DNA Barcode Authentication and Library Development for the Wood of Six Commercial *Pterocarpus* Species: the Critical Role of Xylarium Specimens

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DNA barcoding has been proposed as a useful tool for forensic wood identification and development of a reliable DNA reference library is an essential first step. Xylaria (wood collections) are potentially enormous data repositories if DNA information could be extracted from wood specimens. In this study, 31 xylarium wood specimens and 8 leaf specimens of six important commercial species of *Pterocarpus* were selected to investigate the reliability of DNA barcodes for authentication at the species level and to determine the feasibility of building wood DNA barcode reference libraries from xylarium specimens. Four DNA barcodes (ITS2, *matK*, *ndhF-rpl32* and *rbcL*) and their combination were tested to evaluate their discrimination ability for *Pterocarpus* species with both TaxonDNA and tree-based analytical methods. The results indicated that the combination barcode of *matK* + *ndhF-rpl32* + ITS2 yielded the best discrimination for the *Pterocarpus* species studied. The mini-barcode *ndhF-rpl32* (167–173 bps) performed well distinguishing *P. santalinus* from its wood anatomically inseparable species *P. tinctorius*. Results from this study verified not only the feasibility of building DNA barcode libraries using xylarium wood specimens, but the importance of using wood rather than leaves as the source tissue, when wood is the botanical material to be identified.

Increasing concern about and demand for biodiversity conservation world-wide and substantial declines in biological diversity at various spatial, temporal and biological scales are driving the need for species identification for forensics. For forest systems, illegal logging and the illegal timber trade are major problems domestically and internationally, threatening not just individual species, but entire ecosystems. Illegal logging is both a consumer- and producer-country driven phenomenon, and international efforts to respond to the problem consist of the enactment of laws to prohibit or limit the trade in illegally sourced timber. Broad international trade restrictions are imposed primarily through the Convention on International Trade in Endangered Species of Wild Fauna and Flora (CITES), which lists species in three appendices according to the degree of protection required.

In recent years, several consumer countries and regions have also taken action to reduce the trade in forest products derived from illegally logged sources. The United States amended the Lacey Act in 2008, which makes it unlawful to import into the United States any plant (or plant product) that was illegally harvested. In Australia, the Illegal Logging Prohibition Act (2012) was enacted to restrict the trade of illegally logged timber. The European Union’s E.U. Timber Regulation (EUTR) came into effect in 2013, prohibiting illegally sourced...
timber and timber products in the EU market. These legislative actions and subsequent enforcement of these laws demonstrate the urgent global attention on forest species protection. Enforcement actions to date in the United States have largely focused on ebony and rosewood from Madagascar (Dept. of Justice 2012), Eurasian hardwoods (Dept. of Justice 2016), and tropical woods from Peru (Dept. of Justice 2017) indicating that wood forensic methods, including DNA barcoding reference libraries, for valuable woods from anywhere in the world could play a critical role in law enforcement for forest protection.

Pterocarpus Jacq., is a pantropical genus in the family Leguminosae, containing approx. 70 species. The timber of Pterocarpus is globally valued for its beauty, wood quality, medicinal properties and even valuable bioactive compounds. This high value and increase in demand for the timber has led to illegal and excessive logging resulting in threat to wild Pterocarpus populations. In 1995, CITES listed P. santalinus under Appendix II to regulate trade in logs, wood chips and unprocessed broken material. P. erinaceus was added to Appendix II at the 17th Meeting of the CITES in 2017. Concurrently, the International Union for Conservation of Nature (IUCN) also listed P. santalinus and P. zenkeri as endangered, P. indicus and P. marsupium as vulnerable, and P. angolensis as near threatened. In China, the species P. indicus was listed in the second-class category of the National List of Local Protected Flora issued by the Chinese Government in 1999. Among the Pterocarpus species, P. santalinus, endemic to the Southern parts of Eastern Ghats of India especially in Andhra Pradesh, is known for its characteristic color, texture, quality and the medical value of its timber, which makes it of particular economic importance, especially in China. In recent years, the wood from P. tincto- rius (non-CITES) mostly distributed in Central and Southern Africa, appeared on the international lumber market as a substitute for P. santalinus (CITES App. II). Its macroscopic wood properties, e.g. color, grain, density, and its anatomical structure are very similar to that of P. santalinus. Due to the great difference in economic value, P. tinctorius has often been treated as an adulterant of P. santalinus in the timber market. Thus, developing accurate species-level identification for Pterocarpus wood is significant for natural resource protection.

Traditional wood identification relies on diagnostic anatomical features, either macroscopic or microscopic, but rarely can provide a precise discrimination of wood at the species level, which limits the enforcement of CITES regulations and related laws. Moreover, traditional wood identification requires expert taxonomic and anatomical knowledge that takes years to gain. To overcome such limitations, recent advances in molecular diagnostic tools for plants have the capacity to improve upon traditional methods of species identification.

For the last decade, DNA barcoding has been the subject of extensive research and application as an accurate and convenient tool for species identification. DNA barcoding is a genetic approach based on a short DNA sequence from a standard part of a genome – in animals this is typically a region of the cytochrome c oxidase subunit 1 (CO1) mitochondrial region. In plants, mitochondrial mutation rates are too slow for species-level identification, so plastid and nuclear regions are typically chosen as barcodes. The Consortium for the Barcode of Life (CBOL) proposed a combination of both the chloroplast DNA (cpDNA) ribulose-bisphosphate carboxylase (rbcL) gene and maturase K (matK) genes as the core DNA barcodes for plants. Chen et al. proposed that the ITS2 region could be potentially used as a standard DNA barcode, especially for identifying medicinal plants and their closely related species. Additionally, the ndhF-rpl32 intergenic spacer in the short single copy region of the chloroplast genome, which was noted as highly variable by Timme et al., has also been used for phylogenetic studies. DNA barcodes are established tools for identifying herbal medicinal materials, in quality control, and in forensic science. Additionally, a number of studies relying upon DNA barcoding have verified the utility, potential and importance in the wood market. These legislative actions and subsequent enforcement of these laws demonstrate the urgent global attention on forest species protection. Enforcement actions to date in the United States have largely focused on ebony and rosewood from Madagascar (Dept. of Justice 2012), Eurasian hardwoods (Dept. of Justice 2016), and tropical woods from Peru (Dept. of Justice 2017) indicating that wood forensic methods, including DNA barcoding reference libraries, for valuable woods from anywhere in the world could play a critical role in law enforcement for forest protection.

Results

Wood anatomical separation of P. santalinus and P. tinctorius. The wood anatomical features of P. santalinus and P. tinctorius are almost identical (Fig. 2). Wood diffuse-porous; vessels exclusively solitary, occasionally with radial multiples of 2 to 3, and often filled with dark gums; intervessel pits alternate; vessel-ray pits...
Barcode Recovery and Sequence Characteristics. The recovery success rate was the highest for ndhF-rpl32 (90%), followed by matK (82%) and rbcL (70%), while ITS2 exhibited the lowest rate (67%). In total, 123 sequences generated in this work were deposited to GenBank (Accessions, ITS2: KY829137-KY829162; matK: KY829163-KY829195; ndhF-rpl32:KY829196-KY829232; rbcL: KY829233-KY829259) (Supplementary Table S1).

The features of the four DNA barcodes were shown in Table 1. The length of the aligned rbcL sequences was 350 bp with 47 variable sites and 15 informative sites. The aligned matK sequence was 239 bp long, with 10 variable sites and 10 informative sites. In the ITS2, the sequence was 234 bp in length, with 75 variable sites, 69 informative sites and 16 indels. For the sequence of ndhF-rpl32, the aligned length was 173 bp, with 14 variable sites, 12 informative sites and six indels. Among the four DNA barcodes, ITS2 had the highest proportion of variable (32.05%) and informative (29.49%) sites, followed by rbcL (13.43% and 4.29%) and ndhF-rpl32 (8.09% and 6.94%), with matK showing the lowest values (4.18% and 4.18%).

The pairwise intraspecific distances for the barcodes ranged from a minimum value of 0.0000 for all four barcodes to a maximum value of 0.0962 (ITS2), and the mean intraspecific distances ranged from 0.0026 (matK) to 0.0200 (ITS2). The pairwise interspecific distances for the barcodes ranged from 0.0000 for all four barcodes to 0.1681 (ITS2), and the mean interspecific distances ranged from 0.0073 (rbcL) to 0.0800 (ITS2) (Table 2). ITS2 shows the highest mean intra- and inter-specific distances. The pairwise intraspecific distances for combined barcodes ranged from 0.0000 for all combinations to 0.0638 (ndhF-rpl32 + rbcL), and the mean intraspecific distances ranged from 0.0027 (matK + ndhF-rpl32 + ITS2) to 0.0091 (ndhF-rpl32 + rbcL). The pairwise interspecific distances for combined barcodes ranged from 0.0000 (matK + ndhF-rpl32, matK + rbcL and ndhF-rpl32 + rbcL) to 0.0954 (rbcL + ITS2) and the mean interspecific distances ranged from 0.0133 (matK + rbcL) to 0.0524 (ndhF-rpl32 + ITS2).

DNA Barcoding Gap Assessment. Barcoding gaps, the absence of overlapping regions between intra- and interspecific distances, were evaluated by the results of the distribution graph obtained in the “pairwise summary” function in TaxonDNA (Supplementary Figure S1). In the study, no single- or multi-barcodes exhibited clear barcoding gaps; all barcodes overlapped between the intra- and interspecific distances. However, the mean interspecific divergence was higher than that of the corresponding intraspecific variation for each of the barcodes (Table 2). Among the single barcodes, ITS2 had the highest variation in interspecific divergence compared to the range of intraspecific distances (Table 2). When barcodes were individually analyzed, ITS2 presented the best barcode gap performance, with 69.6% of pairwise interspecific distances greater than 0.05 and 95.9% of pairwise intraspecific distances lower than 0.05. Conversely, unsatisfactory results were observed for matK, ndhF-rpl32 and rbcL separately, with almost total overlap of intra- and interspecific variation (Supplementary Figure S1) for each.

As for the barcode combinations, the best results were found for matK + ITS2 and matK + ndhF-rpl32 + ITS2, with 98.9% and 95.1% of pairwise interspecific distances greater than 0.05, respectively, and 91.8% of pairwise intraspecific distances lower than 0.05, both of which also outperformed any single barcode. All other barcode combinations showed clear overlap (Supplementary Figure S1).
Species Discrimination based on TaxonDNA and Tree-based Analysis. The parameters “best match” and “best close match” from Taxon DNA were used to analyze all sequences generated in this study as well as those downloaded from the GenBank database (Fig. 3). For single-locus barcodes, both the “best match” and “best close match” methods were applied to determine the closest match between the generated sequences and those in the GenBank database. The results showed that the performance of “best match” and “best close match” were comparable, with the “best close match” method yielding slightly better performance in terms of accuracy and specificity. The parameters derived from these two methods were then used to construct phylogenetic trees for the species under study, which further confirmed the accuracy of the TaxonDNA analysis.

Table 1. The characteristics of the four DNA barcode loci.

| DNA marker | Recovery rate (%) | Sequence length (bp) | Aligned sequence length (bp) | G+C ratio (%) | No. variable sites (%) | No. parsimony informative sites (%) | Indel length (bp) |
|------------|-------------------|----------------------|-----------------------------|---------------|-----------------------|-------------------------------------|------------------|
| ITS2       | 66.67             | 219–224              | 234                         | 66.1          | 75 (32.05)            | 69 (29.49)                         | 16               |
| matK       | 82.05             | 273                  | 239                         | 36.0          | 10 (4.18)             | 10 (4.18)                          | 0                |
| ndhF-rpl32 | 89.74             | 167–173              | 173                         | 26.6          | 14 (8.09)             | 12 (6.94)                          | 6                |
| rbcL       | 69.67             | 485                  | 350                         | 40.6          | 47 (13.43)            | 15 (4.29)                          | 0                |

Table 2. Genetic distance generated using Kimura 2-parameter model analysis for the candidate barcode loci and their combinations.

| Barcode loci and combinations | Intraspecific distance | Interspecific distance |
|------------------------------|------------------------|------------------------|
|                              | Minimum | Maximum | Mean | Minimum | Maximum | Mean |
| a) ITS2                      | 0.0000  | 0.0962  | 0.0200 | 0.0000  | 0.1681  | 0.0800 |
| b) matK                      | 0.0000  | 0.0293  | 0.0026 | 0.0000  | 0.0335  | 0.0099 |
| c) ndhF-rpl32                | 0.0000  | 0.0226  | 0.0045 | 0.0000  | 0.0292  | 0.0091 |
| d) rbcL                      | 0.0000  | 0.0914  | 0.0063 | 0.0000  | 0.0943  | 0.0073 |
| e) matK + ITS2               | 0.0000  | 0.0101  | 0.0030 | 0.0000  | 0.0671  | 0.0433 |
| f) matK + ndhF-rpl32         | 0.0000  | 0.0122  | 0.0015 | 0.0000  | 0.0146  | 0.0083 |
| g) matK + rbcL               | 0.0000  | 0.0578  | 0.0060 | 0.0000  | 0.0608  | 0.0133 |
| h) ndhF-rpl32 + ITS2         | 0.0000  | 0.0131  | 0.0035 | 0.0000  | 0.0824  | 0.0524 |
| i) ndhF-rpl32 + rbcL         | 0.0000  | 0.0638  | 0.0091 | 0.0000  | 0.0755  | 0.0173 |
| j) rbcL + ITS2               | 0.0000  | 0.0464  | 0.0067 | 0.0013  | 0.0954  | 0.0410 |
| k) matK + ndhF-rpl32 + ITS2  | 0.0000  | 0.0091  | 0.0027 | 0.0065  | 0.0582  | 0.0392 |
| l) matK + ndhF-rpl32 + rbcL  | 0.0000  | 0.0467  | 0.0058 | 0.0012  | 0.0551  | 0.0159 |
| m) matK + rbcL + ITS2        | 0.0000  | 0.0374  | 0.0053 | 0.0049  | 0.0739  | 0.0335 |
| n) ndhF-rpl32 + rbcL + ITS2  | 0.0000  | 0.0388  | 0.0059 | 0.0010  | 0.0787  | 0.0383 |
| o) matK + ndhF-rpl32 + rbcL + ITS2 | 0.0000 | 0.0327 | 0.0064 | 0.0050 | 0.0638 | 0.0340 |

Figure 2. Wood anatomical features of *P. santalinus* and *P. tinctorius*. (A,B and C) Transverse, radial, and tangential sections of *P. santalinus* wood, respectively. (D,E and F) Transverse, radial, and tangential sections of *P. tinctorius* wood, respectively. Scale bars, 200 μm (A and D) and 100 μm (B,C,E and F).
and “best close match” methods provided the similar species discrimination success rate. ITS2 showed the highest success rate (85.1%), followed by ndhF-rpl32 (20.0%), rbcL (18.2%), while matK exhibited the lowest rate (10.7%). The identification success rates for all barcode combinations were generally higher than those of the single barcodes. The highest success rate (100%) of barcode combinations based on the “best match” and “best close match” analysis was obtained by the two-barcode combination of matK + ITS2 and three-barcode combination of matK + ndhF-rpl32 + ITS2. The ndhF-rpl32 + rbcL combination exhibited the lowest performance for correct identification. All barcode combinations that included ITS2, i.e. matK + ITS2, ndhF-rpl32 + ITS2, rbcL + ITS2, matK + ndhF-rpl32 + rbcL + ITS2 and matK + ndhF-rpl32 + rbcL + ITS2, provided higher identification success rates than other chloroplast DNA barcode combinations (Fig. 3).

Bootstrap support for species-specific clusters based on unrooted neighbor-joining (NJ) trees for the four barcodes and their combinations were calculated (Supplementary Figure S2). When barcodes were individually analyzed, the highest species discrimination successes were obtained by ITS2 and rbcL (16.7%), whereas the barcodes matK and ndhF-rpl32 could not distinguish any Pterocarpus species (Supplementary Table S4). The mini-barcode ndhF-rpl32, 167–173 bps in length, can separate the two anatomically similar species, P. santalinus and P. tinctorius using neighbor-joining tree analysis (Fig. 4B). Furthermore, six continuous diagnostic characters (insertion/deletion) at nucleotide positions from 112 to 117 (TTATTA) were found within the ndhF-rpl32 region (Fig. 4C), which was a distinguishing feature based on the character-based approach. Discrimination of all six species using only one barcode was insufficient to provide an accurate resolution among the Pterocarpus species studied here. When combining two to four barcodes, the highest discrimination rate (100%) was obtained by matK + ndhF-rpl32 + ITS2 and matK + rbcL + ITS2 (Fig. 5). Moreover, the barcode combinations that included ITS2 yielded higher success rates than other chloroplast DNA barcode combinations (Supplementary Table S4).

Discussion

Assessment of DNA Barcodes for Pterocarpus. An ideal DNA barcode should be short making it easy for recovery, and have sufficient information to provide maximal species discrimination. While this is true for any barcode as a general principle, it is a key concern for barcodes for wood identification, because wood...
is a DNA-poor botanical material in the living tree, and the quality and quantity of DNA in wood degrades with industrial processing, necessitating barcodes known to be recoverable from dry wood. In these *Pterocarpus*, shorter amplicons showed a generally higher recovery rate than longer ones, with the shortest fragment *ndhF-rpl32* having the highest success rate, which is in line with several previous studies. We expect that the DNA in xylarium wood specimens is typically highly fragmented. Additionally, the nuclear ribosomal DNA region ITS2 yielded lower recovery success rate (67%) compared to the chloroplast DNA regions although it is present in multiple copies in the genome. In spite of some amplification disadvantages, ITS2 provided the best discrimination ability of *P. santalinus* and *P. tinctorius*. Figure 4. Analysis of discrimination ability of *P. santalinus* and *P. tinctorius* based on the specific mini-barcode *ndhF-rpl32*. (A) PCR amplification and sequencing success rate of the four DNA barcodes, (B) neighbor-joining tree constructed based on the barcode *ndhF-rpl32*, (C) variable sites of the barcode *ndhF-rpl32* between the two species.

Figure 5. Taxon identification tree constructed using neighbor-joining analysis of P-distance showing the best-performing barcode combination *matK + ndhF-rpl32 + ITS2*. Bootstrap values (>50%) are shown above the relevant branches. Photomacrographs (×16) of *Pterocarpus* xylarium specimens.
discrimination performance among the four barcodes. The superior identification power of nuclear DNA region ITS2 over plastid barcodes is also consistent with the results of other previous studies.15,16,19,33–37

Although the chloroplast DNA regions rbcL and matK were proposed as core barcodes for seed plants31, the two regions gave low species resolution for Pterocarpus in this study. Both rbcL and matK are widely used in phylogenetic analyses with over 130,000 sequences available in Genbank. Kress et al.30 showed that the rbcL sequence evolves slowly and this barcode has been recognized as the lowest divergence of studied plastid genes in flowering plants.30 Consequently, on average it is not likely to be useful for identification at the species level.15,31,38–40 It is reported that matK shows different discrimination success rates when it comes to different taxonomic groups (e.g. discriminating more than 90% of species in the Orchidaceae40) but less than 49% of species in the nutmeg family.40,42 Meanwhile, despite its power in phylogenetic studies of other species17,18, ndhF–rpl32 showed low resolution for distinguishing all six Pterocarpus species in this study.

No single barcode was found to be able to distinguish all six Pterocarpus species in this study. Overall, combined barcodes provided higher species resolution than any single barcode, which was consistent with previous studies.15,19,34 The CBOL Plant Working Group recommended the combination barcode of rbcL + matK as the core barcode for land plants. Yan et al.32 also demonstrated that the three barcode combination of ITS + psbA–trnH + matK could give better discrimination performance than single barcodes, and was the best choice for the genus of Rhododendron. In this study of Pterocarpus, the highest success rate of barcode combinations based on the “best match” and “best close match” analysis of TaxonDNA was obtained by matK + rbcL and matK + ndhF–rpl32 + ITS2. When the tree-based analyses (NJ) were conducted, the combination matK + rbcL + ndhF–rpl32 + ITS2 gave the best results. We conclude that the combination matK + ndhF–rpl32 + ITS using these two methods is the best combination DNA barcode to resolve six of Pterocarpus species (Figs 3 and 5).

Although the barcode matK individually or in combination with other chloroplast DNA barcodes yielded a low success rate for species discrimination, interestingly it has the ability to cluster studied Pterocarpus species according to their broad geographic origins (Fig. 6). We found that Asian and African species clustered together except for 1 or 2 samples of P. angolensis (Fig. 6). Here we suggest the two chloroplast locus combination of matK + ndhF–rpl32 as a potential barcode for geographic origin tracking of Pterocarpus species when the recovery success rate is considered. It has been reported that the chloroplast DNA barcodes that are variable enough to reveal geographic structure could be used to differentiate the origin of timber45–47. Additionally, Lee et al.12 also showed that the DNA barcode combination matK + trnL–trnF + ITS2 had the ability of geographic clustering for Aquilaria species.42

Species Discrimination between P. santalinus and P. tinctiorius based on the Special Mini-barcode. Inasmuch as P. santalinus and P. tinctiorius cannot be separated by wood anatomy but are mixed in trade, an effective method to separate these woods is critically needed. A single DNA barcode targeted to this question alone would be an effective tool, especially if the barcode were easily recovered from both species. The DNA mini-barcode ndhF–rpl32 could give good performance for distinguishing the two closely related Pterocarpus species.

DNA mini-barcodes, short DNA sequences of 100–250 bp, are suitable for species identification within a given taxonomic group of old herbarium/museum specimens when high-quality DNA is not available and seriously degraded DNA is retrieved.48,49 We suggest that the DNA mini-barcoding approach is suitable for species identification of woody tissues, especially in narrow cases to separate a small number of anatomically indistinguishable woods. In this study, the recovery success rate of ndhF–rpl32 was highest among the four DNA barcodes in the study (Fig. 4) and this parameter has been used as an important criterion to determine whether DNA could be effectively isolated from wood tissues.11,12,22,23,34 The reduced taxonomic discriminatory power of a mini-barcode compared to that of a full-length barcode and the taxon-specific nature of which mini-barcodes are most effective are the primary detriments of this approach. If for every group of taxa a new mini-barcode is needed, the basic principle of standardization is violated. Therefore, the choice of position of mini-barcodes from DNA genome is significant in their ability of discriminating species.46–50 A good DNA mini-barcode candidate should be of high PCR and sequencing success without much loss of species discrimination power, and as broadly applicable as possible.

Regarding the Utility of DNA Barcoding in the Conservation of and Controlled Trade in Pterocarpus Wood. Biodiversity conservation has rapidly become a focus of attention due to the sharp increase of global forest resources trade, over-exploitation and illegal logging activities. For forest protection and global trade monitoring, developing accurate species-level identification and geographic traceability for wood is a crucial and significant technical prerequisite. The application of DNA barcoding to identify the species and track the geographic origin of internationally traded timber has attracted increasing interest as a potential part of global systems to support sustainable forestry and especially to reduce the behaviors of illegal logging. In addition to this work, previous studies have reported the potential of DNA barcoding to support conservation efforts of wood species, e.g. Aquilaria1,12, Dalbergia31,32 and Populus1,2.

DNA barcoding can play an increased role in identification and conservation of Pterocarpus species, and of wood species worldwide. Availability of a reliable reference DNA barcode library remains the main obstacle of application of DNA barcoding for the next few years. Our study confirms that xylarium wood specimens are rich sources for reliable DNA sequence data. Xylarium wood specimens could certainly enhance the construction of global DNA barcode reference libraries to support species conservation worldwide, and thus continue to play a critical role as repositories of wood anatomical, chemical, and molecular information for the future.
Materials and Methods

Plant Materials. All wood specimens were taken from the xylarium (wood collections) of the Chinese Academy of Forestry (WOODPEDIA), the largest wood collection in China. A total of 39 specimens of 6 species of *Pterocarpus* were sampled. Four types of specimens, i.e., heartwood, sapwood, twig, and silica gel-dried leaf were collected in this study. 4–11 individuals per species were sampled. Details of the collected reference samples, including the location of vouchers, are listed in Supplementary Table S1.

Molecular Methods. Exposed surfaces of xylarium wood specimens were removed with a sterile scalpel to avoid external contamination. Each wood sample of 500 mg was frozen in liquid nitrogen and then ground into a fine powder in a 6770 Freezer/Mill (SpexSamplePrep, Metuchen, NJ, USA).

All DNA isolations were carried out under sterile conditions. DNA from the wood specimens was extracted following the DNeasy Plant Maxi Kit (Qiagen, Hilden, Germany) protocol, modified according to Jiao et al.11. For silica gel-dried leaves, DNA was isolated using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany) following the manufacturer’s recommendations.

PCR amplification was performed in a 30 μl reaction with 15 μl of TaKaRa Premix Ex Taq (containing 0.75 units of Ex Taq DNA polymerase, 2 mM of MgCl₂, and 200 μM of each dNTP), 0.2 μM of each primer and approx. 10 ng of template DNA. The amplification was conducted in a Veriti 96-Well Thermal Cycler (Applied Biosystems, Foster City, CA, USA). PCR Primers and PCR cycling conditions used in this study are listed in Supplementary Table S2. The PCR products were purified using a UNIQ-10 Spin Column DNA Gel Extraction Kit (Sangon, Shanghai, China) and sequenced in both directions with the same primers used for PCR on an ABI PRISM 3730xl (Applied Biosystems, Foster City, CA, USA).

In addition to the sequences generated in this work, we downloaded sequences (from loci ITS2, *matK*, *ndhF-rpl32* and *rbcL*) (Supplementary Table S3) for specimens of *Pterocarpus* from GenBank for analysis.

Light Microscopy. Sectioning blocks [10 mm (L) × 10 mm (R) × 10 mm (T)] were cut with razor blades and then soften at 60 °C for 48 hours. Thereafter, 15 μm thick transversal, radial and tangential sections were cut on a sliding microtome. Sections were stained with 1% aqueous safranin solution, rinsed, then mounted on glass slides and then observed under a light microscope (Olympus BX61, Japan).

Data Analysis. Raw sequences for each region were assembled and edited using ContigExpress in Vector NTI Advance v. 10.1 (Invitrogen InforMax, Frederick, MD, USA), saved in FASTA format and deposited to GenBank (Supplementary Table S1). The edited sequences were then aligned with Clustal X 1.81 followed by a manual
adjustment with BioEdit software. To assess the barcoding gap, the relative distribution of pairwise genetic distances was calculated using TaxonDNA under the K2P-corrected pairwise distance model.

To evaluate species discrimination success, two widely used methods, TaxonDNA and a neighbor-joining tree-based approach, were applied to the four single barcode and all their possible combinations. For the TaxonDNA analysis, we used the “best match” and the “best close match” functions in the software to test the species-level discrimination rates under the K2P-corrected distance model for each barcode singly and all possible combinations of barcodes. The “best close match” method required a threshold value which was calculated for each barcode from pairwise summary. All the results above the threshold were treated as “no match”. For the tree-based method, unrooted neighbor-joining (NJ) trees were constructed in MEGA 5 with pairwise deletion and the P-distance model. Only when all the conspecific individuals were clustered a single clade and at least one specimen in each clade was derived from a botanically vouchered collection was it considered a successful species discrimination.

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A.C.W., Y.Y. wrote the paper. All authors read and approved the manuscript. A.C.W. analyzed the data. L.J., M.Y., T.H., J.L., B.L., Y.Y. contributed reagents/materials/analysis tools. L.J., M.Y., Y.Y., L.J., M.Y., X.J. conceived and designed the study. L.J., M.Y., Y.Y., L.J., M.Y., X.J. performed the experiments. L.J., M.Y., Y.Y., L.J., M.Y., X.J. contributed reagents/materials/analysis tools. L.J., M.Y., Y.Y., L.J., M.Y., X.J. analyzed the data. L.J., M.Y., X.J. wrote the paper.

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Author Contributions
Y.Y., L.J., M.Y., X.J. conceived and designed the study. I.J., M.Y., T.H. performed the experiments. L.J., M.Y., Y.Y. analyzed the data. L.J., M.Y., T.H., J.L., B.L., Y.Y. contributed reagents/materials/analysis tools. L.J., M.Y., A.C.W., Y.Y. wrote the paper. All authors read and approved the manuscript.

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