Resolving phylogenetic relationships and species delimitations in closely related gymnosperms using high-throughput NGS, Sanger sequencing and morphology

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Abstract Plastid genomes have been widely applied to elucidate plant evolution at higher taxonomic levels, but have rarely been considered useful for addressing close relationships. Here, we resolve the phylogeny and taxonomy of the Chinese lianoid Gnetum clade (Gnetales), using high throughput and Sanger sequencing techniques and studies of plant morphology. Despite previous efforts, relationships among taxa and the taxonomy within the clade have remained unclear. We generated 11 plastid genomes representing one arborescent and four lianoid species. Phylogenetic analyses were conducted using (a) the entire plastid genomes and (b) the protein-coding genes only. Sequence divergence among the lianoid species was substantial, with 9345 variable sites. Four variable regions were identified, targeted and sequenced for an additional 54 specimens and analyzed together with one nuclear ribosomal marker. Results from the phylogenetic analyses corroborate G. parvifolium as sister to the remaining lianoid species and support the presence of at least five additional species in the Chinese lianoid clade: G. catasphaericum, G. formosum, G. luofuense, G. montanum and G. pendulum. Following morphological investigations, G. giganteum and G. gracilipes are included in and synonymized with G. pendulum. Gnetum hainanense is included in and synonymized with G. luofuense. Two names, G. indicum and G. cleistostachyum, remain questionable. A taxonomic revision and a key to Chinese lianoid Gnetum are presented. Internal nodes in the Chinese lianoid Gnetum clade are from the Miocene and onwards and coincide with the expansion of tropical to subtropical forests in South China, which may have facilitated speciation in the clade.

Keywords Historical diversification • Next-generation sequencing • Phylogeny • Plant taxonomy • Plastid genomes

Introduction

In recent years, next-generation sequencing (NGS) has revolutionized systematic biology and molecular ecology. Compared to traditional Sanger sequencing techniques, NGS allows us to generate orders of magnitude of more informative characters in a both cost- and time-efficient way (Davey et al. 2011; Ekblom and Galindo 2011; Straub et al. 2012). As a consequence, NGS has allowed for the description of a rapidly increasing number of nuclear, mitochondrial and plastid genomes (Rounsley et al. 2009; Deschamps and Campbell 2010) and is set to contribute to a deeper understanding of both phylogeny and evolution in many lineages. Among the three types of genomes, data of entire plastid genomes have been widely used to seek for markers for the purpose of DNA barcoding (Hollingsworth et al. 2011; Nock et al. 2011), to investigate population genetics, and in phylogeographic studies (Powell et al. 2011).
purposes (Yao and Lin 2005; Yao et al. 2006; Liu et al. 2010), for production of daily necessities (Markgraf 1951) and as food and drinks (Fu et al. 1999a). Other representatives of more than one species (Fu et al. 1999a). Other representatives of more than one species (Fu et al. 1999a). The present study focuses on Chinese lianoid species of Gnetum (clade O sensu, Hou et al. 2015). Markgraf, who monographed the genus in 1930, recognized one arborescent taxon (G. gnemon L.) and two lianoid taxa from China (G. montanum Markgr. and G. montanum f. parvifolium (Warb.) Markgr.). Later, several new taxa were described (Markgraf 1930; Cheng 1964; Cheng et al. 1975; Shao 1994; Fu et al. 1999b), and at least 11 Chinese lianoid taxa have been proposed and discussed in the past. However, the taxonomic identity and delimitation of some of these taxa are unclear, partly as a result of insufficient knowledge of their genotypic and phenotypic variation, but also as a result of formal problems in their original protologues. Markgraf (1930), for example, listed no less than 43 synonyms in the original protologue of G. montanum, a heterogeneous collection that probably includes representatives of more than one species (Fu et al. 1999a). Other names (taxa), such as G. cleistostachyum C.Y.Cheng, G. pendulum f. intermedium C.Y.Cheng, were incorrectly described. Consequently, the names are formally invalid, and if they truly point at taxonomically identifiable units (species), they should be validly published and described. Relationships among the lianoid taxa are also unclear.

Previous phylogenetic studies based on molecular data support that two arborescent species, G. gnemon and G. costatum K.Schum., comprise the sister clade to the remaining Asian (lianoid) Gnetum (Hou et al. 2015). Among the Asian lianoid Gnetum, Chinese species constitute a strongly supported clade (Won and Renner 2003, 2005a, 2006; Hou et al. 2015), but relationships within this clade remain poorly supported, partly due to restricted sampling in previous phylogenetic analyses, but also because of limited sequence variation in the utilized molecular markers. Another problem may be misidentified specimens, resulting in contradictory taxonomic conclusions in the past.

Here, we aim to resolve the relationships and taxonomy of the Chinese lianoid clade of Gnetum using a combination of molecular and morphological data. We have developed new markers with appropriate amount of variation by generating complete plastid genomes of representative species. Developed markers are subsequently sequenced for additional specimens, and relationships of the included specimens are inferred by conducting phylogenetic analyses using all generated sequence data. Results, in combination with morphological investigations, are used to revise the taxonomy of the group. We also assess node ages in the clade and discuss the underlying evolutionary history of Chinese lianoid Gnetum.

Materials and methods

Taxon sampling

Silica material for DNA extraction was collected during field trips to the southern provinces of China (Guangdong, Guangxi, Yunnan and Hong Kong). In addition, material from herbaria in Belgium (BR), Scotland (E), China (HITBC, PE), Indonesia (KRB), USA (MO), Sweden (S) and the Netherlands (U) were studied. Altogether representatives of nine lianoid species (G. catasphaericum H.Shao, G. formosum Markgr., G. giganteum H.Shao, G. gracilipes C.Y.Cheng, G. hainanense C.Y.Cheng, G. costatum K.Schum., comprise the sister clade to the remaining Asian (lianoid) Gnetum. Among the Asian lianoid Gnetum, Chinese species constitute a strongly supported clade (Won and Renner 2003, 2005a, 2006; Hou et al. 2015), but relationships within this clade remain poorly supported, partly due to restricted sampling in previous phylogenetic analyses, but also because of limited sequence variation in the utilized molecular markers. Another problem may be misidentified specimens, resulting in contradictory taxonomic conclusions in the past.

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representing nine Chinese lianoid species, 15 representing remaining Asian taxa and one specimen representing the African species *G. africanum* Welw. Specimens representing *G. cleistostachyum* and *G. indicum* were not available for molecular studies.

**Plastid genome assembly, annotation and alignment**

Total genomic DNA of *Gnetum* was extracted from silica-treated leaves according to the cetyltrimethylammonium bromide (CTAB) method (Doyle 1991) and purified using a QIAquick PCR Purification Kit (Qiagen, Sweden). To comply with the DNA quantity sample requirement of the Illumina platform, silica material of each specimen was sampled twice and manually merged for subsequent DNA library construction. High-throughput sequencing was performed at Science for Life Laboratory (Stockholm, Sweden) following the manufacturer’s instructions for the Illumina HiSeq 2500 platform (Illumina, San Diego, USA). Single raw reads from Illumina sequencing were set in pairs, and low-quality nucleotides were removed with the error probability limit 0.05 in Geneious version 8.0.5.


**Table 1** Primer sequences and sources

| DNA region   | Primer name | Primer sequence 5′–3′ |
|--------------|-------------|-----------------------|
| ITS forwarda | 18SF        | GAA CCT TAT CGT TTA GAG GAA GG |
| ITS reversedb | 1270R      | TGA TTC ACG GGA TTC TCC A |
| matK forwardb | 7262F      | TAC CAG GGA GCT TAG TCC CAA |
| matK reversedb | 7983R      | GGA TTC GAA CCC GGA ACT CTT |
| rps12 forwardb | 101997F    | CAG TTC ACG GGG TAA TCC ACA |
| rps12 reversedb | 102820R    | CAC GTG CGA TCC TAA GGG GAG |
| rpoC1 forwardb | 71673F    | CAA GAG TGG TCC CGC GTC ACT |
| rpoC1 reversedb | 72942R    | TCT CAA TGA ACC GCA CGT GTA |
| trnF-trnV forwardb | 86084F  | CGC AGG GCT ATA GCT CAG TC |
| trnF-trnV reversedb | 87369R | CTA TCC CCA AGC CCG GTT TT |

a Rydin et al. (2004)  
b This study

(Kearse et al. 2012). Trimmed paired reads were mapped onto the plastid genome of *G. parvifolium* (GenBank ID, NC_011942) as the reference for five iterations with medium to low sensitivity in the Geneious mapper (installed in the same software). Eleven draft sequences of plastid genomes were generated, and matched raw reads for each specimen were saved. De novo assembly of matched raw reads was performed for each specimen using assembler Velvet version 1.2.10 (Zerbino and Birney 2008) with raw reads was performed for each specimen using assembler Velvet version 1.2.10 (Zerbino and Birney 2008) with the K-mer value optimized to 99 using Velvet Optimizer (installed in the same software). To evaluate accuracy, generated de novo contigs of each specimen were aligned with the draft plastid genomes from mapping. The quality of draft genomes were assessed and manually adjusted to fill up gaps and unclear nucleotides where necessary. Complete plastid genome sequences were annotated using DOGMA (Wyman et al. 2004) with the default settings. The start and stop codons of protein-coding genes were inferred by comparison with the known plastid genomes of the Gnetales. Annotated tRNA was further assessed using mVISTA (Mayor et al. 2000) by comparison and aligned plastid genomes with annotations were visualized and manually adjusted in the multiple alignments, and aligned plastid genomes with annotations were visualized using mVISTA (Maya et al. 2000) by comparison with the reference plastid genome.

**Pairwise genome comparisons, marker designing and PCR amplification**

To detect divergence hotspots in the plastid genome, the alignment of 11 complete plastid genomes was skimmed using Geneious variant/SNP finder with the minimum variant frequency set to 0.1 and maximum variant *P* value set to 10−5. Divergence hotspots consist of nucleotide substitutions and indels (insertions and deletions), which are marked onto the consensus of the alignment. With regard to sequence variations revealed in mVISTA and detected using variants/SNPs finder, four plastid markers, i.e., *matK, rpoC1, psbB-rps12* and *trnF-trnV*, were chosen to address species relationships of Chinese lianoid *Gnetum*. We made initial attempts to amplify three additional markers (*ycf1*, *trnP*, *ycf1, trnK-trnT*) but encountered problems and these markers were not further used. It is probably due to large numbers of sequence repeats in these regions of the plastid genomes of *Gnetum*. Complementing the four plastid markers, also the nuclear ribosomal marker ITS1 (including partial 5.8S) was utilized, a marker that has been used to resolve species relationships in *Gnetum* previously (e.g., Won and Renner 2005b). All primer information is listed in Table 1. PCR conditions for nrITS regions were set to an initial denaturation at 97 °C for 1 min; followed by 40 cycles of 97 °C for 10 s, 55 °C for 30 s and 72 °C for 20 s (4 s were added at each cycle); and followed by a final extension at 72 °C for 7 min. For amplification of plastid markers, the initial denaturation was set to 95 °C for 2 min; followed by 37 cycles of 95 °C for 30 s, 50 °C for 45 s and 72 °C for 2 min; and followed by a final extension at 72 °C for 7 min. PCR products were purified using Multiscreen PCR purification kits (Millipore, Billerica, MA) and sequenced at Macrogen Europe (Netherlands). Sequences from Sanger sequencing were assembled using the Staden Package version 1.6 (Staden 1996) and aligned using Clustal W version 1.81 (Thompson et al. 1994). Multiple alignments for each marker were visualized and manually adjusted in Mesquite version 2.75 (Maddison and Maddison 2011).

**Phylogenetic analyses of NGS data**

Phylogenetic analyses of the NGS data were conducted using two datasets: dataset (1) comprising the 11 complete
plastid genomes, and dataset (2) comprising a concatenation of all the protein-coding genes from the 11 plastid genomes (Table 2, Online Resource 2). Data in each dataset were treated as a single partition and analyzed using both maximum likelihood and a Bayesian inference approach. The single partition was tested with 22 candidate substitution models using jModeltest version 2.1.3 (Darriba et al. 2012). Best-fitting model was selected according to the Akaike information criterion (AIC), and a general time reversible substitution model with substitution rates drawn from a gamma distribution (GTR + Γ) was selected. Maximum likelihood analyses were conducted using GTR + Γ model implemented in RAxML version 7.2.8 (Stamatakis 2014). The analyses were conducted with 1000 replicates of rapid bootstrapping using a parsimony tree as a starting tree. Two specimens of *Gnetum gnemon* were used as outgroup, based on the results in previous phylogenetic studies (Won and Renner 2006; Hou et al. 2015). Bayesian analyses were conducted using MrBayes version 3.2.3 (Ronquist et al. 2012) via the online service of Cyber-infrastructure for Phylogenetic Research (CIPRES, Miller et al. 2010). Two parallel runs were set, each with four chains and using GTR + Γ as substitution model. The analyses were run for six million generations with a sampling frequency of 1000 generations. Convergence of the two MCMC runs was assessed using software AWTY (Nylander et al. 2008) and Tracer version 1.6 (Rambaut and Drummond 2003). Sampled trees were summarized in a majority rule consensus tree with the initial 600 trees removed as burn-in.

**Phylogenetic analyses of Sanger sequencing data**

Phylogenetic analyses were also conducted on a 66-taxon dataset generated from Sanger sequencing. Two different datasets were applied: dataset 3) a two-partitioned dataset of nrITS and the combined four plastid markers (*matK*, *rpoC1*, *psbB-rps12* and *trnF-trnV*) and dataset 4) a one-partitioned dataset of the combined four plastid markers (thus excluding nrITS) (Table 2, Online Resource 2). The best-fitting substitution models were selected according to the AIC criterion in jModeltest. Selected substitution model for the nuclear ribosomal data was GTR + Γ and for the combined plastid dataset a symmetrical model with gamma distribution (SYM + Γ). The reconstruction of phylogeny based on datasets 3 and 4 was conducted using maximum likelihood analyses with 1000 rapid bootstrap replicates in RAxML. Since the SYM + Γ model is not available in RAxML, the more complex substitution model GTR + Γ was set for the combined plastid dataset. In addition, datasets 3 and 4 were analyzed using Bayesian inference of phylogeny and substitution models GTR + Γ for the nrITS partition and SYM + Γ for the combined plastid markers. MCMC runs were set to six million generations, and generated trees were sampled every 1000 generations. AWTY and Tracer were applied to confirm the convergence of the two MCMC runs. Phylogenies were generated by summarizing the sampled trees in a majority rule consensus tree, and the initial 600 trees were deleted as burn-in. Supported topologies from the maximum likelihood and the Bayesian inference analyses of datasets 3 and 4 were compared to assess incongruence between the nrITS marker and the combined plastid markers.

**Estimation of divergence times**

Divergence times of clades in the Chinese lianoid *Gnetum* clade were estimated based on the two-partitioned dataset of five markers (dataset 3, Table 2) in a Bayesian framework using BEAST version 1.8 (Drummond et al. 2012) via CIPRES. Since the SYM + Γ model is not available in BEAST, the more complex substitution model GTR + Γ was set for both partitions. Four alternative settings were tested: (a) a strict clock model with a pure birth process as tree prior (Yule 1925; Gernhard 2008); (b) a strict clock model with a birth–death speciation process as the tree prior (Kendall 1948; Stadler 2009); (c) an uncorrelated lognormal clock model with a pure birth process; and (d) an uncorrelated lognormal clock with a birth–death speciation process. The optimal setting was selected using a posterior simulation-based analog of the Akaike information criterion (AICM, ...
Raftery et al. 2006; Baele et al. 2012) implemented in Tracer. Two independent MCMC runs were performed and run for 60 million generations with the sampling frequency of 20,000. A random tree was selected as the starting tree in all analyses. Nodes were calibrated to absolute time using normally distributed age priors, with monophyly enforced, based on results in Hou et al. (2015; Table 3). Separate runs excluding the data were also run, and the resulting effective prior distributions were assessed and compared to those specified. Effective sampling sizes for each Bayesian parameter were assessed in Tracer. In each run, the initial 300 trees were discarded as burn-in. The remaining sampled trees from two independent runs were combined in LogCombiner version 1.8 included in the BEAST package. The ultrametric tree was summarized with median node heights and maximum clade credibility from the sampled trees using TreeAnnotator version 1.8 included in the BEAST package.

### Morphological studies

Morphological studies of the Chinese lianoid species were performed using the 11 currently accepted names of species occurring in China and adjacent regions as a framework (Table 3). Forty-one herbarium sheets of type material were studied (of G. cleistostachyum, G. gracilipes, G. hainanense, G. luofuense and G. pendulum in PE and of G. catasphaericum and G. giganteum in PEM) (herbarium acronyms following Thiers 2016). Type material (40 specimens) of the remaining species was reviewed through Global Plants on JSTOR (http://plants.jstor.org/) as well as from requested images from the type localities. In addition, 126 herbarium sheets of non-type material were studied during visits in several additional herbaria in China and elsewhere (A, B, E, HK, HITBC, IBK, KUN, P, PE, PEM, NAS and S). Species determinations of non-type material were verified using available information (protologues, keys and comparison with type material). The gross morphology of specimens was studied by eye and using a dissecting microscope. Most species were also studied in their natural environments in the field. The type localities of G. giganteum and G. catasphaericum in Guangxi were visited. Localities near the type localities of G. gracilipes, G. hainanense and G. luofuense were also visited. Twenty-one characters were defined, seven from vegetative plant parts (characters 1–7), six from male reproductive parts (characters 8–13) and eight from female reproductive parts (characters 14–21). The state of each character was noted for each specimen, and data were summarized for each of the 11 currently accepted names (Table 3). Based on the results, species boundaries were subsequently assessed and taxonomic revisions were made accordingly.

### Results

#### Plastid genome assembly and sequence variation detection

The eleven specimens of *Gnetum*, for which the entire plastid genome was sequenced, generated between 13,547,250 and 77,910,472 paired-end reads with the average read length of 126 bp. A total of 190,899 to 2,003,005 paired-end reads were mapped onto the reference plastid genome, resulting in the average coverage number of 298–2156. Numbers of contigs assembled de novo varied from four to nine with the maximum length 29,004 to 65,957 bp. After manual modification, 11 plastid genomes of *Gnetum* were completed, with the length between 114,238 bp (*G. gnemon*) and 114,983 bp (*G. parvifolium*) (Online Resource 1). Plastid genomes of Chinese lianoid *Gnetum* consist of 115 genes (Online Resource 3), of which 66 were protein-coding genes, eight were ribosomal RNA (rRNA) genes and 40 were transfer RNA (tRNA) genes. In addition, sequence variations across the five *Gnetum* plastid genomes were revealed in the identity map (Fig. 2). A total number of 9345 variable sites were detected, consisting of 5600 polymorphisms (60 %) and 3745 insertions and/or deletions (40 %). Protein-coding regions revealed a conserved pattern in the alignment with the notable exception of *matK*, *ycf1* and *ycf2* where sequence variations existed. Non-protein-coding regions, consisting of intergeneric regions, introns, rRNA and tRNA, exhibited higher variability in the alignment, especially the intergeneric regions *psbD-trnT*, *trnT-trnK*, *trnN-ycf1*, *ycf1-trnP*, *trnH-trnL*, *rpl36-rps11*, *psaC-trnN*, *psbB-rps12* and *trnF-trnV*.

#### Phylogenetic analyses of NGS data

An alignment comprising 66 protein-coding genes (dataset 2; 64,150 aligned sites) was extracted from the full alignment of entire plastid genomes (dataset 1; 116,201 aligned sites). Phylogenetic analyses of the two datasets revealed highly congruent results, with regard to topology and statistical support, in both the maximum likelihood analysis and the Bayesian analysis. Detailed results from the analyses of dataset 1 are reported in Fig. 3. Bootstrap values higher than 70 and posterior probabilities larger than 0.95 are considered supported (Alfaro et al. 2003). Specimens that represent *G. parvifolium* comprise a clade [clade 2, BS (bootstrap support) = 100, PP (posterior probability) = 1.0]. Within *G. parvifolium*, specimen C50 is sister to the remaining two specimens (clade 3, BS = 100, PP = 1.0). *Gnetum parvifolium* is sister to all the remaining Chinese lianoid specimens (clade 5, BS = 100,
Table 3 Results of morphological studies of the 11 currently accepted species of the Chinese lianoid clade of *Gnetum*  

| Characters                        | Species                                      | Plant height (m) | Stem surface                                      | Petiole length (mm) | Leaf shape | Leaf size (mm) | Leaf surface | Leaf apex and base | Male branching pattern | Length of male spike (mm) | Number of male spike collars | Number of sterile units on each collar | Number of male reproductive units | Basal hairs of male spike | Female spike branching pattern | Number of fertile units on each collar | Number of involucre collars | Length between female collars (mm) | Seed shape |
|----------------------------------|----------------------------------------------|------------------|--------------------------------------------------|--------------------|------------|----------------|--------------|------------------|------------------------|-------------------------------|-------------------------------|-----------------------------------|-------------------------------|--------------------------|-------------------------------|-------------------------------|-----------------------------|-------------------------------|-----------------------------|
|                                  | *G. parvifolium* (Warb.) W.C. Cheng          | Liana, to 12     | Bark greyish brown, lenticels conspicuous       | 5–7\(^b\)         | Elliptic to narrowly oblong | 40–110 × 20–40\(^b\) | 5–8 lateral veins, inconspicuous on the adaxial surface | Apex acute to obtuse, base cuneate or subrounded | Simple or once branched | 8–15                      | 9–11\(^b\)                      | 8–12                         | 60–80\(^b\)                      | Few                        | Once branched                | 5–8                        | 6–11\(^b\)                  | 6–9\(^b\)                     | Elliptic, ellipsoid          |
|                                  | *G. formosum* Markgr.                        | Liana, to 25     | Bark black or dark brown, lenticels conspicuous | 9–10\(^b\)        | Elliptic to narrowly oblong | 110–140 × 40–70\(^b\) | 5–8 lateral veins, inconspicuous on the adaxial surface | Apex shortly acuminate to acute, base subacute | Simple or once branched | --                        | --                           | --                           | --                             | --                         | Once branched                | 6–11\(^b\)                  | 6–9\(^b\)                     | 6–9\(^b\)                     | Narrowly oblong, fusiform    |
|                                  | *G. indicum* (Lour.) Merr.\(^a\)           | --               | --                                               | 8–10\(^b\)        | Elliptic to oblong ovate | 60–120 × 40–60\(^b\) | 5–8 lateral veins, conspicuous on the adaxial surface | Apex shortly acuminate to obtuse, base cuneate | Simple or once branched | 20–30\(^b\)              | 12–15\(^b\)                   | 12–18\(^b\)                  | 60–80\(^b\)                      | Dense                       | Once or twice branched        | 9–13\(^b\)                  | 6–10\(^b\)                   | 6–9\(^b\)                     | Broadly ellipsoid to cylindrical |
|                                  | *G. luofuense* C.Y. Chenng ex L.K. Fu, Y.F. Yu & M.G. Gilbert | --               | --                                               | --                | Elliptic to oblong ovate | 45–160 × 30–85\(^b\) | 5–8 lateral veins, conspicuous on the adaxial surface | Apex shortly acuminate to obtuse, base cuneate to rounded | --                       | 20–30\(^b\)              | 12–15\(^b\)                   | 12–18\(^b\)                  | 60–80\(^b\)                      | Dense                       | Once branched                | 9–13\(^b\)                  | 8–13\(^b\)                   | 6–8\(^b\)                     | Broadly ellipsoid to cylindrical |
|                                  | *G. hainanense* C.Y. Cheng                   | --               | --                                               | --                | --                                               | 80–150 × 30–75\(^b\) | 5–8 lateral veins, conspicuous on the adaxial surface | Apex acuminate to obtuse, base cuneate to rounded | --                       | --                        | --                           | --                           | --                             | Few                         | Simple or once branched\(^a\) | --                           | 5–8\(^d\)                   | --                           | --                           | Ovulate                      |
|                                  | *G. cataphlaericum* H. Shao                 | Liana, to 10     | Bark black or dark brown, lenticels conspicuous | 8–13\(^b\)        | --                                               | 70–120 × 40–65\(^b\) | 5–8 lateral veins, conspicuous on the adaxial surface | Apex shortly acuminate to obtuse, base cuneate to rounded | --                       | 20–30\(^b\)              | 12–15\(^b\)                   | 12–18\(^b\)                  | 60–80\(^b\)                      | --                         | Simple or once branched\(^a\) | --                           | 5–8\(^d\)                   | --                           | --                           | Oblong, subglobeose           |
| Characters                  | Species                                      |
|----------------------------|----------------------------------------------|
|                            | *G. parvifolium* (Warb.) W.C.Cheng           |
|                            | *G. formosum* Markgr.                        |
|                            | *G. indicum* (Lour.) Merr.a                  |
|                            | *G. luofuense* C.Y.Cheng ex L.K.Fu, Y.F.Yu & M.G.Gilbert |
|                            | *G. hainanense* C.Y.Cheng                    |
|                            | *G. catasphaericum* H.Shao                   |
| Seed size (mm)             | 13–18 × 4–10b                                |
| Seed ornametation (silver scales) | Absentb                                     |
| Seed stipe                 | Absentb                                      |
|                            | Present; contracted, 2–5 mm when dryb         |
| Plant height (m)           | Liana, to 15                                 |
| Stem surface               | Bark black or dark brown, lenticels conspicuousb |
| Petiole length (mm)        | 9–26b                                        |
| Leaf shape                 | Elliptic to oblong                           |
| Leaf size (mm)             | 100–260 × 150–300                            |
| Leaf surface               | 8–12 lateral veins, conspicuous on the adaxial surfaceb |
| Leaf apex and base         | Apex obtuse, acute or acuminate, base cuneate to roundedb |
| Male branching pattern     | Once or twice branched                       |
| Length of male spike (mm)  | 20–35b                                       |
| Number of male spike collars | 16–25b                                      |
| Number of sterile units on each collar | 17–22b                                      |
| Number of male reproductive units | 20–45                                        |
| Basal hairs of male spike  | Dense                                        |
| Female spike branching pattern | Simple or once or twice branched             |
| Number of fertile units on each collar | 5–8                                           |

*Note: b indicates a specific characteristic.*
Clade 5 includes two specimens of *G. hainanense* (clade 4, BS = 100, PP = 1.0) sister to a group including specimens of *G. montanum* and *G. pendulum* (clade 7, BS = 100, PP = 1.0). *Gnetum montanum* (clade 6, BS = 100, PP = 1.0) is sister to *G. pendulum* (clade 8, BS = 98, PP = 1.0).

### Phylogenetic analyses of Sanger sequencing data

The 66-taxon dataset of Sanger sequencing data comprised a total of 4690 aligned sites (dataset 3), including 3675 aligned sites for the four plastid markers (dataset 4), and 1015 aligned sites for the nrITS. Detailed results from each analysis are reported in Fig. 4a, c (dataset 3) and Fig. 4b, d (dataset 4). There were no statistically supported conflicts between results from an analysis of the nrITS alone (not shown), and the concatenation of four plastid markers used in our study. Maximum likelihood and Bayesian inference analyses of datasets 3 and 4 revealed generally congruent results with regard to both topology and statistical support. These results are also highly congruent with those obtained from the analysis of NGS data (Fig. 3) and from results of the dating analysis (Fig. 5). Noteworthy, however, is that the clade comprising four *G. montanum* specimens yielded slightly lower statistic support when the nrITS marker was included in the Bayesian inference analyses (PP = 0.99 vs. 0.92, Fig. 4). The results considered below are those from the analyses of dataset 3 (including both nrITS and plastid markers).

Ten specimens that represent *G. parvifolium* form a clade (clade P, BS = 100, PP = 1.0). Relationships within the clade are resolved but generally poorly supported. *Gnetum parvifolium* is sister to the remaining taxa of the Chinese lianoid clade (clade R, BS = 100, PP = 1.0). Clade R comprises two specimens of *G. formosum*, which are excluded from clade S (BS = 100, PP = 1.0) comprising all remaining specimens. Clade S exhibits a trichotomy that consists of the clades V (BS = 99, PP = 1.0), W (BS = 97, PP = 1.0) and T (BS = 98, PP = 1.0). Clade V includes five specimens representing *G. catasphaericum*. Clade W includes seven specimens of *G. pendulum*, four specimens of *G. montanum*, three specimens of *G. giganteum* and two specimens of *G. gracilipes*. The clade comprising the four specimens of *G. montanum* was well supported in the Bayesian inference phylogeny (PP = 1.0) but not in the maximum likelihood analyses. Remaining relationships in clade W were either unresolved or poorly supported. Clade T comprises five specimens of *G. luofuense* and twelve specimens of *G. hainanense*. As with clade W, resolution in this clade is poor and the two taxon groups do not constitute separate clades.
Divergence time estimates

The birth–death prior combined with an uncorrelated (lognormal) relaxed clock (scenario d) had the best fit to the data as estimated from bootstrap simulation in AICM, and these settings were used in all subsequent divergence time analyses. Effective age priors were consistent with those specified. Divergence time estimates are shown in Fig. 5 in the form of a chronogram calibrated against the geological time scale (Gradstein et al. 2012), and in Table 4 where mean divergence and credibility intervals, as estimated by the 95 % HPD intervals, are given for major clades. The mean crown age of Chinese lianoid Gnetum was estimated to 21 Ma (HPD: 11–34 Ma corresponding to the Oligocene to early Miocene). The mean crown age of clade P (i.e., G. parvifolium) was 6 Ma (HPD: 2–11 corresponding to the late Miocene to Pliocene), which is younger than the mean crown age of clade R dated at 9 Ma (HPD: 4–17 early Miocene to Pliocene). Following a divergent event, clade Q (i.e., G. formosum) emerged at 2 Ma (HPD: 0–8, late Miocene to extant) and clade S (i.e., the remaining lianoid species) at 6 Ma (HPD: 3–10, late Miocene to Pliocene). The mean age of clade V (i.e., G. catasphaericum) was estimated at 4 Ma (HPD: 2–7 Ma; latest Miocene to Pliocene), which is older than the mean crown ages of clade T (i.e., G. luofuense) at 3 Ma (HPD: 1–5 Ma; Pliocene to Pleistocene) and clade W (i.e., G. pendulum and G. montanum) at 2 Ma (HPD: 1–4 Ma, Pliocene to Pleistocene).

Morphological studies and taxonomic conclusions

Based on the comparative studies of gross morphology, together with the results from our molecular analyses, the Chinese lianoid clade is proposed to comprise six species, i.e., G. catasphaericum, G. formosum, G. luofuense, G. montanum, G. parvifolium and G. pendulum. Gnetum hainanense C.Y.Cheng ex L.K.Fu, Y.F.Yu & M.G.Gilbert is included in and synonymized with G. luofuense C.Y.Cheng. Gnetum giganteum H.Shao and G. gracilipes C.Y.Cheng are included in and synonymized with G. pendulum C.Y.Cheng. New species circumscriptions and studied material are listed in Online Resource 4. Keys to the six recognized Chinese species (modified from Fu et al. 1999a), one based on male plants and one on female plants, are presented in the section of taxonomic treatment.

Fig. 2 Identity plots of plastid genome comparison of five species of Gnetum plotted against the reference genome of G. parvifolium (GenBank ID, NC_011942) in mVISTA. Pairwise comparisons and sequence identity are visualized on the y-axis from 50 to 100 % identity across the plastid genome. All genes are annotated and shown with directions (gray arrows), protein-coding regions (purple), tRNA and rRNA regions (light blue), intron and intergeneric regions (pink) and Sanger sequenced regions (green).

Fig. 3 Phylogenetic relationships of 11 Gnetum specimens based on maximum likelihood (ML) and Bayesian inference (BI) analyses of the entire plastid genomes (dataset 1). The tree was rooted using two specimens of G. gnemon as outgroup. Numbers shown represent support values obtained from 1000 rapid bootstrap replicates from the ML analyses (left) and Bayesian posterior probabilities (right).
Many specimens we developed four new plastid markers. Our morphological investigations corroborate the results from molecular data (with the possible exception of G. catasphaerica) and show for example that species with substantial intraspecific genetic divergence, like G.)

Discussion

According to the currently accepted classification, there are 11 lianoid species in China. However, based on our phylogenetic results and morphological studies, we conclude that there are only six lianoid species in the Chinese Gnetum clade (Fig. 5; Online Resource 4); G. parvifolium, G. catasphaerica, G. formosum, G. luofuense, G. montanum and G. pendulum. Remaining names are synonymous with other taxa. Distinct structural differences of plastid genomes were found among Chinese species of Gnetum. Interestingly, these differences are not only evident between the arborescent and the lianoid species, but also within the clade of Chinese lianoid species. Species delimitations are well resolved using the entire plastid genomes, but for more economic extraction of data from
**parvifolium**, also have distinct intraspecific morphological variation that appears correlated with geographic regions. Similarly, previously recognized species that cannot be separated based on our molecular data (e.g., *G. pendulum*, *G. giganteum* and *G. gracilipes*), also cannot be separated morphologically.

Estimated mean crown ages of species fall within the Pliocene to Pleistocene epochs. Only *G. parvifolium* may be slightly older (latest Miocene). The entire clade of Chinese lianoid *Gnetum* is from the Oligocene to Miocene and deep divergence events in the clade took place in the Miocene (Fig. 5). In the Miocene, tropical and subtropical areas expanded in South China, probably due to elevated mean temperature and annual precipitation (Yao et al. 2011), with expansion of broad-leaved forests as a consequence (Zhao et al. 2004). These transitions should have facilitated speciation and geographic expansion in the Chinese lianoid clade of *Gnetum* and may explain the largely overlapping geographic distribution of extant species. From the Pliocene, the climate gradually became cooler and drier (Ravelo et al. 2004; Xu et al. 2004; Koucies. From the Pliocene, the climate gradually became largely overlapping geographic distribution of extant species (Figs. 3, 4, 5). The analyses also indicate considerable sister to remaining species of the lianoid Chinese clade (Fig. 5). In the Miocene, tropical and subtropical areas expanded in South China, probably due to elevated mean temperature and annual precipitation (Yao et al. 2011), with expansion of broad-leaved forests as a consequence (Zhao et al. 2004). These transitions should have facilitated speciation and geographic expansion in the Chinese lianoid clade of *Gnetum* and may explain the largely overlapping geographic distribution of extant species. From the Pliocene, the climate gradually became cooler and drier (Ravelo et al. 2004; Xu et al. 2004; Koucies. From the Pliocene, the climate gradually became largely overlapping geographic distribution of extant species (Figs. 3, 4, 5). The analyses also indicate considerable sister to remaining species of the lianoid Chinese clade (Figs. 3, 4, 5).

**Phylogenetic relationships and species delimitations**

**Clade P**

Our phylogenetic analyses of both NGS data and Sanger sequence data strongly support that included specimens of *G. parvifolium* are monophyletic and that *G. parvifolium* is sister to remaining species of the lianoid Chinese clade (Figs. 3, 4, 5). The analyses also indicate considerable sequence divergence among the 10 included specimens of *G. parvifolium* (Fig. 4c, d), although there is only weak or no support for the relationships between the different specimens. The pattern seems to corroborate results in an earlier study where *G. parvifolium* was shown to possess high levels of genetic intraspecific diversity (Huang et al. 2010). Our morphological studies, based on material collected across the entire geographic distribution of the species (Fujian, Guangdong, Guangxi, Hainan, Hunan, Jiangxi and Hong Kong), further support the independent status of *G. parvifolium*. A combination of characters separates it from remaining Chinese lianoid *Gnetum* (Table 3), i.e., small leaf size and inconspicuous venation, short petioles (Fig. 1d) and sessile, ellipsoid seeds (Fig. 1g, i). Leaf and seed sizes of the specimens collected in Guangxi are, however, considerably larger than seen in specimens collected in Fujian, Guangdong and Hong Kong, indicating that intraspecific variation also of gross morphology occurs in *G. parvifolium*. This is further supported by variation in their average pollen size as reported by Yao et al. (2004). They documented *G. parvifolium* to have smaller average pollen size than *G. cleistostachyum*, *G. hainanense*, *G. luofuense*, *G. montanum* and *G. pendulum* (Yao et al. 2004).

**Clade Q**

*Gnetum formosum* was described by Markgraf (1930) on material from Vietnam, but the recognition of *G. formosum* and its separation from *G. parvifolium* has, at least in China, been unclear in previous taxonomic work. Markgraf (1930) considered *G. formosum* to be endemic to Vietnam. Also, Fu et al. (1999a) viewed *G. formosum* to be known with certainty only from Vietnam and did not recognize it as a Chinese species in their treatment of the Flora of China (Fu et al. 1999a). Female material, collected in Hainan (China) under the name *G. formosum*, was not accepted by Fu et al. (1999a) who considered the material not to differ significantly from plants of *G. parvifolium*. Wu and Chen (1978), in their Flora of China treatment, did not recognize *G. formosum* as a Chinese species either, and they even raised doubts about the Vietnamese material, questioning if not also this should be included under *G. parvifolium* (Wu and Chen 1978). In contrast, results from our molecular analyses reveal strong support for a separation of *G. formosum* and *G. parvifolium*, and the included specimens of *G. formosum* are resolved sister to the other species in clade R rather than falling with *G. parvifolium* (Figs. 4, 5).

This is true for the Chinese specimen from Hainan (*G. formosum* 14), as well as for a specimen from Laos (*G. formosum* 21). These results not only firmly establish the separation of *G. formosum* from *G. parvifolium*, but in addition expand the distribution of *G. formosum* to also include southern China (Hainan) as well as Laos (in the Vietnamese peninsula). Our investigation of gross morphology lend further support for these results; specimens of *G. formosum* differ from *G. parvifolium* by having a black or dark brown stem surface, larger leaves, longer petioles and narrowly oblong or fusiform seeds (Fig. 1j). They can also be distinguished from the other lianoid species by having 6–10 involucre collars placed in female spikes and sessile fusiform seeds (Table 3). Of potential concern is the lack of support for monophyly of *G. formosum* in our non-clock model-based molecular analyses (Fig. 4a, b). However, the two specimens are resolved together with moderate support in the relaxed clock model-based analysis (PP = 0.92; Fig. 5), they are with good support separated both from *G. parvifolium* and from remaining species in clade S in all analyses (Figs. 4, 5), and they are clearly distinguishable based on morphology (Table 3). We therefore consider it fully justified to treat *G. formosum* as
an independent species. Our result further indicates that its presence in both Laos and China (Hainan) has previously been overlooked.

**Clade V**

*Gnetum catasphaericum* was originally described by Shao (1994), and the type material was collected in Shangsi, Guangxi (China). However, the species status of *G. catasphaericum* has previously never been carefully assessed, probably due to difficulties of accessing plant material. Fu et al. (1999a), for example, indicated that they had not seen any material of this species. For the purpose of the present study, we collected new material of *G. catasphaericum* from three different localities in Guangxi (China), including the type locality. We also reassessed the identity of two previously collected herbarium specimens, one collected in Yunnan (China), and one collected in Vietnam, and the results of our phylogenetic analyses clearly show that all five specimens form a strongly supported monophyletic group (Fig. 4a, b). The analyses also indicate considerable sequence divergence among the investigated specimens (Fig. 4c, d). Unlike the situation in *G. parvifolium*, there is good support for the relationships among different specimens, and the relationships reflect geographical patterns. The three specimens from Guangxi (specimens 32, 33 and 44) are grouped together with the specimen from Yunnan (specimen 31) as sister. Sister to all the Chinese specimens is the specimen from Vietnam (specimen 24). Whether or not these patterns indicate the presence of cryptic and as yet undescribed species in this group should be investigated further, but this will require a more extensive sample of *G. catasphaericum* than included in the present analyses. Morphologically, *G. catasphaericum* can be distinguished from the other Chinese lianoid species by possessing relatively few female units on each collar (5–8) (Fig. 1f), fewer basal hairs and oblong, subglobose seeds with contracted seed stipes 2–6 mm long when dried (Fig. 1k; Table 3). In addition, intraspecific variation of reproductive characters is detected among the four specimens of *G. catasphaericum* used for our molecular studies. For example, the number of fertile ovules in specimen 32 is 5–6, which is fewer than 7–8 of specimens 44 collected in another site of Guangxi.

**Clade W**

The well-supported clade W comprises representatives of four previously recognized species, *G. pendulum*, *G. giganteum*, *G. gracilipes* and *G. montanum*. Our molecular analyses included four Chinese specimens of *G. montanum*, two from Guangxi (specimens 105, 106) and one from Yunnan (specimen 17), as well as a specimen from Thailand (specimen 29). They are always resolved as monophyletic, although only some analyses (Figs. 4b, 5) provide strong support for the *G. montanum* clade. *Gnetum montanum* was described by Markgraf (1930), and he listed no less than 43 different syntypes in the original protologue. The listed types indicate a broad geographical distribution of the species and include material collected in China (Guangxi, Yunnan), India (Assam, Sikkim), Burma, Thailand and Vietnam. However, Fu et al. (1999a) pointed at several problems regarding the application of the name *G. montanum*. For example, the 43 syntypes are

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Table 4 Results of the dating analysis (dataset 3, uncorrelated lognormal clock, birth-death tree prior)

| Node names | Prior type (distribution) | Monophyly enforced | Prior setting (CI 95 %) | Posterior height (95 % HPD) | Posterior probability (PP) |
|------------|--------------------------|--------------------|------------------------|-----------------------------|-----------------------------|
| Clade D (root) | Secondary (normal)* | Yes | 74 (57–91) | 70 (53–88) | 1 |
| Clade F | Secondary (normal)* | Yes | 65 (48–82) | 56 (43–69) | 1 |
| Clade H | Secondary (normal)* | Yes | 55 (38–72) | 48 (36–60) | 1 |
| Clade J | Secondary (normal)* | Yes | 36 (20–52) | 38 (27–49) | 1 |
| Clade K | Secondary (normal)* | Yes | 26 (13–39) | 29 (19–39) | 1 |
| Clade O | – | – | – | 21 (11–34) | 1 |
| Clade P | – | – | – | 6 (2–11) | 1 |
| Clade Q | – | – | – | 2 (0–8) | 0.92 |
| Clade R | – | – | – | 9 (4–17) | 1 |
| Clade S | – | – | – | 6 (3–10) | 1 |
| Clade T | – | – | – | 3 (1–5) | 1 |
| Clade V | – | – | – | 4 (2–7) | 1 |
| Clade W | – | – | – | 2 (1–4) | 1 |

* – Not applicable
* Based on results in Hou et al. (2015)
morphologically highly variable and include specimens not only of this taxon but also of *G. latifolium* Blume (Fu et al. 1999a). They also pointed out that the name *G. montanum* commonly has been used in SE China, especially in Hong Kong, for material of *G. luofuense*. Following the protologue as well as current usage of the name, Fu et al. (1999a) considered *G. montanum* as best characterized by having male involucral collars with fewer microsporangia and smaller stipitate seeds. They applied the name *G. montanum* accordingly but at the same time argued that this (current) application of the name should be conserved by selecting an appropriate lectotype (Fu et al. 1999a). This, however, they never did. Morphologically, the species is distinct as assessed by our observations, with more collars on the male spikes (Fig. 1c), fewer reproductive units on female spikes and relatively small stipitate seeds (Fig. 11) compared to other Chinese species. Based on this, and the strong support for monophyly of the species in at least some of our analyses of molecular data, we find it justified to recognize *G. montanum*. We also agree with Fu et al. (1999a) that lectotypification is warranted. Unfortunately, we have not been able to see any of the type specimens from China listed by Markgraf (1930), because they are located in herbaria outside of China not visited by us. However, the specimen from Thailand included in our molecular analyses was collected from the exact same area (Chiang Mai) as was Kerr n. 602, a syntype specimen listed by Markgraf (1930). We have therefore chosen Kerr 602 (K) as the designated lectotype for *G. montanum* (see nomenclature below), conserving the current application of the name as suggested by Fu et al. (1999a).

The remaining specimens of clade W represent three additional species that have been recognized in treatments of the Chinese flora: *G. pendulum, G. gracilipes* and *G. giganteum* (Cheng et al. 1975; Wu and Chen 1978; Fu et al. 1999a). The first two, *G. pendulum* and *G. gracilipes*, were originally described by Cheng et al. (1975) on collections from Yunnan (*G. pendulum*) and from Guangxi (*G. gracilipes*). However, the validity of the original diagnoses was questioned by Fu et al. (1999b) as Cheng et al. (1975) include no direct information on the species being described. They only indicate how some other species differ from it, and both were compared to species found well outside of China. *Gnetum pendulum* was compared to *G. oblongum* Markgr., a species described by Markgraf (1930) from the Chittagong Hill Tracts in Bangladesh, and *G. gracilipes* was compared to *G. arboreum* Foxw., a species that does not even have a lianoid habit and was described from the Philippines by Foxworthy (1911). Both comparisons are of questionable value, but *G. pendulum* and *G. gracilipes* have remained recognized in more recent work (Wu and Chen 1978; Fu et al. 1999a). One feature that *G. pendulum* (Fig. 1m) and *G. gracilipes* (Fig. 1n) seem to share is long-stipitate seeds, although Cheng et al. (1975) considered the character so variable in *G. pendulum* that they described different forms: *G. pendulum f. intermedium* C.Y. Cheng and *G. pendulum f. subsessile* C.Y. Cheng. The first name, *G. pendulum f. intermedium* is invalid as two different types (male and female) were designated in the original protologue (Cheng et al. 1975).

Fu et al. (1999a) considered *G. pendulum* as closely related to, and *G. gracilipes* as showing many similarities to, *G. latifolium*. This is a putatively widespread and highly variable species described by Blume (1834) from Java, Indonesia, for which a large number of different forms and varieties have been described. Our analyses clearly indicate a different pattern (Fig. 4a, b). The included specimen of *G. latifolium* is resolved together with other species from Southeast Asia in clade K, nowhere close to the *G. pendulum* and *G. gracilipes* specimens. Also in previous work, including additional specimens of *G. latifolium* (Won and Renner 2006; Hou et al. 2015) are specimens of *G. latifolium* monophyletic and included in the equivalent of our clade K, i.e., clearly separated from the Chinese lianoid clade.

The third species included in clade W is *G. giganteum*, described by Shao (1994) based on type material collected in Guangxi. It was described to differ from other species by having large, sessile and broadly ellipsoid seeds, and female involucral collars separated by up to 3 cm when mature (Shao 1994). The status also of this species has not previously been carefully assessed. There are very few collections of *G. giganteum* in Chinese herbaria, and Fu et al. (1999a), for example, indicated that they had not seen any material of this species.

Based on our morphological work, it is clear that the three taxonomic entities *G. pendulum, G. gracilipes* and *G. giganteum* share characters with overlapping states and are morphologically indistinguishable (Table 3). Further, although the representing specimens are unresolved within clade W in our non-clock model-based molecular analyses (Fig. 4a, b), they form a clade in the relaxed clock model-based analysis (Fig. 5). We therefore conclude that it is relevant to assign material previously referred to any of the three names *G. pendulum, G. gracilipes* and *G. giganteum* to the same single species (i.e., *G. pendulum*). From a sequence divergence point of view, there is a striking pattern with almost no variation at all in the entire clade W (Fig. 4c, d). Strictly speaking, no conclusions can be drawn based on this almost complete absence of molecular variability, but a comparison with the divergence patterns seen in the *G. parvifolium* and *G. catasphaericum* clades indicates that other processes are active in clade W (and clade T) than in clades P and V regarding the exchange of genetic material among individuals.
Clade T

Also, another clade with almost no sequence divergence among included specimens even though the specimens represent more than one taxon, is identified in our analyses. The taxa concerned are G. hainanense and G. luofuense and specimens representing these two taxa are all resolved in the well-supported clade T (Figs. 4, 5). Morphologically, G. hainanense and G. luofuense can be distinguished from the other Chinese lianoid species by possessing longer male spikes (Fig. 1d), more fertile units on each collar (Fig. 1e), seed ornamentation (Fig. 1h) and short-stipitate seeds (Fig. 1o, p) (Table 3). Both species were originally described by Cheng et al. (1975) based on material from China, G. hainanense from Hainan and G. luofuense from Guangdong. Their description of G. hainanense pointed, however, at both a female and a male type specimen and the name was for this reason invalid. Fu et al. (1999b) noticed this problem and supplied a valid description of G. hainanense based on the female type specimen from Cheng et al. (1975). They described G. hainanense as possessing apparently larger seeds with smooth outer seed coats (Fig. 1o) compared with smaller wrinkled seeds of G. luofuense (Fig. 1p) (Fu et al. 1999b). However, a separation of G. hainanense and G. luofuense is not supported in the analyses conducted here. Our molecular analyses indicate almost no sequence divergence whatsoever between the included specimens (Fig. 4c, d), and also the results of our morphological studies reveal no differentiation in the features investigated (Table 3). These include the number of lateral veins on the leaves, and the size and shape of the seeds, features that were used in the original protologue of G. hainanense to separate it from G. luofuense (Fu et al. 1999b). Yao et al. (2004) investigated the exine ultrastructure and size of pollen grains from Chinese species of Gnetum, and they too found no differentiation between G. hainanense and G. luofuense. We therefore consider G. hainanense synonymous to the older name G. luofuense.

Gnetum indicum (Lour.) Merr

The type material of G. indicum was collected in Vietnam in the early 1770s. The name is possibly synonymous with G. parvifolium, but the taxonomic history of G. indicum is complex and the name has been applied in several ways in the past, often in the sense of G. montanum according to Fu et al. (1999a). Markgraf (1930) argues that the type material is vegetative and difficult to interpret, and we largely agree although there actually is a young seed on the sheet. The seed is cracked and important information on its original shape is not preserved. Therefore, and in the absence of DNA data from the type, we find it difficult to assess the affinity of the type material of G. indicum with certainty. Based on available information, it may represent G. parvifolium, G. formosum or a species of its own.

Gnetum cleistostachyum C.Y.Cheng

This name is invalid because both a male and a female specimen were indicated as the type specimen, and the protologue only states a few sentences on how the material differs from G. pendulum (Cheng et al. 1975). More importantly, however, very little material of this species has ever been collected and Fu et al. (1999b) considered it wise to postpone a validation of the name. Based on our morphological investigations, it seems possible that the type material of G. cleistostachyum in fact represents G. montanum. The “closed” nature of the reproductive spikes of G. cleistostachyum is most probably unimportant and a developmental difference; all known spikes of G. cleistostachyum are at an early developmental stage. There are, however, one apparent difference between G. cleistostachyum and G. montanum (the number of male reproductive units on each collar), but in the absence of additional morphological data and/or DNA data from G. cleistostachyum, we refrain from assessing whether or not the material represents a unique species.

Taxonomic notes

Characters of the seed, such as size, shape, surface ornamentation and length of the stipe, have previously been considered useful in identifying species and forma of Chinese lianoid Gnetum (Cheng et al. 1975; Wu and Chen 1978; Fu et al. 1999a). Some seed characters are, however, variable within species and/or overlap between species. For example, the fleshy seed coat may be both coarsely reticulate-wrinkled and smooth in G. luofuense; the basal part of seeds contract into a “false” stipe when dried in G. luofuense and G. catasphaericum, which then becomes indistinguishable from the “true” seed stipe of G. montanum formed during the development; and seed ornamentation of silver scales are often present in both G. luofuense and G. pendulum. Further, we find details of hairs that surround female reproductive spikes uninformative and not useful for species identification in the Chinese lianoid clade; hairs are usually brown, short and inconspicuous in these species. The new key to female specimens presented here (see below) therefore makes use of other reproductive characters (e.g., spike length, length between involucre collars and number of fertile units on each collar, see Table 3). The key to male specimens is based on characters such as spike length, number of sterile units on each collar and amount of basal hairs. Vegetative characters such as measurements

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of leaves size, leaf surface pattern, color of leaves when dried, are broadly applied for distinguishing species among South American species (Markgraf 1965) and Malaysian species (Markgraf 1951), but are not useful to identify Chinese lianoid Gnetum because the characters are largely overlapping (see Table 3). Nevertheless, such characters can be useful in combination with reproductive characters.

**Taxonomic treatment**

*Gnetum parvifolium* (Warb.) W.C.Cheng, Acta Phytotax. Sin. 9: 386. 1964. ≡ *Gnetum scandens* Roxb. var. *parvifolium* Warb., Monsunia 1: 196. 1900. ≡ *Gnetum montanum* Markgr. f. *parvifolium* (Warb.) Markgr., Bull. Jard. Bot. Buitenzorg Ser. III, 10(4): 466. 1930. ≡ *Thoa catasphaerica* (Warb.) Doweld, Turczaninowia 3(4): 32. 2000. — TYPE: China, Guangxi, Shangsi, 8 Sep 1987, Shao 87053

= *Gnetum gracilipes* C.Y.Cheng Acta Phytotax. Sin. 13(4): 88. 1975. — TYPE: China, Guangxi, Sup Man Ta Shan, Liang 69874 (lectotype designated here, or perhaps holotype: PE barcode PE00002919!; isolectotypes: P barcode P00722134 [photo!], P barcode P00722135 [photo!], B barcode B100001469 [photo!], IBK barcode IBK00041278 [photo!]).

*Gnetum luofuense* C.Y.Cheng, Acta Phytotax. Sin. 13(4): 89. 1975. — TYPE: China, Guangdong, Luofushan, 10 Aug 1930, Chen 41527 (lectotype designated here, or perhaps holotype: PE 00002913!; isolectotype: MO barcode MO954421 [photo!]).

*Gnetum hainanensis* C.Y.Cheng ex L.K.Fu, Y.F.Yu & M.G.Gilbert, Novon 9(2): 187, 1999 ≡ *Thoa hainanensis* (C.Y.Cheng ex L.K.Fu, Y.F.Yu & M.G.Gilbert) Doweld, Turczaninowia 3(4): 33. 2000. — TYPE: China, Hainan Island, Hung Mo Shan, 22 Jul 1929, Tsang et Fung 554 (holotype: PE barcode PE00002917!; isotypes: A barcode A00022589 [photo!], K barcode K000454089 [photo!], PE barcode PE00002918).

**Key based on male plants**

1a. Shrubs or small trees; leaves yellowish when dried; sterile ovules and axis visible between involucral collars ........................................... G. gnemon

1b. Liana; leaves dark brown or black when dried; spikes condensed with axis invisible between involucral collars ........................................... 2

2a. Sterile ovules 7–12; surrounding basal hairs few…… 3

2b. Sterile ovules 13–22; surrounding basal hairs dense ........................................... 4

3a. Male spike 0.5–1.5 cm in length; inconspicuous lateral veins on the adaxial surface of leaves ........................................... G. parvifolium

3b. Male spike ca. 2 cm in length; apparent lateral veins on the adaxial surface of leaves ........................................... G. catasphaericum

4a. Male spike 1–1.5 cm in length; male spikes branching three to five times .................. G. pendulum

4b. Male spike 2–3.5 cm in length; male spike simple, or once or twice branching .................... 5

5a. Involucral collars 12–15; male reproductive units 60–80 ........................................... G. luofuense

5b. Involucral collars 16–25; male reproductive units 20–45 ........................................... G. montanum

**Key based on female plants**

1a. Shrubs or small trees; leaves yellowish when dried; involucral collars flat ..................... G. gnemon

1b. Liana; leaves dark brown or black when dried; involucral collars funnel-shaped .............. 2

1 The type sheets assigned in the protologue were not found while visiting the PEM herbarium.
2a. Seed sessile; inconspicuous lateral veins on the adaxial surface of leaves .............................................. 3

2b. Sterile ovules 13–22; surrounding basal hairs dense ................................................................. 4

3a. Seed 1.3–1.8 cm in length, 0.4–1 cm in width; leaf 4–11 cm in length, 2–4 cm in width …… G. parvifolium

3b. Seed 2–2.5 cm in length, 0.8–0.9 cm in width; leaf 2–2.5 cm in length, 0.5–0.9 cm in width …… G. formosum

4a. Fertile ovules 9–13; collars 11–25 mm apart ......... 5

4b. Fertile ovules 5–8; collars 6–10 mm apart ................................................................. 6

5a. Seeds 3–4 cm in length, 1.7–1.8 cm in width; seed stipe 8–30 mm long ……… G. pendulum

5b. Seeds 1.8–2.5 cm in length, 1.3–1.7 cm in width; seed stipe contracted into 2–5 mm long when mature …… G. luofuense

6a. Seeds oblong, subglobose; seed stipe contracted into 2–6 mm long when mature …… G. catasphaericum

6b. Seeds oblong, subglobose; seed stipe contracted into 2–6 mm long when mature .......... G. montanum

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Compliance with ethical standards

Conflict of interest We declare that we have no conflicts of interest.

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Information on Electronic Supplementary Material

Online Resource 1. Sequence information for high throughput sequencing (top): we list voucher information, collection places of each specimen, the sizes of assembled plastid genomes, number of matched reads and average coverage during mapping. Information on Sanger sequencing (followed), includes voucher information, collection places of each specimen and GenBank accessions.

Online Resource 2. We list alignments of the four datasets (dataset 1–4) used in the present study.

Online Resource 3. We list 115 genes identified in the plastid genomes of Chinese lianoid Gnetum categorized according to gene function. Other relevant information, such as pseudogenes, number of introns and copies in IRs are also listed.

Online Resource 4. Here we list all lianoid species of Gnetum present in China. The descriptions are based on original descriptions of species, notes in Flora of China (Fu et al. 1999a), and results of the present study including investigations of type material of all listed species. One additional species of Gnetum is present in China, the arborescent G. genmon. It is variable with many described subspecies and variants and will be dealt with in a forthcoming study.

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