Research Article

Stability and Accuracy Assessment of Identification of Traditional Chinese Materia Medica Using DNA Barcoding: A Case Study on Flos Lonicerae Japonicae

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DNA barcoding is a novel molecular identification method that aids in identifying traditional Chinese materia medica using traditional identification techniques. However, further study is needed to assess the stability and accuracy of DNA barcoding. Flos Lonicerae Japonicae, a typical medicinal flower, is widely used in China, Korea, and other Southeast Asian countries. However, Flos Lonicerae Japonicae and its closely related species have been misused and traded at varying for a wide range of prices. Therefore, Flos Lonicerae Japonicae must be accurately identified. In this study, the ITS2 and psbA-trnH regions were amplified by polymerase chain reaction (PCR). Sequence assembly was performed using CodonCode Aligner V 3.5.4. The intra- versus inter-specific variations were assessed using six metrics and “barcoding gaps.” Species identification was conducted using BLAST1 and neighbor-joining (NJ) trees. Results reveal that ITS2 and psbA-trnH exhibited an average intraspecific divergence of 0.001 and 0, respectively, as well as an average inter-specific divergence of 0.0331 and 0.0161. The identification efficiency of ITS2 and psbA-trnH evaluated using BLAST1 was 100%. Flos Lonicerae Japonicae was formed into one clade through the NJ trees. Therefore, Flos Lonicerae Japonicae can be stably and accurately identified through the ITS2 and psbA-trnH regions, respectively.

1. Introduction

Lonicera japonica Thunb. (Caprifoliaceae) is a commonly used herb that grows widely in China, Japan, Korea, and Southeast Asian countries [1]. The flower bud, called Flos Lonicerae Japonicae, is the most important part of the plant for medicinal use [2, 3], with more than 3000 years of medical history. Flos Lonicerae Japonicae was first reported in one of the earliest pharmacopoeias worldwide, Shen-Nong’s Herbals Classic [4]. Flos Lonicerae Japonicae is considered a herbal medicine with heat-clearing, detoxifying, and anti-inflammatory effects [5, 6]. In addition, extracts of Flos Lonicerae Japonicae have shown efficacy against various diseases such as H1N1 influenza [4, 7–9], hand-foot-and-mouth disease (HFMD) [4, 10], and severe acute respiratory syndromes, among others [11–13]. In recent years, Flos Lonicerae Japonicae has also become the main material of a Chinese “cooling” beverage [14]. Flos Lonicerae, which comes from L. macranthoides Hand-Mazz, L. fulvotomentosa Hsu et S.C. Cheng, L. confusa (Sweet) DC., and L. hypoglauca Miq., is another medicinal material from Lonicera [6]. More than 30% of the current traditional Chinese medicine prescriptions have recently used Flos Lonicerae Japonicae [4]. Consequently, Flos Lonicerae and other species of Lonicera,
such as L. japonica var. chinensis, L. similis, and L. acuminata, have been misused as Flos Lonicerae Japonicae because of high market demand for Flos Lonicerae Japonicae.

High-quality botanical materials are important for the safe and effective use of herbal products [15]. Several criteria and techniques have been used in the identification of Flos Lonicerae Japonicae to ensure its quality and therapeutic effects [16–19]. Li et al. introduced the morphologic and microstructural methods for identifying L. japonica and Lonicerae [20]. Despite its ability to preliminarily identify L. japonica, the method needs the integrity of the material structure. The geoherbalism of L. japonica has been identified by sequence divergence of the 5S-rRNA gene spacer region [21]. However, Flos Lonicerae Japonicae is difficult to identify according to morphology, microstructural methods, and certain molecular markers.

DNA barcoding can quickly and reliably identify species using a standard genetic sequence that can be amplified by universal primers from a small fragment of the genome [22–25]. Several DNA regions have been evaluated for potential application as DNA barcodes, such as matK, rbcL, psbA-trnH, and ITS2 [26–28]. In the recent years, the DNA barcoding technology has been increasingly used in several fields including species identification [29–33], biodiversity study [34, 35], misidentification detection [36], food safety testing [14], and tackling illegal trade of endangered species [37, 38]. Chen et al. suggested that ITS2 can be used as a novel DNA barcode for the identification of medicinal plants and their closely related species [27]. The potential identification ability of ITS2 in plants has been assessed by analyzing 50,790 plant ITS2 sequences [28]. The China Plant BOL Group has also shown that ITS2 can be used to identify seed plants [39]. ITS2 sequences are potential general phylogenetic markers and have been widely used for phylogenetic reconstructions at both the genus and the species levels [40, 41]. Previous studies used the psbA-trnH intergenic spacer region from plastid DNA to successfully number a relationship of several species identification cases [42–45]. Chen et al. recommended psbA-trnH as a complementary barcode to ITS2 for a broad series of plant taxa [27].

In our previous study, the genomic DNA extracted from the leaves of L. japonica was used for genomic identification. Sun et al. tested several potential DNA regions and proposed that psbA-trnH can be used for the identification of L. japonica and its closely related species [46]. In the present study, the ITS2 and psbA-trnH barcoding regions of Flos Lonicerae Japonicae and closely related species were evaluated for the suitability for species discrimination.

2. Materials and Methods

2.1. Materials. Flower bud samples from 45 individuals belonging to seven species, which were used as Flos Lonicerae Japonicae and Flos Lonicerae in Chinese markets, were collected from various geographical areas in China, including Henan, Shandong, and Guizhou (Table 1). All samples were identified by Professor Yulin Lin from the Institute of Medicinal Plant Development (IMPLAD), Chinese Academy of Medical Sciences. Voucher specimens were deposited in the IMPLAD herbarium in Beijing, China. Subsequently, eight ITS2 sequences of L. japonica were downloaded from GenBank for further analysis.

2.2. DNA Extraction, PCR Amplification, and Sequencing. Total genomic DNA was extracted from 25 mg of dry flower bud by the Plant Genomic DNA Kit (Tiangen Biotech Co., Beijing, China) after grinding for 1 min at 30 oscillations/s using a DNA Extraction Grinder (MM 400; Retsch, Haan, Germany). PCR amplifications of the ITS2 region were performed in a Peltier Thermal Cycler PTC200 (BioRad Lab Inc., USA) using a pair of primer ITS2F (5'-ATGCAGACCTTGTTGTAAT-3') and ITS3R (5'-GAGGCTTCTAGACCATCAAT-3') [27]. PCR reactions were performed in a 25 μL volume containing approximately 20–50 ng of genomic DNA, 2x EasyTaq PCR SuperMix (TransGen Biotech Co., China), 1.0 μL of each primer (2.5 μM), and distilled deionized water. The reaction conditions were as follows: 95°C for 5 min, 40 cycles at 94°C for 30 s, 58°C for 30 s, 72°C for 45 s, and 72°C for 10 min. The psbA-trnH was conducted as described previously [27, 42]. The desired PCR products were separated by electrophoresis in 1.2% agarose gels. The purified PCR products were directly sequenced bidirectionally by the sequencing center at Molecular Biology and Chemical (Mbchem, Shanghai, China).

2.3. Data Analysis. The forward and reverse trace files were trimmed and assembled after sequencing using the Codon-Code Aligner V 3.5.4 (CodonCode Co., USA). The ITS2 region was annotated with the HMMer software using ITS2 database [47]. The genetic distances were then calculated using MEGA 5.0 [48], according to the Kimura 2-Parameter (K2P) model. The average intra-specific distance, theta, and coalescent depth were computed to evaluate the intraspecific variation by the K2P model [49]. The average interspecific distance, minimum interspecific distance, and theta primer were used to represent interspecific divergences [24, 27]. The distribution of intra- versus inter-specific variability was assessed using the DNA barcoding gaps [50]. Species identification efficiency was determined based on BLAST1 method [51]. NJ trees were constructed to identify Flos Lonicerae Japonicae and its closely related species.

3. Results

3.1. Efficiency of PCR Amplification. DNA was isolated from 45 individuals. Some smear bands can only be found by DNA gel electrophoresis because of DNA degradation. The PCR amplification success rates for ITS2 and psbA-trnH were both 100% (Figure 1). The sequencing results showed that high-quality bidirectional trace files were obtained from the ITS2 regions.

3.2. Analysis of DNA Divergence. The ITS2 sequence lengths of 24 Flos Lonicerae Japonicae samples were 228 bp, and the average GC content was 75.4%. Only one nucleotide variation site was found at the 27th base, which is the substitution of C/T. Two new haplotypes of ITS2 sequences...
Table 1: Samples used in the present study.

| Latin name       | Haplotype (for ITS2) | Voucher no. | GenBank no. ITS2, psbA-trnH | Locality                  |
|------------------|----------------------|-------------|-----------------------------|---------------------------|
| L. japonica A1   | YC0014MT15           | JX669516, JX885527 |                            | Pingyi, Shandong         |
| L. japonica A1   | YC0014MT60           | JX682917, JX885540 |                            | Pingyi, Shandong         |
| L. japonica A1   | YC0014MT91           | JX682918, JX885545 |                            | Kaixian, Chongqing       |
| L. japonica A1   | YC0014MT104          | JX682919, JX885525 |                            | Huanggang, Hubei         |
| L. japonica A2   | YC0014MT16           | JX669517, JX885528 |                            | Pingyi, Shandong         |
| L. japonica A2   | YC0014MT29           | JX682898, JX885529 |                            | Xixia, Henan             |
| L. japonica A2   | YC0014MT31           | JX682899, JX885530 |                            | Xixia, Henan             |
| L. japonica A2   | YC0014MT32           | JX682900, JX885531 |                            | Xixia, Henan             |
| L. japonica A2   | YC0014MT33           | JX682901, JX885532 |                            | Xixia, Henan             |
| L. japonica A2   | YC0014MT41           | JX682902, JX885533 |                            | Fengqiu, Henan           |
| L. japonica A2   | YC0014MT43           | JX682903, JX885534 |                            | Fengqiu, Henan           |
| L. japonica A2   | YC0014MT54           | JX682904, JX885535 |                            | Donghai, Jiangsu         |
| L. japonica A2   | YC0014MT55           | JX682905, JX885536 |                            | Donghai, Jiangsu         |
| L. japonica A2   | YC0014MT57           | JX682906, JX885537 |                            | Donghai, Jiangsu         |
| L. japonica A2   | YC0014MT58           | JX682907, JX885538 |                            | Donghai, Jiangsu         |
| L. japonica A2   | YC0014MT59           | JX682908, JX885539 |                            | Donghai, Jiangsu         |
| L. japonica A2   | YC0014MT62           | JX682909, JX885541 |                            | Pingyi, Shandong         |
| L. japonica A2   | YC0014MT63           | JX682910, JX885542 |                            | Pingyi, Shandong         |
| L. japonica A2   | YC0014MT64           | JX682911, JX885543 |                            | Nanning, Guangxi         |
| L. japonica A2   | YC0014MT81           | JX682912, JX885544 |                            | Liupanshui, Guizhou      |
| L. japonica A2   | YC0014MT92           | JX682913, JX885546 |                            | Wanzhou, Chongqing       |
| L. japonica A2   | YC0014MT98           | JX682914, JX885547 |                            | Zhoukou, Henan           |
| L. japonica A2   | YC0014MT101          | JX682915, JX885524 |                            | Shucheng, Anhui          |
| L. japonica A2   | YC0014MT108          | JX682916, JX885526 |                            | Julu, Hebei              |
| L. japonica A2   | —                    | JQ737172     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ737175     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780993     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780991     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780986     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780987     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780988     | GenBank                     | GenBank                   |
| L. japonica A2   | —                    | JQ780989     | GenBank                     | GenBank                   |
| L. macranthoides B1 | YC0163MT10           | JX669518, JX885548 |                            | Liuzhi, Guizhou          |
| L. macranthoides B1 | YC0163MT20           | JX682920, JX885549 |                            | Banan, Chongqing         |
| L. macranthoides B1 | YC0163MT21           | JX682921, JX885550 |                            | Longhui, Hunan           |
| L. macranthoides B1 | YC0163MT25           | JX682922, JX885551 |                            | Nanjiang, Sichuan        |
| L. macranthoides B1 | YC0163MT26           | JX682923, JX885552 |                            | Puer, Yunan              |
| L. fulvotomentosa B2 | YC0164MT01           | JX669519, JX885553 |                            | Anlong, Guizhou          |
| L. fulvotomentosa B2 | YC0164MT07           | JX682924, JX885554 |                            | Xingren, Guizhou         |
| L. fulvotomentosa B2 | YC0164MT08           | JX682925, JX885555 |                            | Zhenfeng, Guizhou        |
| L. fulvotomentosa B2 | YC0164MT09           | JX682926, JX885556 |                            | zerong, Guizhou          |
| L. fulvotomentosa B2 | YC0164MT11           | JX682927, JX885557 |                            | Xingyi, Guizhou          |
| L. hypoglauca B3  | YC0165MT01           | JX669520, JX885558 |                            | Nanning, Guangxi         |
| L. hypoglauca B3  | YC0165MT02           | JX682928, JX885559 |                            | Hezhou, Guangxi          |
| L. confusa B4     | YC0166MT04           | JX669521, JX885560 |                            | Shipan, Sichuan          |
| L. confusa B4     | YC0166MT07           | JX682929, JX885561 |                            | Jianyang, Sichuan        |
| L. similis B5     | YC0169MT04           | JX669522, JX885562 |                            | Chengkou, Chongqing      |
| L. similis B5     | YC0169MT07           | JX682930, JX885563 |                            | Kaixian, Chongqing       |
| L. similis B5     | YC0169MT10           | JX682931, JX885564 |                            | Chengkou, Chongqing      |
generated in this study were submitted to GenBank, with the following accession numbers: JX669516 and JX669517. The ITS2 regions of haplotype A1 were the same as that of JX669516, whereas that of haplotype A2 was the same as that of JX669517. According to the K2P model, the intraspecific distance of Flos Lonicerae Japonicae ITS2 ranged from 0 to 0.0044. The ITS2 sequence length of Flos Lonicerae and other samples ranged from 228bp to 229bp; the average GC content was 73.9%. A total of 14 nucleotide variation sites and one insertion/deletion mutation were detected in 53 ITS2 sequences (Table 2). In this study, the ITS2 region of the other Lonicera species except L. japonica was divided into six haplotypes (Table 1). These ITS2 regions were also submitted to GenBank, and GenBank accession numbers were obtained (Table 1).

Using the same approach, also psbA-trnH sequences of the 45 Lonicera samples were analyzed (Table 2). The sequence length of the psbA-trnH of Flos Lonicerae Japonicae was 339 bp, and the average GC content was 29.5%. No variation sites were found in the psbA-trnH regions. Additional characteristics of the psbA-trnH sequences of 45 samples are shown in Table 2.

Ideal barcode sequences should provide a distinct inter-specific divergence and relatively small intra-specific variation. They must concurrently show significant differences between these two types of variation to distinguish individual intra-specific variation from variation between species. According to the K2P model, the genetic distance of Flos Lonicerae Japonicae from its closely related species was evaluated using the ITS2 regions. The inter-specific distance was 0.0331, which is larger than the intra-specific distance of 0.0010. The minimum inter-specific distance was 0.0142, which is larger than coalescent depth of 0.0019 (Table 3). Likewise, the minimum inter-specific distance for the psbA-trnH was 0.0060, which is larger than coalescent depth of 0.0004 (Table 3).

3.3. Assessment of Barcoding Gap. We investigated the distribution of genetic distance in classes of 0.010 distance units for ITS2 and psbA-trnH sequences to perform a preliminary examination of intra- versus inter-specific variation. The results showed apparent gaps between the intra- and inter-specific variability of ITS2. Obvious gaps were also found between the intra- and inter-specific variability of psbA-trnH using the same method (Figure 2).

3.4. Applicability for Species Discrimination. Two methods of BLAST1 and NJ tree techniques were used to evaluate the reliability of species identification using DNA barcoding. The results indicated that the ITS2 and psbA-trnH regions exhibited the highest identification efficiency (100%) by BLAST1 (Table 3). NJ tree effectively determines the power of a given locus combination to discriminate among species. In this study, a phylogenetic tree was constructed using the NJ method, with 1000 bootstrap replicates for the ITS2 and psbA-trnH sequences (Figure 3, Supplementary Figure 1 available online at: http://dx.doi.org/10.1155/2013/549037). The results showed that L. japonica formed into one clade, which can be distinguished successfully from its closely related species.
The applicability of the ITS2 regions for species identification is cost effective, and easy to use even for a nonspecialist [52]. DNA barcode should be easily amplified using one pair of primers. Within species, the ITS2 regions can be used as an efficient and powerful marker for distinguishing various species in Fabaceae. Pang et al. accurately identified the Ephedrae herb and its closely related species by the ITS2 DNA barcoding technique [57]. The stability and accuracy of identifying Notopterygii Rhizoma et Radix has been tested using the ITS/ITS2 barcodes [55]. All results consistently showed that the ITS2 DNA barcode exhibits good stability and accuracy in identifying Chinese materia medica.

A previous study by Sun et al. used the leaves of L. japonica and proposed the psbA-trnH regions for the identification of L. japonica and its related species [46]. We obtained genomic DNA from the flowers of Flos Lonicerae Japonicae and assessed their PCR amplification and identification efficiency to test the psbA-trnH regions. The results revealed that psbA-trnH can also be used to identify Flos Lonicerae Japonicae and verified the results of Sun et al. [46]. This study simultaneously showed that ITS2 regions can be used to distinguish Flos Lonicerae Japonicae. However, in the study by Sun et al., the resulting PCR product of ITS2 was not amplified using the approach employed by Chen et al. [27]. Compared with the study by Sun et al., the present study obtained two findings. One is that genomic DNA, which can be improved for follow-up experiments, was extracted by extending the pyrolysis time. The other is that the ITS2 regions were amplified successfully by increasing the annealing temperature from 56°C to 58°C using a universal primer and a 2× EasyTaq PCR SuperMix (TransGen Biotech Co., China). Sequence analysis indicated that the average GC contents of Flos Lonicerae Japonicae was 75.4%, which may be that reason Sun et al. failed to obtain the ITS2 regions by the Chen et al. method [27].

Several studies have reported the existence of multiple copies of ITS2 in plants and animals [58]. Questions have been raised whether the sequence obtained by PCR would be stable and representative as a DNA marker. Alvarez and Wendel suggested that the presence of different copies of ITS2 can result in misleading phylogenetic inferences [58]. However, Coleman indicated that ITS2 can be effectively treated as a single locus [59]. The most recent study also revealed that using the major variants alone is sufficient for

### Table 2: Sequence characteristics of ITS2 and psbA-trnH of Flos Lonicerae Japonicae and its related species.

| Sequence characteristics          | ITS2       | psbA-trnH  |
|-----------------------------------|------------|------------|
| Length range in Flos Lonicerae Japonicae (bp) | 228        | 339        |
| Length in all taxa (bp)           | 228–229    | 332–356    |
| Average of GC content in Flos Lonicerae Japonicae (%) | 75.4       | 29.5       |
| Average of GC content in all taxa (%) | 73.9       | 29.6       |
| No. of variable sites in Flos Lonicerae Japonicae | 1          | 0          |
| No. of variable sites in all taxa  | 14         | 10         |

### Table 3: Inter- and intra-specific genetic divergence and identification efficiency in ITS2 and psbA-trnH sequences of L. japonica and its related species.

| Parameter                              | ITS2            | psbA-trnH         |
|----------------------------------------|-----------------|-------------------|
| All inter-specific distance            | 0.0331 ± 0.0119 | 0.0161 ± 0.0066   |
| Theta primer                           | 0.0253 ± 0.0093 | 0.0126 ± 0.0050   |
| The minimum inter-specific distance    | 0.0142 ± 0.0105 | 0.0060 ± 0.0060   |
| All intra-specific distance            | 0.0010 ± 0.0018 | 0.0000 ± 0.0003   |
| Theta                                  | 0.0011 ± 0.0093 | 0.0002 ± 0.0004   |
| Coalescent depth                       | 0.0019 ± 0.0024 | 0.0004 ± 0.0011   |
| Identification efficiency (BLAST1)     | 100%            | 100%              |

### 4. Discussion

An ideal barcoding system should have sufficient variation among the sequences to discriminate species; however, it also needs to be sufficiently conserved to achieve less variability within species than between species [52]. In addition, the DNA barcode should be easily amplified using one pair of universal primers; identification should be rapid, accurate, cost effective, and easy to use even for a nonspecialist [53]. The applicability of the ITS2 regions for species identification has been tested by Chen et al. [27]. In recent years, the DNA barcoding technique has become the main method for species identification [54–56].

In the present study, the ITS2 region was used for the first time to identify Flos Lonicerae Japonicae and its closely related species. The samples were collected from the main producing areas of the species. All samples were obtained from the flower bud, which is the medicinal part of L. japonica. The following results were obtained from this study. First, the length of the ITS2 of Flos Lonicerae Japonicae was 228 bp, which is short and relatively easy to amplify using one pair of universal primers. Second, the ITS2 regions of Flos Lonicerae Japonicae obtained from different areas were similar; only one variable site was found. The minimum inter-specific distance was 0.0142, which is significantly larger than coalescent depth of 0.0019. Divergence at the inter-specific level and relatively low divergence at the intra-specific level (Table 3) were found. Third, the identification efficiency of the 53 samples was 100% by the BLAST1 method. The results obtained using the NJ tree can also be used to distinguish Flos Lonicerae Japonicae successfully. In summary, the ITS2 sequence can be used as a DNA barcode for identifying Flos Lonicerae Japonicae and its closely related species. To date, ITS2 regions have been widely used to distinguish related species. The study by Gao et al. [31] indicated that ITS2 regions can be used as an efficient and powerful marker for distinguishing various species in Fabaceae. Pang et al. accurately identified the Ephedrae herb and its closely related species by the ITS2 DNA barcoding technique [57]. The stability and accuracy of identifying Notopterygii Rhizoma et Radix has been tested using the ITS/ITS2 barcodes [55]. All results consistently showed that the ITS2 DNA barcode exhibits good stability and accuracy in identifying Chinese materia medica.
phylogeny construction and species determination in most cases although intragenomic multiple variants are frequently found within each genome. Inclusion of minor variants further improves the resolution of species identification [60]. In the present study, the PCR amplification success rates for ITS2 was 100% and high-quality sequence stability from the 45 samples was obtained. This result indicates that the major variants alone are sufficient for PCR amplification of the ITS2 region. Therefore, the ITS2 DNA barcode can be used to identify Flos Lonicerae Japonicae and its closely related species.

The results of this study can help identify Flos Lonicerae Japonicae and Flos Lonicerae in the market and ensure safe and stable supply of traditional medicines derived from *L. japonica*. These findings can be used as reference for the authentication of other medicinal herbs, especially the flower types. The results can also be valuable in resource protection, industrial production, customs, forensic examination, and plant quarantine.

5. Conclusions

ITS2 and *psbA-trnH* were examined for their stability and accuracy in identifying Chinese materia medica in the case study with Flos Lonicerae Japonicae. The results indicate that using the ITS2 and *psbA-trnH* regions as DNA barcodes can stably and accurately distinguish Chinese materia medica as well as provide a new technique to ensure clinical safety in using traditional Chinese medicines.

Conflict of Interests

The authors have declared that no competing interests exist in this paper.
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