Morphological analyses suggest a new taxonomic circumscription for *Hymenaea courbaril* L. (Leguminosae, Caesalpinioideae)

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Abstract

*Hymenaea* is a genus of the Resin-producing Clade of the tribe Detarieae (Leguminosae: Caesalpinioideae) with 14 species. *Hymenaea courbaril* is the most widespread species of the genus, ranging from southern Mexico to southeastern Brazil. As currently circumscribed, *H. courbaril* is a polytypic species with six varieties: var. *altissima*, var. *courbaril*, var. *longifolia*, var. *stilbocarpa*, var. *subsessilis*, and var. *villosa*. These varieties are distinguishable mostly by traits related to leaflet shape and indumentation, and calyx indumentation. We carried out morphometric analyses of 14 quantitative (continuous) leaf characters in order to assess the taxonomy of *H. courbaril* under the Unified Species Concept framework. Cluster analysis used the Unweighted Pair Group Method with Arithmetic Mean (UPGMA) based on Bray-Curtis dissimilarity matrices. Principal Component Analyses (PCA) were carried out based on the same morphometric matrix. Two sets of Analyses of Similarity and Non Parametric Multivariate Analysis of Variance were carried out to evaluate statistical support (1) for the major groups recovered using UPGMA and PCA, and (2) for the varieties. All analyses recovered three major groups coincident with (1) var. *altissima*, (2) var. *longifolia*, and (3) all other varieties. These results, together with geographical and habitat information, were taken as evidence of three separate metapopulation lineages recognized here as three distinct species. Nomenclatural adjustments, including reclassifying formerly misapplied types, are proposed.

Keywords

Fabaceae, Detarieae, integrative taxonomy, nomenclatural changes
Introduction

*Hymenaea* L. is a genus of caesalpinioioid legumes with 14 species (Lee and Langenheim 1975). The genus is distributed throughout tropical America, from Mexico to Paraguay, with one species in coastal East Africa (Mackinder 2005). *Hymenaea* is included in the Resin-producing Clade of the tribe Detarieae, and is most closely related to the genera *Guibourtia* Benn. and *Peltogyne* Vogel (Bruneau et al. 2001, 2008; Fougère-Danezan et al. 2007), all presenting similar leaf morphologies, with two asymmetrical pellucid-punctate leaflets. *Hymenaea* can be differentiated from these related genera by its rather larger and more massive bat-pollinated flowers with a robust hypanthium, and indehiscent, pulpy and woody pods with very large seeds.

The current taxonomy of the genus is largely based on Lee and Langenheim’s (1975) revision. In addition to recognizing the 14 currently accepted species, they reduced several species to varietal rank. These polytypic species were viewed by Lee and Langenheim (1975) as showing complex relationships with other species of *Hymenaea*. They hypothesized, for example, that *H. oblongifolia* Huber var. *oblongifolia* is more closely related to *H. aurea* Lee and Lang. and *H. eriogyne* Benth., while *H. oblongifolia* var. *davisi* (Sandwith) Lee and Lang. is probably more closely related to *H. parvifolia* Huber, *H. rubriflora* Ducke, and *H. reticulata* Ducke. According to their concepts, *H. oblongifolia* should be regarded as a polyphyletic species. Another example of a putative polyphyletic species, according to Lee and Langenheim’s (1975) conceptual framework, is *H. courbaril*, with var. *villosa* Lee and Andrade-Lima hypothesized as being more closely related to *H. martiana* Hayne, and var. *longifolia* (Benth.) Lee and Andrade-Lima to *H. velutina* Ducke and *H. stigonocarpa* Mart. ex Hayne.

*Hymenaea courbaril* is the most widely distributed species of the genus, almost matching the geographic range of *Hymenaea* in the New World. It also has the greatest economic importance in the genus, due to the high quality of its wood and its resin, the latter being used by native populations as incense, cement, in the manufacture of varnishes, and for medicinal purposes. Its nutritive fruits are sought after by mammals and birds (Rizzini 1971; Langenheim 1967; Lee and Langenheim 1975). *Hymenaea courbaril* is the most taxonomically complex species, with six varieties: var. *altissima* (Ducke) Lee and Lang., var. *courbaril*, var. *longifolia*, var. *stilbocarpa* (Hayne) Lee and Lang., var. *subsessilis* Ducke, and var. *villosa*. These varieties are differentiated by their leaflet sizes, shapes, and indumentation, calyx indumentation, petal shapes, ovary stipe sizes, and pod sizes and shapes. These variations in several diagnostic features make the boundaries of putatively related taxa rather imprecise. The widely circumscribed *H. courbaril*, as defined by Lee and Langenheim (1975), is hereafter referred to as the *H. courbaril* complex.

The species and varietal limits of *H. courbaril*, *H. stigonocarpa*, and *H. martiana* were investigated by Pestana (2010). This author did not employ objective analytical methods and used the same classical taxonomic approach as Lee and Langenheim (1975), thus coming to similar conclusions as the latter authors in keeping *H. courbaril* as a polytypic species with six varieties.
Delineating precise species boundaries is a key task in plant taxonomy. This process has direct impacts on society, as there is a growing demand for credible taxonomic information that allows us to conserve, manage, and understand natural biodiversity (Wheeler et al. 2004). However, questions of species recognition can be affected by several theoretical, methodological, and practical issues. De Queiroz (2005, 2007) defined species as separately evolving metapopulation lineages (the Unified Species Concept—USC), and proposed that all other previously considered properties of species should be reinterpreted as contingent rather than critical. These additional contingent properties, such as phenetic distinctiveness, reciprocal monophyly, genetic coalescence, or ecological distinctiveness, are acquired during speciation and should be considered as different lines of evidence relevant to assessing lineage separation. The USC, by treating species conceptualization and species delimitation as clearly separate issues, allows the use of properties formerly treated as secondary criteria in species delimitations. More importantly, it allows for currently accepted species limits to be considered as hypotheses to be tested using the presence of any one of those secondary properties as evidence for the existence of a species.

Analyses of morphometric data can be useful in objectively demonstrating species limits, especially when combined with molecular markers (Andrés-Sánchez et al. 2009; Newmaster and Ragupathy 2009). Additionally, the use of morphological information represents the fastest and least expensive manner of assessing taxonomic complexes—and has been used to solve problems of species limits in many different plant groups, especially when molecular data was not easily available (Handerson 2006; Estrella et al. 2009; Pedersen 2010; Ceolin and Miotto 2012; Rahman and Rahman 2012; Castello and Galetto 2013; Scrivanti et al. 2013), as was the case of the group studied here.

The problem of defining species limits in the polytypic *H. courbaril* complex is revisited here under the USC conceptual framework by exploring morphometric, geographical, and ecological patterns as lines of evidence for the existence of separate metapopulation lineages. Specifically, we sought to test Lee and Langenheim’s (1975) hypothesis that *H. courbaril* should be treated as a polytypic species with six varieties.

**Materials and methods**

A total of 96 specimens of the *H. courbaril* complex were examined in this study (vouchers listed in Appendix 1). All analyzed materials were sheets from the following herbaria: B, CEN, CEPEC, HRB, HUEFS, IBGE, IPA, LAGU, M, MBM, NY, RB, SP, SPF, U, UB, and UC. We selected specimens that displayed branch ends with fully-developed (mature) leaves. This criterion avoided considering young leaves from the tips of the branches or leaves at the bases of the branches that are often much larger. Each specimen analyzed was considered an individual, and identifications strictly followed Lee and Langenheim (1975), although these names were only used as nomenclatural references in this study.

The leaves in the group studied here are bifoliolate; the leaflets range from oblong to ovate or obovate, with rounded, acute or obtuse apices; the main vein is displaced...
towards the inner margin resulting in an asymmetrical base, the outer portion being wider than the inner and extending beyond the attachment to the petiolule (Figure 1). Fourteen quantitative (continuous) characters were examined (Table 1; Figure 1). Only leaf traits were measured and quantified, as most herbarium sheets lacked flowers and/or fruits. In any case, flower morphology is much conserved in the species studied and the herbarium material examined usually contained only incomplete or damaged flowers. Measurements were taken of two fully developed leaves per dried herbarium sheet, using a graduated ruler (precision 1 mm).

All multivariate analyses were carried out using Past software (Hammer et al. 2001). Cluster analyses used the Unweighted Pair Group Method with Arithmetic Mean (UP-
Morphological analyses suggest a new taxonomic circumscription...

GMA) based on Bray-Curtis dissimilarity matrices. Principal Component Analyses (PCA) were carried out based on the same morphometric matrix. Two sets of Analyses of Similarity (ANOSIM; Warwick, Clarke and Suharsono 1990) and Non Parametric Multivariate Analysis of Variance (NPMANOVA; Anderson 2001) were carried out to evaluate statistical support for: (1) the major groups recovered in the UPGMA and PCA; and (2) the varieties as defined by Lee and Langenheim (1975). *Hymenaea courbaril* var. *villosa* was not included in the second analysis because it is known from only two specimens (Lee and Langenheim 1975) and only one was located during this study. Both ANOSIM and NPMANOVA used Bonferroni corrections, 10,000 permutations, and Bray-Curtis distances. As *H. courbaril* var. *villosa* was represented by only a single specimen, we carried out a second set of UPGMA, PCA, and similarity analyses for the major groups without including this variety in order to test its influence on the results.

Distribution maps of the specimens studied were prepared using DIVA-GIS software (Hijmanns et al. 2005), based on the geographic coordinates recorded on the herbarium sheet labels. For material lacking original coordinates, a central coordinate for the municipality was used as provided by the Species Link website (available at http://specieslink.org.br).

Species limits were tested following the USC framework (de Queiroz 2005, 2007). Species ranks were ascribed to groups that showed morphological and habitat distinctiveness and geographical consistency. We considered as morphologically distinct groups those that were recovered in UPGMA and PCA and that exhibited statistical significance in both ANOSIM and NPMANOVA tests. Habitat distinctiveness was assessed from the vegetation type where the taxon occurs, following the UNESCO (1973) classification. Species diagnoses were prepared based on vegetative and reproductive characters.

### Table 1.

<table>
<thead>
<tr>
<th>Leaf characters (cm)</th>
<th>Measurements in Figure 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 - Total length of the leaflet</td>
<td>A</td>
</tr>
<tr>
<td>2 - Leaflet length /width ratio</td>
<td>A / B</td>
</tr>
<tr>
<td>3 - Length of the distal sixth</td>
<td>E</td>
</tr>
<tr>
<td>4 - Inner width in distal third</td>
<td>C</td>
</tr>
<tr>
<td>5 - Outer width in distal third</td>
<td>D</td>
</tr>
<tr>
<td>6 - Inner width in middle third</td>
<td>F</td>
</tr>
<tr>
<td>7 - Outer width in middle third</td>
<td>G</td>
</tr>
<tr>
<td>8 - Inner width in basal third</td>
<td>H</td>
</tr>
<tr>
<td>9 - Outer width in basal third</td>
<td>I</td>
</tr>
<tr>
<td>10 - Main vein displacement</td>
<td>(G–F) / (I–H)</td>
</tr>
<tr>
<td>11 - Base extension (length of the base from the inner attachment of the petiolule)</td>
<td>J</td>
</tr>
<tr>
<td>12 - Petiole length</td>
<td>M</td>
</tr>
<tr>
<td>13 - Petiolule length</td>
<td>L</td>
</tr>
<tr>
<td>14 - Distance between inner and outer attachment of the petiolule</td>
<td>K</td>
</tr>
</tbody>
</table>

Species diagnoses were prepared based on vegetative and reproductive characters.
Results and discussion

Both UPGMA and PCA recovered three major groups (Figure 2): Group 1 included all specimens of var. *altissima*; Group 2 all specimens of varieties *courbaril*, *stilbocarpa*, *subsessilis*, and the single specimen of *villosa*; and Group 3 all specimens of var. *longifolia*. Within Group 2, individuals of the different varieties did not cluster together and appeared intermixed in UPGMA, or formed highly overlapping groups in two first axes of PCA. The first PCA axis accumulated 88.3% of the total variance, with the two first axes summing 93% of the observed variation. Leaflet length was the trait that explained most of the variation found in first axis, and the three major groups were sorted mostly by leaflet size. These results indicate that Group 3 includes specimens with largest leaflets, and Group 1 the smallest leaflets (Figure 3).

The results of the UPGMA and PCA were consistent with both ANOSIM and NPMANOVA. Comparisons of the varieties of *H. courbaril* showed significant differences (p < 0.05) between the varieties *altissima* and *longifolia* and all other varieties, but no significant differences between the varieties *courbaril*, *stilbocarpa*, and *subsessilis* that clustered in Group 2 (Table 2). The same analyses comparing the three groups recovered in UPGMA and PCA found significant morphological differences between them (Table 3).

The complex morphological variations found in Group 2 appear to reflect its wide geographic range and large genetic variability (Ramos et al. 2009). Phylogeographic studies in part of the geographic range of *H. courbaril* var. *stilbocarpa* clearly demonstrated the wide genetic base and geographic structure of this genetic variation (Ramos et al. 2009). The morphological distinctions among the varieties clustered in Group 2 are made by rather continuous characters, such as the color of the calyx lobes (ochraceous to golden or rusty brown), leaflet shape (broadly to narrowly falcate), and the shapes of the outer sides of the leaflet base (straight to nearly straight or rounded). The var. *subsessilis* presents a short ovary stipe (c. 2 mm long) that grades to a medium-sized stipe in var. *stilbocarpa* (3–4 mm) and then large in var. *courbaril* (4–6 mm).

Leaflet indument distinguishes var. *villosa* from the remaining varieties of Group 2 (Lee and Andrade-Lima 1974, 1975). The ovary is also described as slightly pilose on one side, a condition not observed in other varieties of *H. courbaril*. The exclusion of var. *villosa* from the similarity analyses did not alter the results obtained with its inclusion (Table 3). The var. *villosa* is known from only two specimens from the eastern coastal areas of the states of Paraíba and Pernambuco (northeastern Brazil) where the range of var. *stilbocarpa* overlaps with that of *H. martiana*. This species (*H. martiana*) has tomentose leaflets and an ovary with a tuft of trichomes near its base. The rarity of *H. courbaril* var. *villosa*, as well as the transitional nature of the leaflets and ovary indumentation in relation to *H. martiana* and the other varieties of the *H. courbaril* complex clustered in Group 2, all suggest that var. *villosa* could represent a hybrid between *H. martiana* and *H. courbaril*.

The habitats of the taxa of Group 2 include mostly tropical ombrophilous alluvial (gallery) forests in areas subject to seasonally dry climates from Mexico and the
Figure 2. A Distribution of the Hymenaea courbaril complex in the Neotropics. The major biomes where this complex occurs are shown in color: wet forests (green), seasonally dry forests and woodlands (yellow), and savannas (purple). The varieties of H. courbaril recognized by Lee and Langenheim (1975) are represented by different symbols: var. altissima (blue squares), var. courbaril (yellow triangles), var. longifolia (red squares), var. stilbocarpa (blue triangles), var. subsessilis (purple circles), and var. villosa (white circles) B UPGMA analysis of the H. courbaril complex based on 14 quantitative leaf measures (see Figure 1 and Table 1 for measurement details and Bray-Curtis distances). Varieties are represented by the same symbols used for the map. C Scatter diagram showing the first two axes of the PCA using the same data matrix as the UPGMA analysis. Ellipses represent the varieties as recognized by Lee and Langenheim (1975), and they are represented by the same symbols used for the map.
Table 2. Results of ANOSIM and NPMANOVA testing the consistence of the varieties of *Hymenaea courbaril*, using 10,000 permutations and Bray-Curtis distances. ANOSIM R-values are given above the diagonal and NPMANOVA F-values are given below the diagonal. Numbers between brackets are Bonferroni-corrected p-values. Significant differences are indicated by shadowed cells.

<table>
<thead>
<tr>
<th></th>
<th>stilbocarpa</th>
<th>courbaril</th>
<th>subsessilis</th>
<th>altissima</th>
<th>longifolia</th>
</tr>
</thead>
<tbody>
<tr>
<td>stilbocarpa</td>
<td>0.09933 [0.0195]</td>
<td>-0.1343 [1]</td>
<td>0.9325 [0.0015]</td>
<td>0.8858 [0.0015]</td>
<td></td>
</tr>
<tr>
<td>courbaril</td>
<td>5.313 [0.0855]</td>
<td>0.09375 [1]</td>
<td>0.8524 [0.0015]</td>
<td>0.9471 [0.0015]</td>
<td></td>
</tr>
<tr>
<td>subsessilis</td>
<td>0.8937 [1]</td>
<td>3.296 [0.489]</td>
<td>0.9972 [0.0045]</td>
<td>0.9329 [0.0075]</td>
<td></td>
</tr>
<tr>
<td>altissima</td>
<td>79.79 [0.0015]</td>
<td>57.97 [0.0015]</td>
<td>49.86 [0.003]</td>
<td>1 [0.0015]</td>
<td></td>
</tr>
<tr>
<td>longifolia</td>
<td>67.13 [0.0015]</td>
<td>94.08 [0.0015]</td>
<td>30.27 [0.006]</td>
<td>235.5 [0.0015]</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. ANOSIM and NPMANOVA (10,000 permutations and Bray-Curtis distances) for the groups recovered in UPGMA and PCA analyses of 78 individuals of *Hymenaea courbaril* complex. Group 1 included all specimens of var. *altissima*, Group 2 all specimens of the varieties *courbaril*, *stilbocarpa*, *subsessilis*, and *villosa*, and Group 3 all specimens of var. *longifolia*. Values before the slash resulted from the analyses including var. *villosa* and after one excluding it. ANOSIM R-values are given above the diagonal and NPMANOVA F-values are given below the diagonal. Numbers in brackets are Bonferroni-corrected p-values. Significant differences are indicated by shadowed cells.

<table>
<thead>
<tr>
<th></th>
<th>Group 1</th>
<th>Group 2</th>
<th>Group 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1</td>
<td>0.8762 / 0.8771 [0.0003 / 0.0003]</td>
<td>0 / 1 [0.0003 / 0.0003]</td>
<td></td>
</tr>
<tr>
<td>Group 2</td>
<td>73.65 / 74.35 [0.0003 / 0.0003]</td>
<td>0.9045 / 0.9055 [0.0003 / 0.0003]</td>
<td></td>
</tr>
<tr>
<td>Group 3</td>
<td>235.5 / 235.5 [0.0003 / 0.0003]</td>
<td>84.47 / 85.58 [0.0003 / 0.0003]</td>
<td></td>
</tr>
</tbody>
</table>

Caribbean islands to central Brazil, but not the Amazonian region (var. *courbaril* and *stilbocarpa*), tropical ombrophilous lowland forests in central and eastern Amazon basin (var. *subsessilis*), or tropical ombrophilous submontane forests in the northeastern Brazilian state of Paraíba (var. *villosa*) (Figure 2).

The var. *altissima* (Group 1) constitutes a morphologically, geographically and ecologically consistent taxon. It is distinguished from the other varieties of *H. courbaril* complex by having smaller, falcate and acuminate leaflets, not exceeding 6.5 cm long on the flowering branches (Table 4). The leaflets of this taxon have a thinner texture than the remaining varieties, allowing its tertiary venation to appear as raised and reticulate veins. The flowers of var. *altissima* are also the smallest within the *H. courbaril* complex, measuring less than 15 mm long (Lee and Langenheim 1975, Pestana 2010). This variety is distributed in tropical ombrophilous lowland and submontane forests of the Atlantic Forest phytogeographical domain along the eastern coast of Brazil (Figure 2). It was cited for the southeastern Brazilian states of São Paulo and Rio de Janeiro (Mattos 1968, Lee and Langenheim 1975, Rizzini 1978, Pestana 2010), but we found specimens occurring northwards into Espírito Santo and Bahia states. It was originally described as *H. altissima* Ducke (Ducke 1935). Lee and Langenheim
Morphological analyses suggest a new taxonomic circumscription...

(1974: 448) considered that the foliar characters and relative flower size “do not appear sufficiently significant to warrant specific status for this taxon” and considered it as a variety of *H. courbaril*.

Var. *longifolia* (Group 3) comprises the morphs with largest leaflets. This variety also differs from the remaining groups by having oblong leaflets with an obtuse apex, clawed petals, and fruits compressed and enlarged toward the apex (Table 4). It occupies a very particular habitat in submontane tropical drought-deciduous thorny forests in the Caatinga and Cerrado phytogeographical domains that extend from western Bahia and Piauí states to the Araripe mountain range in the border area between the states of Ceará and Pernambuco (Figure 2). It was first proposed as a variety of *Hy- menaea splendida* Vogel. Lee and Langenheim (1974, 1975) considered *H. splendida* var. *splendida* to be synonymous with *H. courbaril* var. *stilbocarpa* and segregated *H. splendida* var. *longifolia* as another variety of *H. courbaril*.

In addition to morphological distinctions, ecological and geographical information can be used for interpreting the three groups recovered by UPGMA and PCA as separately evolving lineages that would be considered different species under the USC framework (see Taxonomy section). With respect to the variation in Group 2, studies of wider samplings, including especially the collections made after Lee and Langenheim’s (1975) work, shows that the diagnostic features used for defining the varieties included in Group 2 have more complex variations. Indumented leaflets, for example, the diagnostic character for var. *villosa*, can be found to varying degrees in var. *stilbocarpa*. A short stipe was used as a diagnostic feature of var. *subsessilis*, but this trait is variable and grades into var. *courbaril*. Finally, the distinctions between vars. *courbaril* and *stilbocarpa* rely on the types and colors of the calyx lobe indumentation, traits that tend to change during the duration of the flower. Thus, a more parsimonious way to treat the taxonomy of Group 2 is to consider it a variable species, without recognizing varieties.

### Table 4. Morphological comparison between *Hymenaea courbaril*, *H. altissima*, and *H. longifolia*.

<table>
<thead>
<tr>
<th></th>
<th><em>H. courbaril</em></th>
<th><em>H. altissima</em></th>
<th><em>H. longifolia</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Leaflet outline</td>
<td>Elliptic or ovate, not or slightly falcate</td>
<td>Falcately elliptic</td>
<td>Oblong or narrowly elliptic, not falcate</td>
</tr>
<tr>
<td>Inner margin of the leaflet</td>
<td>Straight or slightly concave</td>
<td>Strongly concave</td>
<td>Straight</td>
</tr>
<tr>
<td>Outer base of the leaflet</td>
<td>Slightly rounded</td>
<td>Truncate</td>
<td>Rounded</td>
</tr>
<tr>
<td>Apex</td>
<td>Acute, rarely obtuse</td>
<td>Abruptly acuminate</td>
<td>Obtuse, rarely rounded</td>
</tr>
<tr>
<td>Petiole length (cm)</td>
<td>1.1–1.7 (– 2)</td>
<td>1–1.3 (– 1.7)</td>
<td>2.6–3</td>
</tr>
<tr>
<td>Petiolule length (mm)</td>
<td>4–5</td>
<td>3–4 (– 5)</td>
<td>7–8</td>
</tr>
<tr>
<td>Leaflet size (cm)</td>
<td>6–9 (– 10.2) × 2.8–4 (– 5.1)</td>
<td>4–5.7 (– 6.4) × 2.0–2.5</td>
<td>10.1–12.5 (–15.4) × 4.5–6.5 (– 6.9)</td>
</tr>
<tr>
<td>Flower length (cm)</td>
<td>2.2–3.1</td>
<td>2–2.7</td>
<td>2.6–3.8</td>
</tr>
<tr>
<td>Fruit shape</td>
<td>Cylindrical, rounded in cross section</td>
<td>Cylindrical, rounded in cross section</td>
<td>Oblongoid, compressed in cross section</td>
</tr>
<tr>
<td>Seeds per fruit</td>
<td>Mostly &gt; 4</td>
<td>Mostly 1–2</td>
<td>Mostly &gt; 4</td>
</tr>
</tbody>
</table>
Taxonomic treatment


= Hymenaea resinifera Salisb., Prodr. 327. 1796, nom. nud.


= Hymenaea candolleana Kunth [spelled ‘candolliana’], Nov. Gen. and Sp. 6: 323, pl 556. 1824. Type: México, Acapulco, Bonpland 3875 (holotype P!).

= Hymenaea confertifolia Hayne, Getreue Darstell. Gew., 11, pl. 9. 1830. Lectotype (designated here): Sellow s.n., s.d. (lectotype W 48857!).


= Hymenaea splendidula Vogel, Linnaea 11: 409. 1837. Type: Sellow 1025 (holotype B†).


= Inga megacarpa M.E. Jones, Contr. West Bot. 15: 140. 1929. Type: México, Nayarit, M.E. Jones 23012 (holotype RSA!, isotypes F!, GH!, NY!).

= Hymenaea courbaril var. villosa Y.T. Lee and Andrade-Lima, J. Arnold Arbor. 55: 446. 1974, syn. nov. Type: Brasil, Paraíba, Paquevira de Natuba, Lee and Andrade-Lima 83 (holotype UC!).

Description. Large trees, to 30 m tall. Petiole 1.1–1.7 (–2) cm long; petiolule 4–5 mm long; leaflets 6–9 (–10.2) × 2.8–4 (–5.1) cm, elliptic, rarely ovate, straight or slightly falcate due to central vein with an angle c. 10°, apex acute or obtuse, rarely acuminate, base acute along the inner margin and rounded along the outer margin, distance from the inner margin to central vein 7–10 mm in the basal region. Flower buds 2.1–2.5 cm long; flower 2.2–3.1 cm long; hypanthium 7–15 mm long; petals 1.1–2.0 cm long. Fruit cylindrical, of uniform width, apex mostly rounded and apiculate.
Morphological analyses suggest a new taxonomic circumscription...

Hymenaea courbaril is defined here more narrowly than the circumscription adopted by Lee and Langenheim (1975), as we are proposing the exclusion of the varieties altissima and longifolia and their recognition as distinct species. In this narrower sense, H. courbaril is characterized by leaflets with slightly convex inner margins and midrib slightly arched, resulting in an elliptic or ovate outline, not or only slightly falcate, apex mostly acute or obtuse (Figure 3). It presents flowers larger than the other species of the H. courbaril complex, and cylindrical fruits mostly with more than five seeds (Table 4). It has a wide geographical range, mostly in different nuclei of Seasonally Dry Forests in Mexico, Central America, Caribbean, northern South America, Bolivia, and Brazil (from the Amazon region to Paraná State in the south).

Nomenclatural problems with several names associated with H. courbaril were not amended by Lee and Langenheim (1975) when they undertook their taxonomic revision of the genus. No formal type for H. stilbocarpa was cited by Lee and Langenheim (1975), who stated that no specimen was cited in the original description of Hayne (1830). However, Hayne (1830) based H. stilbocarpa on material collected by Martius, citing “Wäscht in Brasilien in Wäldern der Provinzen S. Paulo, Minas Geraes und Bahia (Martius)”. We found a specimen in the M herbarium collected by Martius in the Brazilian state of São Paulo (Santana farm) with an attached label indicating that this plant was distributed over the “Prov. Rio de Jº., S. Paulo, Minas Geraes, Bahia”, which probably served as the original material for Hayne’s description of H. stilbocarpa. Thus, we are lectotypifying this species with Martius’ specimen held in M under the barcode number M-0215314.

Hymenaea confertifolia Hayne was based on material collected by Sellow and Olfers in Brazil (“Wächst in Brasilien (Olfers u. Sellow)”; Hayne 1830: table 9). Lee and Langenheim (1975: 88) stated that this name was based on Sellow 1025 and that the “holotype” in

Figure 3. Leaf morphology of the species recognized in the Hymenaea courbaril complex: A Hymenaea longifolia B Hymenaea courbaril C Hymenaea altissima. Scale bar = 1 cm.
the B herbarium was destroyed. However, as Hayne (1830) did not refer to one particular specimen, all materials collected by Sellow or Olfers that can be linked with *H. confertifolia* should be considered syntypes. No such specimens can be found in the B herbarium, and were probably destroyed. A duplicate from B collected by Sellow was found in W annotated as *Hymenaea confertifolia*, and is designated here as the lectotype of this name.

*Hymenaea retusa* Willd. ex Hayne was published as a homotypic synonym of *H. candelleana* (Hayne 1830). It is thus illegitimate under the Article 52 of the International Code of Botanical Nomenclature (McNeill et al. 2011).

*Hymenaea courbaril* var. *obtusifolia* Ducke was published based on a tree cultivated in the Pará Botanical Garden (now Museu Paraense Emílio Goeldi, Belém, state of Pará, Brazil). Lee and Langenheim (1975) misinterpreted this as evidence that the holotype was the specimen in the herbarium of this museum (MG). However, Ducke (1925: 47) explicitly stated that “specimina florifera and fructus in herb. Jard. Bot. Rio n. 16.906”, and thus the RB material should be considered as the holotype of this name.

Ducke (1925: 265) did not cite any specimen when describing *H. courbaril* var. *subsessilis* Ducke. Lee and Langenheim (1975) did not explicitly designate a type for this variety, but chose a “representative specimen” collected “in the same general area believed to be the type locality”. We are accepting it as an inferential typification, and the status of this material should be a neotype since no other specimen was refereed in the protologue.


**Description.** Large trees, up to 38 m tall. Petiole 1–1.3 (–1.7) cm long; petiolule 3–4 (–5) mm long; leaflets 4–5.7 (–6.4) × 2–2.5 cm, oval, narrow-elliptic or elliptic, strongly falcate, due to central vein with an angle c. 35°, apex acuminate, rarely acute, base acute along the inner margin and truncate along the outer margin, distance from the inner margin to central vein 4–5 mm in the basal region. Flower buds 1.5–2.5 cm long; flower 2–2.7 cm long; hypanthium 7–12 mm long; petals 1.2–1.6 cm long. Fruit cylindrical, of uniform width, apex mostly rounded and apiculate.

*Hymenaea altissima* shows the smallest leaflets and flowers in the species group related to *H. courbaril* (Table 4). Additionally, its leaflets are strongly falcate due to the strongly curved and displaced midvein, with an abruptly acuminate apex. The base of the leaflet is acute along the inner margin and truncate along the outer margin (Figure 3). The fruits are mostly cylindrical, as in *H. courbaril*, but usually shorter than those of this species (4–5 vs. 8–20 cm long) because they have only one or two seeds, while in *H. courbaril* they usually have six or more seeds. This species is restricted to
the coastal rain forests of eastern Brazil, from southern Bahia State to São Paulo and Paraná states.

This species was described by Ducke (1935) as *H. altissima*. Lee and Langenheim (1974) treated it as a variety of *H. courbaril*. Later, these latter authors highlighted the differences between this taxon and the other varieties as having smaller and falcate leaflets and smaller flowers (Lee and Langenheim 1975: 86), but kept it as a variety of *H. courbaril*, a position also adopted by Pestana (2010). Our results indicated that the leaf traits of *H. altissima* are clearly distinct of those of *H. courbaril* and *H. longifolia*. These results, together with the distinctive flower and fruit traits and the coherent distribution and habitat data, all give support to the original view of Ducke (1935) in considering it a different species.

Lee and Langenheim (1975: 84) stated that the lectotype of *H. altissima* was designated by Egler (1963) as the specimen “A. Ducke RB 30306”. However, Egler (1963: 51) simply indicated that the type should be “Type: RB 23.306”. We could not track any specimens of *Hymenaea* in the RB herbarium with the numbers RB 30306 or RB 23306. We encountered, however, a specimen annotated by Ducke as *H. altissima* that was collected in the state of “Rio de Janeiro, Avelar” by G. M. Nunes in 1925 and could be linked to the protologue of *H. altissima* (Ducke 1935). This specimen is numbered RB 20306, and we are assuming that both Egler (1963) and Lee and Langenheim (1975) erroneously noted the RB number when referring to the lectotype of *H. altissima*.

*Hymenaea longifolia* (Benth.) I.M. Souza, Funch & L.P. Queiroz, comb. et stat. nov.
urn:lsid:ipni.org:names:77140226-1


**Description.** Medium-sized trees c. 5–12 (–18) m tall. Petiole 2.6–3 cm long; petiolule 7–8 mm long; leaflets 10.1–12.5 (–15.4) × 4.5–6.5 (–6.9) cm, elliptic, narrowly-elliptic, or oblong, not falcate, due to central vein with an angle c. 24°, apex obtuse or acute, base acute or rounded along the inner margin and rounded or slightly rounded along the outer margin, distance from the inner margin to central vein 10–11 mm in the basal region. Flower buds 2.2–3.3 cm long; flower 2.6–3.8 cm long; hypanthium 9–14 mm long; petals 1.3–2.1 cm long. Fruit oblongoid, compressed, wider at distal region, apex slightly truncate and apiculate.

*Hymenaea longifolia*, as circumscribed here, is characterized by large leaflets, both longer and wider than those of the related species *H. courbaril* and *H. altissima*, with longer petioles and petiolules (Table 4). The leaflets are elliptic or oblong with obtuse (rarely acute) apices and rounded bases along the outer margin (Figure 3). This species occurs in Seasonally Dry Forests within the Caatinga and Cerrado phytogeographical domains in northeastern Brazil, from Ceará State to the northern portion of Bahia State.
It was first described by Bentham (1870) as *H. splendida* var. *longifolia*, and considered related to *H. courbaril* and *H. stilbocarpa* because of the glabrous leaflets, but differing from them by having larger leaflets. Lee and Langenheim (1975) treated all these taxa under a more widely circumscribed *H. courbaril*, a position not supported by the results presented here. Lee and Langenheim (1975: 86) misinterpreted the specimen *Blanchet 3135* (R) as the holotype of *H. splendida* var. *longifolia*. However, when describing this variety, Bentham (1870: 236) cited two syntypes, *Blanchet 3135* (“ad Villa da Barra prov. Bahia”) and *Gardner 1938* (“Serra da Araripé, prov. Ceará”). Thus, the material cited by Lee and Langenheim (1975) as the holotype should be considered as a lectotype.

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Appendix 1


*Hymenaea courbaril var. villosa*—BRAZIL, Paraíba, Near Paquivera de Netuba, Y-T. Lee 82 (NY).