinensis |  |  |
| Hiép \& Chan 405 (P). Hoa Binh, Mai Chôu, Pà Co, 900-1500 m | 13 | 8-13 |
| Taxus aff. chinensis |  |  |
| Poilane 4150 (P). Phu Khanh: Nha Trang, |  |  |
| 1500 m | 11 | 4 |
| Poilane 4150 (A). Nha Trang, 1500 m | 10-11 | 4 |


|  | SR | MC |
| :---: | :---: | :---: |
| Schmind (P). Dalat: Dak Tria, 1400 m | 15 | 8 |
| Soulie 1523 (P). "Haut Mekong" | 15 | 12 |
| CHINA |  |  |
| Tibet \& Yunnan |  |  |
| Sumatrana Group |  |  |
| Taxus kingstonii |  |  |
| Soulie 1411 (P). "Tackou et Nekou ("Haut Mekong" ) | 11 | 12 |
| Wallichiana Group |  |  |
| Taxus florinii |  |  |
| R.C. Ching 21505 (A). Soc. W. Sikiang: Tamichung | 10 | - |
| C. W. Wang 65475 (A). Sikang, Me-kong, Tsa-wa rung, 2500 m | 8-9 | 16 |
| Handel-Mazzetti 2602 (K). Ngaitschekou, $2800-3500 \mathrm{~m}$ | 11 | 2 |
| Fleigner et al. 1129 (K). Sahlie Valley on Muzhiyan Shan, 2980 m | 10 | 10 |

## Taxus wallichiana var. yunnanensis

Zhang 916 (PE: type). Tibet, Zayul, 2100 m 153
Kingdon Ward 6292 (BM). Zayul, 7000-8000 ft 153
Sichuan

| Sumatrana Group <br> Taxus celebica |  |  |
| :--- | :--- | :--- |
| H. Smith 10401 (BM). Huangnipu, Malingtsang, |  |  |
| 1000 m | 12 | 36 |
| Wang 20541 (A). South of Kuan-Hsien, |  |  |
| 1160 m | $11-12$ | 36 |
| Farges $1895-1897$ (P). Tchenkéou Tin | 20 | 32 |
| Farges 128 (P). Tchenkéou Tin | 14 | 24 |
| $\quad$ Taxus kingstonii |  |  |
| Cheng 1001 (BM). Tachienlu | 12 | 18 |
| Cheng 1475 (P). Tachienlu | 12 | 18 |

## Taxus mairei

Wilson 1265 (A). Western: Nin Ya-chou Fu,
$2000 \mathrm{ft} \quad 21 \quad 25$
Wilson 1265 (US) 1516
Fang 5811 (P). Nanchuan-Hsien 1712
Fang 5811 (A). Nanchuan-Hsien - 16
Hwa 229 (K). Metasequoia area 1612
Fan \& Class 91 (A). Kuan-Hsien,
Chien-Chang-Shan, 1000 m

Farges 1436 (P). NE 1615
Farges 100 (P) 1516
Law 65 (K). Pei pah 15 8-12
Hwa 27 (A). Li-chuan, Jian-Nan-Hsien, Ta-pen-Ying, 3800 ft Huangnipu and Yaan (Yachou), Malingtsang, $900 \mathrm{~m} \quad 1414$
Smith 10402 (S) $15 \quad 27$
Hwa 27 (A) $14 \quad 21$
$\begin{array}{ll}\text { Hwa } 27(\mathrm{~K}) & 14 \quad 17\end{array}$

|  | SR | MC |  | SR | MC |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Legendre 586 (P). Pao Shan NE, 600 m | 14 | 17 | Chiao \& Fan 464 (A) | 10-11 | 7 |
| Fang 3461 (A). Tienchuan Hsien, |  |  | Chiao \& Fan 464 (P) | 12 | 8 |
| Tienchuanchow, 2500-3000 ft | 14 | 20 | Chiao \& Fan 464 (US) | 13 | 9 |
| Fang 3461 (P) | 13 | 17 | Hu 8176 (A) | 14 | 9 |
| Fang 12205 (A). Kuan-Hsien, Mt. Tsing-cheng, Chengtu and Kuan-Hsien, 1390 m | 14 | 13 | Wang 20993 (A). W of Wen-chuan Hsien, 2800 m | 11 | 9 |
| Wang 20600 (A). Wah-Hsien, Mou-tao-chi, Metasequoia area, 1390 m |  |  | Hu 8497 (A) | - | 10 |
|  | 12-14 | 16 | Fang 18310 (A) | 13 | 10 |
| Hu 1563 (A). Shikong:Tien-Chuan Ling-Kwan3000 ft |  |  | Yu 8166 (A). 2400 m | 12 | 10 |
|  | 14-16 | - | Fang 15128 (A) | 16 | 11 |
| Farges 128 (P). Tchenkéou Tin | 14 | 17 | Fang 15128 (A) | 14 | 11 |
| Farges 128 (P). Tchenkéou Tin | 12 | 24 | Wilson 624 (K). S. Wushan, ravine | 12 | 12 |
| Fang 3442 (A). Tienchuan-Hsien, <br> Tienchuanchow, 2500-3000 ft <br> Fang 3796 (A). UnqLing-Hsien, 5000 ft <br> Cao 0152 (BM). Jiabigon, Zhao Quing-sheng, $2500 \mathrm{~m}$ | 11 | 12 | Taxus aff. chinensis |  |  |
|  | 12 | - | Cheng 2890 (A, Taxus OCR, in Spjut 200 W of Lung-an-fu | 13-15 | 8 |
|  | 12 | 10 | Cheng 2890 ( P ) <br> Hu 8619 (A). Emei-Hsien, Mt. Emei |  | 4 8 |
| Wallichiana Group |  |  | Taxus wallichiana |  |  |
| Taxus chinensis |  |  | Sichuan: Mt Emei |  |  |
| Harry Smith 10398 (BM). Tachsiangling, $2600 \text { m }$ | 16 | 6 | Hu 8166 (A) |  | 4 |
| Peng 502 (Biol. WCUU) (A) Yachow 1600 ft | 15 | 8 | Hu 8542 (A) | 15-16 | 4 |
| d'Legendre (P). 2500 m | 14 | 5 | Feng 3945 (A) | 14 | 8 |
| Henry 7155 (US: type): E Sichuan: Wushan-Hsien, 2000-3000 m |  |  | Lee 4465 (A) | 13 | 8 |
| Wushan-Hsien, 2000-3000 m | 13 | 4 | Wilson 4053 (A). W Pan-lan-shan W of Kuan |  |  |
| Henry 7097 (US) | 14 | 5 | Hsien, 5000-6000 ft | 14 | 4 |
| Henry 7097 (A) | 12 | 7 | Gansu (Kansu) |  |  |
| Farges 128. NE Sichuan, Tschen-kuu-tin Dist.: |  |  | Sumatrana Group |  |  |
| Farges 128 (P) | 16 | 8 | Taxus kingstonii |  |  |
| Farges 128 (P) | 11 | 7 | Meyer 1790 (P) | 14 | 8-12 |
| Wang 1930 (A) | 12 | 4 | Ningxia Huizu |  |  |
| Wang 22602 (A). Kwang-yun Hsien, 1800 m | 12 | 4 |  |  |  |
| Sichuan (Emei-Hsien: Mt. Emei) |  |  | Taxus celebica |  |  |
| (T. chinensis, by increasing number of bare | rginal | cells) | Chao 1223 (A). Sikong: Lung Dung An, |  |  |
| Feng 1941 (A) | 16-17 | 4 | 1000 m | 13 | 36 |
| Fang 16082. (A) | 16-17 | 4 | Shaanxi (Shensi) |  |  |
| Yu 667 (A). Mt. slope, 2600 m | 15 | 4 | See also Cuspidata Alliance, T. biternata, Pur | m s.n. |  |
| Wilson 6200 | 14-15 | 4 | Sumatrana Group (T. kings | nii) |  |
| Lee 3237. (A) | 14 | 4 | Davis 1872 (P). Tsin-lin au Lao-lin, 3000 m | 13 | 12 |
| Hu 8243. (US) | 13 | 4 | Davis 1872 (P). ThinlinauLaoli,3000 |  |  |
| Hu 8243. (A) | 14 | 5 | Wallichiana Group |  |  |
| Lee 4445 (A) | 13 | 4 | Taxus chinensis |  |  |
| Yu 669 (A). 1000 m | 12 | 4 | Chens 1893 (P). Central | 15 | 10 |
| Hu 8786. (A) | 14 | 5 | Yunnan |  |  |
| Yu 869 (A). 2500 m | 12 | 5 |  |  |  |
| Fang 18420. (A). 2335 m | 12-13 | 6 | Sumatrana Group |  |  |
| Lee 4500 (A). | 21 | 7 | Taxus celebica |  |  |
| Ching 1676 (A). Siachu, 2600 ft | 16 | - | Forrest (A) | 11 | 27 |
| Ching 1676 (P). Siachu, 2600 ft [rectangular cells, papillose midrib in upper half] |  |  | Forrest 7798 Gaoligongshan (K) | 14 | 18 |
|  | 16 | 7 | Taxus kingstonii |  |  |
| Wang 23656 (A). 2000 m | 14 | 7 |  |  |  |
| Wilson 479 (A). | 12 | 7 | Forrest 11789 (BM). Shweli-Salween Divide, |  |  |
| Chiao \& Fan. 604 (A). 1000 m | 10-11 | 7 | Forrest 11788 (K) | 13 | 12 |
| Fang 10940 (A). 1200 m | 13 | 7 | Forrest 11789 (K) | 13 | 7 |
| Fang 15940 (A) | 13 | 8 | Forrest 15945 (K). Schweli-Salween Divide Forrest 15945 (BM). Schweli-Salween Divide | 13 12 | 16 12 |


|  | SR | MC |
| :---: | :---: | :---: |
| Forrest 9462 (K). Ma-Chang-Kai, valley, |  |  |
| $25^{\circ} 30^{\prime} \mathrm{N}, 8000 \mathrm{ft}$ | 10 | 14 |
| Forrest (A). Yunnan, no other data | 12 | 16 |
| Forrest (A). Yunnan, no other data | - | 14 |
| Forrest 12087 (S). Schweli-Salween Divide | 15 | 12-14 |
| Forrest 12087 (K) | 13-14 | 14-20 |
| Forrest 9339 (BM) | 13 | 16-18 |
| Rankin 1913 (K). "Yung Chun" | 12 | 14 |
| Rock 7587 (US). Salween E of Tengyueh, to summit of Shweli, Shweli River | 12 | 9 |
| Taxus mairei |  |  |
| Forrest 15053 (K) | 16 | 12 |
| Maire 131 (BM) | 16 | 15 |
| Maire 1913 (P). Tie'tchang Keol, 700 m | 14 | 28 |
| Maire. (A: type). Dongchuan, 700-800 m | 13 | 17 |
| Wallichiana Group Taxus chinensis |  |  |

Feng 630 (A). Ta-hon-shan near Ta-koo, NE of Likiang Snow Range

12 -
Cavalerie 7823 (K)
Taxus florinii
Alpine Gard. Soc. Exped. 309 (K). Zhongdian; haba Shan, 3347 m
Rock 18502 (A). NW: Mt Ludu, NW of Li-Kiang,
W of Yangtze 84
Rock 18502 (US) 9-10 4
Forrest 19967 (S). NW: Mekong-Salween Divide 8-9 4
Schneider 2918 (A). 3000-3200 m 7 -
Schneider 2918 (K)
Schneider 1429 (A). 3500 m
9-10 5-6

Schneider 1429 (K) 78
Yu 11076 (A). sine locality 7-8 9
Yu 7848 (BM); Dokerla, 3100 m 8
Rock 11573 (A). Litiping Range, MekongYangtze divide, E of Weihsi 10-11 -
Rock 11573 (A) 9 -

Ching 21980 (A). Litiping, between Likiang and Weihai

10 -
Feng 1809 (A). S Chungtien, Kung- shiang-shu, Snow Mt to Kai-Lou-wei, Yangtze bank 3200 m

816
Wang 67735 (A). Wei-si Hsien, 2500 m 10-11 -
Wang 67414. (A). Lung-pan la Champu fung 10-12 5

## Taxus aff. chinensis

Feng 11937 (A: type in adnot., Taxus OCR in Spjut 2007). Si-chour-Hsien, Faa-doou, 1500 m

16
Feng 12105 (A) l.c. 14 -
Tsai 59874 (A, T. phytonii). Wei-se Hsien 2800 m

124
Tsai 58464 (A: type in adnot, Taxus SCU in Spjut 2007) Che-tse-lo, 3200 m 14
Tsai 58464 (P) 126

|  | SR | M |
| :---: | :---: | :---: |
| Taxus wallichiana var. yunnanensis |  |  |
| SB 1981 Exped., Cangshan 0419 (K). W |  |  |
| Shangschang, above Yangbi, 2700 m | 17 | 2 |
| Wang 67412 (A). Champu, 2120 m | 19 | 4 |
| 1984 SAB Exped 388 (A). Xangbi Xian, W side of Diancang Shan Mt Range, Malultang, Chang Shan, 2700 m, $25^{\circ} 46^{\prime} 100^{\circ} 01^{\prime}$ | 19 | 6 |
| Handel-Mazzetti 6408 (A). Dji-shan ad boreo- <br> orientem urbis Dali (Talifu) 3200 m 184 |  |  |
| SB 1981 Exped., Cangshan 0419 (A). W Shangschang, above Yangbi, 2700 m | 17 | 0 |
| SB 1981 Exped., Cangshan 0227 (A). |  |  |
| Kiemiu-ingdi above Yangbi, 3000 m | 16 | 4 |
| Wang 72417 (A). Chen-Kang Hsien | 15 | 4 |
| Yu 21036 (A). Salween, Kiukiang Divide, Shawlongwang, 2600 m | 14 | 4 |

## Guizhou (Kweichow)

## Sumatrana Group <br> Taxus mairei

Steward et al. 328 (A). Ta Ho Yen, Kianakou Hsien, 980 m 1420
Steward et al. 328 (US) 1412
Steward et al. 328 (P) 1318
Cheng 7525 (A: type, T. speciosa Florin).
Kiangkow, 450 m 1624

Steward et al. 154 (US). Liang Feng Yah,
Tsunyi Hsien, 900 m
Steward et al. 154 (A) $12 \quad 18$
Tsiang 8987 (P). Pichish 1413
Tsiang 8987 (A). Pichish 11-13 -
SAG Exped. 1981 (GH). Songtao Xian,
Lengjiaba, Xiaohe and Dahe Rivers, NE Fanjing Shan mt range, 820-1120 m 1218

## Wallichiana Group

Taxus chinensis

| Cavalerie \& Foriupat 2604 (P) | 20 | 7 |
| :--- | :--- | :--- |
| Cavalerie \& Foriupat 2604 (P) | 12 | 5 |
| Cavalerie \& Foriupat 2604 (P) | 11 | 5 |
| SAG Exped. 1981 (US). | 12 | 6 |

SAG Exped. 1981 (US). 126
SAGB 1986 Exped. 1854 (A). Yinjiang Xian, Xiapingsho, W Fanjing Shan range, 1 100-1400m 16-17 10
SAGB Exped. 1046 (A). Jiangkou Xian, Daiyenpeng, Kaitu River, SW Fanjing Shan range, 750-1000 m
Hubei (Hupeh)

## Wallichiana Group

Taxus chinensis
Chow 76099 (A). Shenlungkai
SA 1980 Exped. 1540 (A). S of Jiuhuping
Forest along Jizigou canyon, $1900 \mathrm{~m} \quad 136$

Wilson 1265b (A). Western: Nin Ya-chou Fu, 2000 ft

134

|  | SR | MC |
| :---: | :---: | :---: |
| SA 1980 Exped. 777 (A).Western: Shennongjia |  |  |
| For. Dist., NE Guanmenshan, S of Shicao river, 1150 m | 13 | 7 |
| SAB 1980 Exped. 1824 (GH). Shibapan, 1850 m |  | 7 |
| SAB 1980 Exped. 1824 (A) | 12 | 6 |
| Wilson 716 (A) | 12 | 7 |
| Taxus aff chinensis (Taxus sp. SCU in Spjut 2007) |  |  |
| SA 1980 Exped. 585 (A). Western: Shennongjia |  |  |
| For. Dist. $331^{\circ} 30^{\prime} \mathrm{N} 110^{\circ} 30^{\prime} \mathrm{E}, 1200-1400 \mathrm{~m}$ |  | 8 |
| Sumatrana Group |  |  |
| Taxus mairei |  |  |
| Gressitt 2507 (A). Metasequoia Area, between |  |  |
| Ta-yin-pin \& Chunglo, Shui-sa-pa, 900 m | 12 | 14 |
| Anhui (Anhwei) |  |  |
| Sumatrana Group |  |  |
| Taxus mairei |  |  |
| Ching 3168 (A). Southern, Chanen, 300 ft | 12-13 | 12 |
| Wallichiana Group |  |  |
| Taxus chinensis |  |  |
| R-C Ching 2622 (A). S Anhui, Clas Hara Shan | 17 | 5 |
| R-C Ching 2622 (US) | 13 | 4 |
| Cheng 4026 (BM). Wangshan | 11 | 6 |
| Henan (Hunan) |  |  |
| Taxus mairei |  |  |
| Fan \& Li 644 (A). Ma-Ling-Tung, Sinning Hsien, |  |  |
| 600 m | 15 | 18 |
| Fan \& Li 296 (A). Changning Hsien, Yang-Shan, |  |  |
| 680 m | 15 | - |
| Fan \& Li 296 (BM) | 14 | 12 |
| Jiangsu (Kiangsu, Kiangshi) |  |  |
| Taxus mairei |  |  |
| Wang-Te-Hui 445 (A). Ningdu, Yuntungtschi | 9-12 | 25 |
| Y.K. Hsiung 6443 (A). NW, Si-ho, Hwang- <br> kong-shan Mt |  |  |
| Wang-Te-Hui 458 (A). Lienhwa-shan, 800 m | - | 10-1 |
| Chow 80325 (BM). Nanking, 75 m | 12 | 16 |
| Guangxi (Kwangsi) |  |  |


| Sumatrana Group |  |  |
| :---: | :---: | :---: |
| Taxus celebica |  |  |
| Ching 5976 (US). Bin Long, Miu Shan, N Luchen, |  |  |
| border of Kweichow, 4000 ft | 15-16 | 32 |
| Ching 5976 (A) | 14-15 | 31 |
| Taxus mairei |  |  |
| Chiao 18795 (US). Lu Shan | 16 | 6-10 |

Wallichiana Group
Taxus chinensis

Steward \& Cheo 947 (P). 2110 m 154
Steward \& Cheo 947 (BM). San Chiang Hsien, $2110 \mathrm{~m} \quad 124$

## Zhejiang (Chekiang)

|  | SR | MC |
| :---: | :---: | :---: |
| Taxus mairei |  |  |
| Cheng 3617 (US). Eastern: Tien-Mu-Shang | 14-16 | 10 |
| Keng 317 (A). Taishun-Hsien | 13 | 18 |
| Hu 342 (A). Tien-Tai-Shan, 1300 m | 13 | 15 |
| S. Chen 1063 (A) | 12 | 18 |
| (US: 145110). Tien Tai Shan | 12 | - |
| Ching 2489 (A). S: King Yuan, 300-800 m | 11 | 20 |
| Ching 2489 (US) | 13 | 14 |
| Taxus sumatrana |  |  |
| Hu 1628 (A). Lin-an Hsien, 1200 ft | 14 | 16 |
| Hu 550 (A). Y-Chien Hsien, 1000 ft | 12 | 15 |
| Guangdong |  |  |
| Taxus celebica |  |  |
| Nanling Exped. 1838 (A). Ruyuan Xian | - | 31 |
| Taxus mairei |  |  |
| Tsang 20694 (US). Loh Ch'ang Dist., Chong |  |  |
| Uen Shan near Kau Fung | 13-15 | 14 |
| Tsang 20694 (A) |  | 24 |
| Chiao 14510 (A). Tien-Tai-Shan, 1300 m | 12 | 24 |
| Chiao 14510 (US) | 12 | 14 |
| Tsiang Ying 1425 (A). Hung-mio to Mio- lan, Jui-feng, Lokohong Hsien N.R. Region, 1340 m | 12 | 14 |
| Tsiang Ying 1425 (A, different label) | 14 | 24 |
| Tsiang Ying 1425 (P, specimen does not appear to be the same plant as in A) | 13 | 5-7 |
| Fujian (Fukien) |  |  |
| Sumatrana Group |  |  |
| Taxus mairei |  |  |
| Price 12586 (K). Ing-dan E. Fookma | 12 | 26 |
| Sheng 1544 (K). Naping, 800 m$16$ |  |  |
| Chung 2865 (A). Yeuping, Shih-Sun-Keng, |  |  |
| Chung 2865 (K) | 12 | 16 |
| David (P). W: Mts | 12 | 18 |
| Chung 3581 (A). Buong Kang, mt slope, 700 m | 9 | 24 |
| He-Guosheng 1544 (US). Naping, 800 m | 16 | 6 |
| Wallichiana Group |  |  |
| Taxus aff chinensis |  |  |
| H.H. Chung 3866 (A). Puchen | 11 | 5 |
| Taiwan |  |  |
| Sumatrana Group |  |  |
| Taxus kingstonii |  |  |
| Hsu 1651 (PH). Mt. Pasein-san, Taichang Hsien | 16 | 8 |
| Liu 389 (PH). Mt. Ammashan, Taichung Hsien | 15 | 8 |
| Liu 389 (A) | 14 | - |
| Hsu (PH). Mt. Pasan-shan, Taichang Hsien | 14 | 9 |
| Wilson 9738 (BM, isotype). Arisan Prov.: Kagi, |  |  |
| 2833 m | 13 | 9 |
| Wilson 9738 (A: holotype) | 10-11 | 8 |
| Liu 437 (US). Taiklang, Shih-wan-hsi, Pa-Hsien-shan, 2250 m | 12 | 9 |


|  | SR | MC |
| :--- | :--- | :--- |
| Nakahara (PH). Arizan Prov. | 11 | 9 |
| C-j Chang, Tongshi 6 (wba) |  |  |
| 27 Sep 93 | 14 | 9 |
| 26 Nov 93 | 13 | 7 |
| 06 Dec 93 | 12 | 9 |
| 09 Dec 93 | 13 | 9 |
| 27 Jan 94. | 13 | 8 |
| 13 Jan 94 (new growth) | 13 | 8 |
| 13 Jan 94 (old growth) | 11 | 9 |

## Taxus mairei

C-j Chang 1-2,4-5, 7-10 (wba). Hua-lien

| 116 May 94 | 12 | 18 |
| :--- | :--- | :--- |
| 103 Aug 94 | 12 | 18 |
| 216 May 94 | $11-12$ | 27 |
| 203 Aug 94 | $11-12$ | 20 |
| 416 May 94 | 12 | 21 |
| 403 Aug 94 | 12 | 17 |
| 516 May 94 | $10-11$ | 23 |
| 503 Aug 94 | 12 | 24 |
| 916 May 94 | 15 | 19 |
| 903 Aug 94 | 12 | 24 |
| 1016 May 94 | 12 | 14 |
| 1003 Aug 94 | 11 | 18 |
| 716 May 94 | 14 | 8 |
| 703 Aug 94 | $14 / 15$ | 6 |
| 816 May 94 | 16 | 9 |
| 803 Aug 94 | 16 | 8 |


| Taxus sumatrana |  |  |
| :---: | :---: | :---: |
| de Laubenfels P671 (A). Tai-shu Shan For. Dist., |  |  |
| 2000 m (Rt. 210, 7km) | 10-11 | 6 |
| de Laubenfels P 670 (A). I.c. | 10 | 8 |
| C-j Chang |  |  |
| 3 (wba) 16 May 94 | 11 | 13 |
| 303 Aug 94 | 10-11 | 12 |

## Wallichiana Group <br> Chinensis Subgroup

Wilson 11154D (A: type in adnot.). Karenko
Prov:: mts W of Karenko 12-13 6
C-j Chang, Tongshi 5 (wba)
27 Sep 9314 4
26 Nov 93125
06 Dec 93 11 5
09 Dec $93 \quad 12 \quad 5$
13 Jan $94 \quad 114$
C-j Chang, Tongshi 7 (wba).
27 Sep $9314 \quad 7$
26 Nov $9314 \quad 9$
06 Dec 93145
09 Dec $9314 \quad 5$
27 Jan $94 \quad 145$
C-j Chang 6 (wba). Hua-lien
16 May 94 09-10 8
03 Aug 94 11 8-9

Origin?
Y. Sugilara, Ex TUS (GH).[arbitrarily placed here]

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THE PHILIPPINES

## Sumatrana Group Taxus sumatrana

| Merrill 4595 (US). Lepanto Dist., Mt. Data de Laubenfels P650 (GH). Benguet, 58 km | 12-13 | 14 |
| :---: | :---: | :---: |
| N of Baguio, 2100 m | 11-13 | 13 |
| Whitehead 1896 (BM). NW central Luzon: $5000-7000 \mathrm{ft}$ | 12 | 12 |
| Williams 1002 (US). Benguet: Mt. St. Tomas | 12-13 | 10 |
| Leano 20672 (US). Benquet Prov. |  |  |
| Mt. St. Tomas | 11-13 | 9 |
| Vidal 623 (GH); Mt. Banahao, Pr. Tayabas | 11 | 7 |

## Wallichiana Group

Chinensis Subgroup
E.C. Leano 25128 (US). Luzon: Benquet Prov.,
Mt. St. Tomas 164

Elmer 6244 (P). Mt. St. Tomas 144
Elmer 6244 (US) 124
Loher 4850 (K). Luzon: central. 144
Loher 4850 (US) 124
Curran 5015 (P). Benquet Prov., Mt. Tonglan 147
Curran 5015 (PH) 144
Jacobs 7171 (K). Luzon: Mt. Pulog,
2200-2300 m 14 2-3

Ramos \& Edaño 40234 (K). Luzon: Lepanto,
Mt. Data 142

Ramos \& Edaño 40234 (P) 124
Wilkes Exped. 1838-1842 (GH). Luconia: Mt Mahaihai
$13 \quad 2$
Curran 7911 (US). Luzon: Benquet Prov.,
Mt. Banajao
Merrill 839 (US). Luzon, Benquet Prov. 12 3-4
Merrill 839 (US) 124
Ocampo 27920 (A). Mt. Banajao 11 -
Ocampo 27920 (P) 11 -
de Laubenfels P669 (GH). Luzon: Laguna Prov.,
Mt. Banajao, 2100 m 114
de Laubenfels P668. (GH: type in adnot.) - 4
Herb. Hook. (K). Luconia, 7600 ft 10-11 4
Loher 7139 (US) - 3
Sulit 2350 (A). Luzon: Benguet Prov., Mt. Pauai,
2450 m 158

Alvarey 18369 (BM). Benquet Prov. 12-14 8
INDONESIA
Sulawesi (Celebes)

## Sumatrana Group Taxus celebica

Everett 35 (K: type, Podocarpus celebicus
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Teysmann 14190 (U). Bonthian
11-13 16


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## REFERENCES

Alry Shaw, H.K. 1973. A dictionary of the flowering plants and ferns. 8th ed. Revised from J.C. Willis. Cambridge University Press, London.
André, D. 1956. Contribution à l'étude morphologique du cône femelle de quelques gymnospmermes (Cephalotaxacées, Juniperoidées, Taxacées). Nat. Monspliensia (Bot.) 8:3-35.
Axelrod, D.I. 1958. Evolution of the Madro-Tertiary Geoflora. Bot. Rev. 24:433-509.
Axelrod, D.I. 1975. Evolution and biogeography of Madrean-Tethyan sclerophyll vegetation. Ann. Missouri Bot. Gard. 62:280-334.
Axelrod, D.I. 1976. History of the coniferous forests, California and Nevada. Univ. Calif. Press, Berkeley, 1-62.
Axelrod, D.I. 1983. Biogeography of oaks in the Arcto-tertiary Province. Ann. Missouri Bot. Gard. 70:629-657.
Axelrod, D.I. 1986. Cenozoic history of some western American pines. Ann. Missouri Bot. Gard. 73:565-641.
Axelrod, D.I., I. Al-ShehbaZ, and P.H. Raven. 1998. History of the modern flora of China. In: Z. Aoluo and W. Sugong, eds. Proceedings of the First International Symposium on Floristic characteristics and diversity of East Asian plants, July 25-27, 1996, Kunming, Yunnan. China Higher Education Press, Beijing and Springer-Verlag Berlin Heidelberg. Pp. 43-55.
Balley, L.H. 1933. The cultivated conifers of North America. MacMillan Co., New York.
Bartholomew, B. 1999. Re-evaluation of the biological effect of plate movement.—Impact of Shan—Malay plate displacement (The movement of Burma-Malaya Geoblock) on the biota of the Gaoligong Mountains. Acta Bot. Yunnanica 21:407-425.
Bennike O. 1990. The Kap København Formation: stratigraphy and paleobotany of a Plio-Pleistocene sequence in Peary Land, North Greenland. Meddel. Grønland Geosci. 23:1-85.
Bertrand, C.E. 1874. Anatomie comparée des tiges et des feuilles chez les Gnétacées et les Conifères. Ann. Sci. Nat. Bot. sér. 5, 20:6-153.
Bobola, M.S., R.T. Eckert, A.S. Klein, K. Stapelfe, D.E. Smith, and D. Guenette. 1996. Using nuclear and organelle DNA markers to discriminate among Picea rubens, Picea mariana, and their hybrids. Canad. J. For. Res. 26:433-443.
Bobrov, A.V.F. C.H., A.P. Melkian, S. Mikhall, S. Romanov, and A.N. Sorokin. 2004. Seed morphology and anatomy of Austrotaxus spicata (Taxaceae) and its systematic position. Bot. J. Linn. Soc. 145:437-443.
Boufford, D.E. and S.A. Spongberg. 1983. Eastern Asian-eastern North American phytogeographical relationships—A history from the time of Linnaeus to the twentieth century. Ann. Missouri Bot. Gard. 70:421-439.
Braun, L. 1950. Deciduous forests of eastern North America. The Blankiston Co., Toronto.
Burnham, R.J. and A. Graham. 1999. The history of Neotropical vegetation: New developments and status. Ann. Missouri Bot. Gard. 86:546-589.
Chaney, R.W. 1947. Tertiary centers and migration routes. Ecol. Monogr. 17:139-148.
Chaw, S-M, H. Long, B-s Wang, A. Zharkikh, and W-h LI. 1993. The phylogentic position of Taxaceae based on 18s rRNA sequences. J. Molec. Evol. 37:624-630.

Chaw, S-m, H-m Sung, H. Long, A. Zharkikh, and W-н. Lı. 1995. The phylogenetic position of the conifer genera Amentotaxus, Phyllocladus, and Nageia inferred from 18S rRNA sequences. J. Molec. Evol. 41:224-230.
Cheng (Zheng), W-c. 1934. An enumeration of vascular plants from Chekiang, III. Gymnospermae. Contrib. Biol. Lab. Chinense Assoc. Adv. Sci. 9:240-241.
Cheng (Zheng), W-c., and L-k. Fu. 1978. Taxaceae. In: Fl. Reipub. Pop. Sin. 7, Gymnospermae. Agendae Academiae Sinicae. [English Translation].
Cheng, Y., R.G. Nicolson, K. Tripp, and S-M. Chaw. Phylogeny of Taxaceae and Cephalotaxaceae genera inferred from chloroplast matK gene and nuclear rDNA ITS region. 2000. Molec. Phylogen. Evol. 14:353-365.
Collins, D., R.R. Mill, and M. Möller. 2003. Species separation of Taxus baccata, T. canadensis, and T. cuspidata (Taxaceae) and origins of their reputed hybrids inferred from RAPD and cpDNA data. Amer. J. Bot. 90:175-182.
Cope, E.A. 1998. Taxaceae: The genera and cultivated species. Bot. Rev. 64:291-322.
Culberson, C.F., W.L. Culberson, and A. Johnson. 1990. The Ramalina americana complex. Bryologist 93:167186.

Dempsey, D. and I. Hook. 2000. Yew (Taxus) species—chemical and morphological variations. Pharm. Biol. 38:274-280.
Deryugina, T.F. and N.D. Nesterovich. 1981. Peculiarities of the morphological and anatomical structure of conifer needles of some Taxus L. species. Dokl. Akad. Nauk Bel. S.S.R. 25(7):652-655 [Russian with English summary].
Dilcher, D.L. 1969. Podocarps from the Eocene of North America. Science 164:299-301.
Dung, V., Van (ed.). 1996. Vietnam forest trees. Ten contributors, Agric. Publ. House, Hanoi.
Dupler, A. 1920. Ovuliferous structures of Taxus canadensis. Bot. Gaz. 69:492-520.
El-Kassaby, Y.A. and A.D. Yanchuk. 1994. Genetic diversity, differentiation, and inbreeding in Pacific yew from British Columbia. J. Hered. 85:112-117.
El-Kassaby, Y.A. and A.D. Yanchuk. 1995. Genetic variation of Pacific Yew in British Columbia and its conservation. In: Ph. Baradat, W.T. Adams, and G. Müller-Starck, eds. Population genetics and genetic conservation of forest trees. Academic Publishing, Amsterdam, The Netherlands. Pp. 227-235.
Fady, B., M. Arbez, and A. Marpeau. 1992. Geographic variability of terpene composition in Abies cephalonica Loudon and Abies species around the Aegean: hypotheses for their possible phylogeny from the Miocene. Trees 6:162-171.
Fady-Welterlen, B. 2005. Is there really more biodiversity in Mediterranean forest ecosystems? Taxon 54: 905-910.
Farjon, A. 1990. Pinaceae. Koeltz Scientific Books, Königstein, Germany.
Farjon, A. 1998. World checklist and bibliography of conifers. The Royal Botanic Gardens, Kew.
Farjon, A. 2001. World Checklist and Bibliography of Conifers. 2nd edition. The Bath Press, Bath, United Kingdom.
Ferguson, D.K. 1978. Some current research on fossil and recent taxads. Rev. Palaeobot. Palynol. 26:213226.

Florin, R. 1931. Untersuchungen zur Stammesgeschichte der Coniferales und Cordaitales. Kongl. Svenska Vetenskapsakad. Handl. 10:45-588.
Florin, R. 1948a. Enumeration of gymnosperms collected on Swedish expeditions to western and northwestern China. Acta Horti Berg. 14:343-384.
Florin, R. 1948b. On the morphology and relationships of the Taxaceae. Bot. Gaz. 110:31-39.
Florin, R. 1948c. On Pseudotaxus, a new genus of the Taxaceae, from eastern China. Acta Horti Berg. 14:385-395, plates 1-3.
Florin, R. 1948d. Nothotaxus or Pseudotaxus? Bot. Not. 1948:270-272.
Florin, R. 1951. Evolution in cordaites and conifers. Acta Horti Berg. 16:285-388, plate.
Florin, R. 1958. On Jurassic taxads and conifers from north-western Europe and eastern Greenland. Acta Horti Berg. 17:257-402, 56 plates.

Florin, R. 1963. The distribution of conifer and taxad genera in time and space. Acta Horti Berg. 20:121312.

Franco, J.A. 1964. Taxus. Fl. Europaea 1:39.
Frederiksen, N.O. 1994. Paleocene flora diversities and turnover events in eastern North America and their relation to diversity models. Rev. Palaeobot. Palynol. 82:225-238.
Frederiksen, N.O. 1995. Differing Eocene floral histories in southeastern North America and western Europe: Influence of paleogeography. Hist. Biol. 10:13-23.
Frimmel, F. von. 1911. Die untere Kutikula des Taxus- Blattes—ein Lichtreflektor. Oesterr. Bot. Z. 61:216-223.
Fryday, A.M. 2006. New and interesting North American lichen records from the alpine and sub-alpine zones of Mt. Katahdin, Maine. The Bryologist 109:570-578.
Fu, L-k, N. Lı and R.R. Mıl.. 1999. Taxaceae. Fl. China 4:89-96, Missouri Botanical Garden Press, St. Louis.
Gaussen, H. 1979. Les gymnospermes actuelles et fossiles: fasc. 15. Les taxines. Toulouse, Unversité Paul-Sabatier Faculté des Sciences.
Gelderen, D.M. van and J.R.P. van Hoey Smith. 1996. Conifers. An illustrated encyclopedia. 2 Vols. Timber Press, Portland, OR.
Gernandt, G.S., G.G. López, S.O. García and A. Liston. 2005. Phylogeny and classification of Pinus. Taxon 54:29-42.
Good, R. 1964. The geography of the flowering plants. John Wiley \& Sons, New York.
Givulescu, R. 1973. Die fossil Koniferen des Fundortes Chiuzbaia F. Inst. Géol. Mém. 19:31-34.
Graham, A. 1972. Outline of the origin and historical recognition of floristic affinities between Asia and eastern North America. In: A. Graham, ed., Floristics and paleofloristics of Asia and eastern North America, Elsevier Publ. Co., Amsterdam. Pp. 1-18.
Graham, A. 1973. History of the arborescent temperate element in the northern Latin American Biota. In: A. Graham, ed., Vegetation and vegetational history of northern Latin America. Elsevier Scientific Publ. Co., Amsterdam. Pp. 301-314.
Graham, A. 1993. History of the vegetation: Cretaceous (Maastrichtian)—Tertiary. FI. North America 1:57-70.
Graham, A. 1999. Late Cretaceous and Cenozoic history of North American vegetation north of Mexico. Oxford Univ. Press, New York.
Hageneder, F. 2007. Yew. A history. Sutton Publishing Ltd., Thrupp-Stroud-Goucestershire.
Hamllton, W. 1983. Cretaceous and Cenozoic history of northern continents. Ann. Missouri Bot. Gard. 70: 440-458.
HARA, H. 1972. Corresponding taxa in North America, Japan and the Himalayas. In: D.H. Valentine, ed. Taxonomy, phytogeography and evolution. Academic Press, New York. Pp. 61-72.
Harris, T.M. 1976a. Two neglected aspects of fossil conifers. Amer. J. Bot. 63:902-910.
Harris, T.M. 1976b. The Mesozoic gymnosperms. Rev. Paleobot. and Palyn. 21:119-134.
HayAsh, Y. 1954. The natural distribution of important trees, indigenous to Japan. Conifers Report 3. Bull. For. Exp. Sta. Meguro, Tokyo 75:166-173 (English summary).
HEINZE, B. 2004. Zur Populationsbiologie der gemeinen Eibe (Taxus baccata). Centralblatt für das gesamte Forstwesen 121:47-59.
HILL, K.D. 1998. Gymnosperms—The paraphyletic stem of seed plants. Fl. Australia 48:505-526.
Hils, M. 1993. Taxaceae Gray. Yew family. Fl. North America 2:423-427.
Holmgren, P.K., N.H. Holmgren, and L.C. Barnett (eds.). 1990. Index Herbariorum. Part I: The herbaria of the world. $8^{\text {th }}$ ed., W. Junk, Hague.
Hou, H-Y. 1983. Vegetation of China with reference to its geographical distribution. Ann. Missouri Bot. Gard. 70:509-548.
Hsü, J. 1983. Late Cretaceous and Cenozoic vegetation in China, emphasizing their connections with North America. Ann. Missouri Bot. Gard. 70:490-508.
Humphries, C.J. 1982. Vacariance biogegoraphy in Mesoamerica. Ann. Missouri Bot. Gard. 69:444-463.
Hunt, R.S. 1993. Abies Miller. Fir. Fl. North America 2:354-362.

Jacobs, B.F., J.D. Kingston, and L.L. Jacobs. 1999. The origin of grass-dominated ecosystems. Ann. Missouri Bot. Gard. 86:590-643.
Jinxing, L. and H. Yuxı (eds.). 2000. Atlas of structure of gymnosperms. Science Press, Beijing.
Klaus, W. 1989. Mediterranean pines and their history. Pl. Syst. Evol. 162:133-163.
KlıCKA, J. and R.M. Zink. 1997. The importance of recent ice ages in speciation: A failed paradigm. Science 277:1666-1669.
Kornas, J. 1972. Corresponding taxa and their ecological background in the forest of temperate Eurasia and North America. In: D.H. Valentine, ed. Taxonomy, phytogeography and evolution. Academic Press, New York. Pp. 37-59.
Kotyk, M.E. A., J.F. Basinger, and E.E. McIver. 2003. Early Tertiary Chamaecyparis Spach from Axel Heiberg Island, Canadian High Arctic. Canad. J. Bot. 81:113-130.
Krüssmann, G. 1985. Manual of cultivated conifers. Translated by M. E. E, eds. H.-D. Warda \& G. S. Daniels. Timber Press, Portland.
Krishnan, M.S. 1974. III. Geology. In: M.S. Mani, ed. Ecology and biogeography of India. W. Junk, Hague. Pp. 60-98.
Kubitzkı, K. and W. Krutzsch. Origins of East and Southeast Asian plant diversity. 1998. In: Z. Aoluo and W. Sugong, eds. Proceedings of the First International Symposium on Floristic characteristics and diversity of East Asian plants, July 25-27, 1996, Kunming, Yunnan. China Higher Education Press, Beijing and Springer-Verlag Berlin Heidelberg. Pp. 56-70.
KvačEk, Z. [1984 fide author]. Tertiary taxads of NW Bohemia. 1982 Acta Univ. Carol., Geol., Pokorny 4:471-491.
KvaČek, Z. and H. Walther. 1998. The Oligocene flora of Kundratice near Litoměřice, České středohoří Volcanic Complex (Czech Republic)—a review. Acta Mus. Nat. Pragae, B, 54:1-42.
Kvaček, Z. and W.C. Rember. 2000. Shared Miocene conifers of the Clarkia flora and Europe. Acta Universitatis Carolinae - Geologica. 44(1):75-85.
Kvaček, Z., and W.C. Rember. Under Review. Calocedrus robustior (Cupressaceae) and Taxus howardii (Taxaceae): Two new conifers from the Middle Miocene of the western North America (Clarkia area, Latah Formation, northern Idaho). Paleobios.
KwEI, Y-L. and S.-Y. Hu. 1974. [Epidermal feature of leaves of Taxus in relation to taxonomy]. Acta Phytotax. Sin. 12(3):329-334, plate 67 [Chinese with English summary].
Larson, D.W., U. Matthes, J.A. Gerrath, N.W.K. Larson, J.M. Gerrath, J.C. Nekola, G.L. Walker, S. Porembski, and A. Charlton. 2000. Evidence for the widespread occurrence of ancient forests on cliffs. J. Biogeogr. 227:319-331.

Laubenfels, D.J., De. 1988. Coniferales. Fl. Malesiana 10(3):337-453.
Leopold, E.B. and H.D. MacGinitie. 1972. Development and affinities of Tertiary floras in the Rocky Mountains. In: A. Graham, ed. Floristics and paleofloristics of Asia and eastern North America. Elsevier Publ. Co., Amsterdam. Pp. 147-200.
LePAge, B.A. 2003. A new species of Thuja (Cupressaceae) from the Late Cretaceous of Alaska: implications of being evergreen in a polar environment. Amer. J. Bot. 90:167-174.
LI, H-L. 1952. Floristic relationships between eastern Asia and eastern North America. Trans. Amer. Philos. Soc. 42:371-429.
LI, J., C.C. Davis, P.D. Tredicl, and M.J. Donoghue. 2001. Phylogeny and biogeography of Taxus (Taxaceae) inferred from sequences of the internal transcribed spacer region of nuclear ribosomal DNA. Harv. Pap. Bot. 6:267-274
LI, X-w and J. LI. 1997. The Tanaka-Kaiyong Line—An important floristic line for the study of the flora of East Asia. Ann. Missouri Bot. Gard. 84:888-892.
Liston, A., W.A. Robinson, D. Piñero, and E.R. Alvarez-Buylla. 1999. Phylogenetics of Pinus (Pinaceae) based on nuclear ribosomal DNA internal transcribed spacer region sequences. Molec. Phylogen. Evol. 11:95-109.
LiTtLE, D.P. 2006. Evolution and circumscription of true cypresses (Cupressaceae: Cupressus). Syst. Bot. 31: 461-480.
Liu, T-s. 1971. A monograph of the genus Abies. Dept. For., College Agric., Natl. Taiwan Univ., Taipei.

Loudon, J.C. 1844. Arboretum et fruticetum Britannicum. Longman, Brown, Green and Longmans, London. MAI, D.H. 1989. Development and regional differentiation. PI. Syst. Evol. 162:79-91.
MAl, D.H. 1998. Contribution to the flora of the middle Oligocene Calu Beds in Brandenburg, Germany. Rev. Palaeobot. Palynol. 101:43-70.
Manchester, S.R. 1994. Fruits and seeds of the Middle Eocene Nut Beds Flora, Clarno Formation, Oregon. Palaeontographica Americana 58:1-205.
Manchester, S.R. 1999. Biogeographical relationships of North American Tertiary Floras. Ann. Missouri Bot. Gard. 86:472-522.
Manı, M.S. 1974. Biogeographical evolution in India. In: M.S. Mani, ed. Ecology and biogeography of India, W. Junk, Hague. Pp. 698-722.
Mclver, E.E. 1994. An early Chamaecyparis (Cupressaceae) from the Late Cretaceous of Vancouver Island, British Columbia, Canada. Canad. J. Bot. 72:1787-1796.
Mclver, E.E. 1999. Paleobotanical evidence for ecosystem disruption at the Cretaceous-Tertiary boundary from Wood Mountain, Saskatchewan, Canada. Canad. J. Earth Sci. 36:775-789.
Mclver, E.E. and J.F. Basinger. 1989. The morphology and relationships of Thuja polaris sp. nov. (Cupressaceae) from the early Tertiary Ellesmere Island, Arctic Canada. Canad. J. Bot. 67:1903-1915.
Mclver, E.E and J.F. BAsinger. 1999. Early Tertiary floral evolution in the Canadian High Arctic. Ann. Missouri Bot. Gard. 86:523-545.
McKenna, M.C. 1983. Holarctic landmass rearrangement, cosmic events, and Cenozoic terrestrial organisms. Ann. Missouri Bot. Gard. 70:459-489.
Meyen, S.V. 1984. Basic features of gymnosperm systematics and phylogeny as evidenced by the fossil record. Bot. Rev. 5:1-111.
Meyer, H.W. and S.R. Manchester 1997. The Oligocence Bridge Creek flora of the John Day Formation, Oregon. Univ, Calif. Publ. Geol. Sci 141:1-195, 75 pl.
Millar, C.I. 1993. Impact of the Eocene on the evolution of Pinus L. Ann. Missouri Bot. Gard. 80: 471-498.
Miller, C.N., Jr. 1977. Mesozoic conifers. Bot. Rev. 43:217-280.
Mılıer, C.N., Jr. 1988. The origin of modern conifer families. In: C.B. Beck, ed., Origin and evolution of gymnosperms. Columbia Univ. Press, New York. Pp. 448-486.
Miranda, F. and A.J. Sharp. 1950. Characteristics of the vegetation in certain temperate regions of eastern Mexico. Ecology 31:313-333.
Mirov, N.T. 1953. Taxonomy and chemistry of the white pines. Madroño 12:81-89.
NAKAI, T. 1938. Indigenous species of conifers and taxads of Korea and Manchuria, and their distribution. Chôsen Sanrin Kaihô (J. Kor. For. Soc.) 158:21.
Najman, Y. and E. Garzanti. 2000. Reconstructing early Himalayan tectonic evolution and paleogeography from Tertiary foreland basin sedimentary rocks, northern India. GSA Bulletin; 112 (3):435-449.
Nicolosi, R.T. 1982. Morphological features of leaves and pollen as in aid in separating selected species and cultivars within the genus Taxus. Thesis, Ph.D., Ohio State University.
Novacek, M.J. 1999. 100 million years of land vertebrate evolution: The Cretaceous-Early Tertiary transition. Ann. Missouri Bot. Gard. 86:230-258.
Ohwi, J. 1965. Flora of Japan. English translation, eds. F. G. Meyer \& E. H. Walker, Smithsonian Institution, Washington, DC.
O'Keefe, J.D. and T.J. Ahrens. 1989. Impact production of $\mathrm{CO}_{2}$ by the Cretaceous/Tertiary extinction bolide and the resultant heating of the Earth. Nature 338:247-249.
Ouden, P. den and B.K. Boom. 1965. Manual of cultivated conifers. M. Nijhoff, Hague.
Palamarev, E. 1989. Paleobotanical evidence of the Tertiary history and origin of the Mediterranean sclerophyll dendroflora. PI. Syst. Evol. 162:93-107.
PHIPPs, J.B. 1983. Biogeographic, taxononmic, and cladistic relationships between east Asiatic and North American Crataegus. Ann. Missouri Bot. Gard. 70:667-700.

Pilger, R. 1903. Taxaceae-Taxoideae—Taxeae. Taxus. In: Engler, Das Pflanzenreich IV:110-116.
Pilger, R. 1916. Die Taxales. Mitt. Deutsch. Dendrol. Ges. 25:1-28.
Pilger, R. 1926. Taxaceae. In: Engler, A. and K. Prantl, eds. Die natürlichen Pflanzenfamilien, 2nd ed., 13:199-211. Prager, E.M., D.P. Fowler and A.C. Wilson. 1976. Rates of evolution in conifers (Pinaceae). Evolution 30:637-649.
Prakash, U., D. Nal-zheng, and P.P. Tripathi. 1995. Fossil woods from the Miocene sediments of China with remarks on environmental implications of Miocene floras of the region. Barbal Sahni Centenary 1995:341-360.
Price, R.A. 1990. The genera of Taxaceae in the southeastern United States. J.Arnold Arbor. 71:69-71.
QiAn, H. 2002. Floristic relationships between eastern Asia and North America: Test of Gray's hypothesis. Amer. Nat. 160 (3):317-332.
Qian, H. and R.E. Ricklefs. 1999. A comparison of taxonomic richness of vascular plants in China and the United States. Amer. Nat. 154 (2):160-181.
Rao, A.S. 1974. The vegetation and phytogeography of Assam-Burma. In: M.S. Mani, ed. Ecology and biogeography of India. W. Junk, Hague. Pp. 204-246.
Rau, M.A. 1974. Vegetation and phytogeography of the Himalaya. In: M.S. Mani, ed. Ecology and biogeography of India. W. Junk, Hague. Pp. 247-280.
Rehder, A. 1919. New species, varieties and combinations from the herbarium and the collections of the Arnold Arboretum. J. Arnold Arbor. 1: 44-60.
Rehder, A. 1940. Manual of cultivated trees and shrubs. 2nd ed. MacMillan Co., New York.
Rosen, D.E. 1975. A vicariance model of Caribbean biogeography. Syst. Zool. 24:431-464.
SAHNI, B. 1920. On certain archaic features in the seed of Taxus baccata, with remarks on the antiquity of the Taxineae. Ann. Bot. 34:117-133.
Sargent, C.S. 1914. Plantae wilsonianae. Vol. 2. Cambridge University Press, Cambridge, MA.
Schmid, M. 1974. Végétation du Viet-nam. Mém. Orstom 74, Paris.
Schuster, R. 1976. Plate tectonics and its bearing on the geographical origin and dispersal of angiosperms. In: C.B. Beck, ed. Origins and early evolution of angiosperms. Columbia Univ. Press, New York. Pp. 48-138.
Sharp, A.J. 1966. Some aspects of Mexican phytogeography. Ciencia (México) 24:229-232.
SilbA, J. 1984. An international census of the coniferae, I. Phytologia Mem. 7:1-79.
Singh, H. 1961. The life history and systematic position of Cephalotaxus drupacea Sieb. \& Zucc. Phytomorphology 11: 153-196.
Spıut, R.W. 1992. A taxonomic key to the species of Taxus. NCI Workshop on Taxus, Taxol, and Taxotere, Rockville, MD (Abstract only).
Spıut, R.W. 1993. Reliable morphological characters for distinguishing species of Taxus (Abstract). In: International yew resource conference. Yew (Taxus) conservation biology and interactions. Berkeley, CA. Pp. 39-40.
Sp.ut, R.W. 1994. A systematic treatment of fruit types. Mem. New York Bot. Gard., Vol 70.
Spıut, R.W. 1998. Two papers presented at the AIBS Annual Meeting, American Systematic Plant Taxonomists, Baltimore Convention Center, MD, 5 Aug. 1998. Abstracts published on the Internet, Botanical Society of America.
(a) Morphological evolution in the Taxus leaf and its significance to recognizing ecological species within the genus. http://www.ou.edu/cas/botany-micro/bsa-abst/section13/abstracts/107.shtml.
(b) Species of Taxus. http://www.ou.edu/cas/botany-micro/bsa-abst/section13/abstracts/112.shtml.

Spuut, R.W. 2000. Three papers presented at the joint meetings of the Botanical Society of America and American Systematic Plant Taxonomists, Portland, OR, August, abstracts published online and Amer. J. Bot.
(a) A phytogeographical analysis and classification of leaf morphological features in Taxus (Taxaceae). http:// www.ou.edu/cas/botany-micro/botany2000/section13/abstracts/27.shtml.
(b) The morphological relationships of Taxus canadensis (Taxaceae) in North America and Eurasia. http://www. ou.edu/cas/botany-micro/botany2000/section13/abstracts/28.shtml.
(c) A revised taxonomic key to species and varieties of Taxus (Poster). http://www.ou.edu/cas/botany-micro/ botany2000/section13/abstracts/166.shtml. (Note key also presented on the WBA website in 2003, extracted from poster document of 24 pages that included 270 figures).

SpJut, R.W. 2006. Biogeographical data on putative hybrids for species of Taxus (Taxaceae) in the Himalayas and North America. Abstract, Botanical Society of America, Chico, CA, July 29-Aug 2. http://www.botanyconference.org/engine/search/index.php?func=detail\&aid=37
Spıut, R.W. 2007. Taxonomy and nomenclature of Taxus (Taxaceae). J. Bot. Res. Inst. Texas 1:203-289.
Srivastava, S.K. 1994. Evolution of Cretaceous phytogeoprovinces, continents and climates. Rev. Paleobot. Palynol. 82:197-224.
Stafleu, F.A. and R.S. Cowan. 1976-1988. TL-2. Taxonomic literature. 7 Vols. W. Junk, Hague.
Strauss, S.H., A.H. Doerksen, and J.R. Byrne. 1989. Evolutionary relationships of Douglas-fir and its relatives (genus Pseudotsuga) from DNA restriction fragment analysis. Canad. J. Bot. 68:1502-1510.
Suyama, Y., H. Yoshimaru, and Y. Tsumura. 2000. Molecular phylogenetic position of Japanese Abies (Pinaceae) based on chloroplast DNA sequences. Mol. Phylogen. Evol. 16:271-277.
Thomas, P.A. and A. Polwart. 2003. Biological flora of the British Isles. Taxus baccata L. J. Ecol. 91:489-524.
Thorne, R.F. 1972. Major disjunctions in the goegraphic ranges of seed plants. Quart. Rev. Biol. 47:365-411.
Thorne, R.F. 1978. Plate tectonics and angiosperm distribution. Notes Roy. Bot. Gard. Edinburgh 30:297-315.
Tiffney, B.H. 1985a. Perspective on the origin of the florisitc similarity between eastern Asia and eastern North America. J. Arnold Arbor. 66:73-94.
Tiffney, B.H. 1985b. The Eocene North Atlantic Land Bridge: Its importance in Tertiary and modern phytogeography of the northern hemisphere. J. Arnold Arbor. 66:243-273.
Tschudy, R.H., C.L. Pillmore, C.J. Orth, J.S. Gilmore, and J.D. Knight. 1984. Disruption of the terrestrial plant ecosystem at the Cretaceous-Tertiary Boundary, western interior. Science 255:1030-1032.
Tsumura, Y., K. Yoshimura, N. Tomaru, and K. Ohba. 1995. Molecular phylogeny of conifers using RFLP analysis of PCR-amplified specific chloroplast genes. Theoret. Applied Genet. 91:1222-1236.
Veitch, J. and Sons. 1881. A manual of the Coniferae. Published by the authors, Kings Road, Chelsea.
Wang, Q., D.L. Dilcher, X-y. Zhu, Y-l. Zhou, and T.A. Lott. 2006. Fruit and leaflets of Wisteria (Leguminosae, Papilionoideae) from the Miocene of Shandong Province, eastern China. Int. J. Plant Sci. 167:1061-1074.
Wang, T., Y-ı. Su, J-m. Zhu, G-k. Fan, and C. Huang. 2000. RAPD Analyses of Taxaceae and it related taxa. Acta Sci. Nat. Univ. Sunyatseni 39:129-130.
Ward, P.D., J.M. Hurtado, J.L. Kirschvink, and K.L. Verosub. 1997. Measurements of the Cretaceous paleolatitude of Vancouver Island: Consistent with the Baja-British Columbia Hypothesis. Science 277:1642-1645.
Whittaker, R.E. 1961. Vegetation history of the pacific coast states and the "central" significance of the Klamath Region. Madroño 16:5-21.
Wilkinson, R.C., J.W. Hanover, J.W. Wright, and R.H. Flake. 1971. Genetic variation in the monoterpene composition of white spruce. For. Sci. 17:83-90.
Wolfe, J.A. 1969. Neogene floristic and vegetational history of the Pacific Northwest. Madroño 20:83-110.
Wolfe, J.A. 1975. Some aspects of plant geography of the northern hemisphere during the late Cretaceous and Tertiary. Ann. Missouri Bot. Gard. 62:264-279.
Wood, C.E., Jr. 1972. Morphology and phytogeography:The classical approach to the study of disjunctions. Ann. Missouri Bot. Gard. 59:107-124.
Xiang, Q-y., S.R. Manchester, D.T. Thomas, W. Zhang, and C. Fan. 2005. Phylogeny, biogeography, and molecular dating of Cornelian cherries (Cornus, Cornaceae): Tracking Tertiary plant migration. Evolution 59:1685-1700.
Xiang, Q-Y., D.T. Thomas, W. Zhang, S.R. Manchester, and Z. Murrell. 2006. Species level phylogeny of the genus Cornus (Cornaceae) based on molecular and morphological evidence-implications for taxonomy and Tertiary intercontinental migration. Taxon 55:9-30.
Zhengyı, W. and W. Sugong. 1998. A proposal for a new floristic kingdom (realm). In: Z. Aoluo and W. Sugong, eds. Proceedings of the First International Symposium on Floristic characteristics and diversity of East Asian plants, July 25-27, 1996, Kunming, Yunnan. China Higher Education Press, Beijing and Springer-Verlag Berlin Heidelberg. Pp. 3-42.

