An authenticity survey of herbal medicines from markets in China using DNA barcoding

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Adulterant herbal materials are a threat to consumer safety. In this study, we used DNA barcoding to investigate the proportions and varieties of adulterant species in traditional Chinese medicine (TCM) markets. We used a DNA barcode database of TCM (TCMD) that was established by our group to investigate 1436 samples representing 295 medicinal species from 7 primary TCM markets in China. The results indicate that ITS2 barcodes could be generated for most of the samples (87.7%) using a standard protocol. Of the 1260 samples, approximately 4.2% were identified as adulterants. The adulterant focused on medicinal species such as Ginseng Radix et Rhizoma (Renshen), Radix Rubi Parvifolii (Maomeigen), Dalbergiae odoriferae Lignum (Jiangxiang), Acori Tatarinowii Rhizoma (Shichangpu), Inulae Flos (Xuanfuhua), Lonicerae Japonicae Flos (Jinyinhua), Acanthopanacis Cortex (Wujiapi) and Bupleuri Radix (Chaihu). The survey revealed that adulterant species are present in the Chinese market, and these adulterants pose a risk to consumer health. Thus, regulatory measures should be adopted immediately. We suggest that a traceable platform based on DNA barcode sequences be established for TCM market supervision.

Counterfeit drugs are frequently sold on the market, resulting in adverse effects, drug resistance, and death1–3. The proportion of counterfeit drug sales in developing countries is approximately 10%, and the proportion may be greater than 50% when considering medicines purchased online4–6. Drug counterfeiting causes health problems, particularly in developing countries where drug regulatory systems are weak or ineffective7. Although organizations are working to develop methods to protect the drug supply chain, counterfeit drug use has dramatically increased in recent years. Each week, new cases of counterfeit medicines are reported around the world8. According to numerous official sources, the proportion of counterfeit medicines in African countries has reached 80%9. Because of a lack of suitable identification methods, the number of reported cases of counterfeit medicine seems to be rising9.

Traditional Chinese medicine (TCM) plays an important role in disease prevention and treatment, and research has demonstrated the clinical efficacy of TCM against certain diseases for which conventional therapy is ineffective or has associated side effects10. In recent years, there has been a huge increase in the use of herbal products; however, there are also numerous reports of adulterant herbal medicine use in many developing countries, which poses a major public health risk. The World Health Organization (WHO) defines a counterfeit product as one that is mislabelled deliberately and fraudulently with respect to its identity or source. Crude materials provide the basis for genuine drug production, but recently, there have been many alarming reports of counterfeit or adulterant drugs that caused life-threatening poisonings. For example, a type of tea adulterant containing Adenostyles alliariae caused serious liver disease after long-term use11,12. Adulterant tea mixed with Illicium anisatum (which contains neurotoxic substances)13 and cases of toxicity caused by Aconitum14 and Datura metel15 have also been reported. Moreover, approximately 50% of artesunate (extracted from Artemisia annua L.) tablets sampled in Southeast Asia were reported to be counterfeit16; Severe kidney damage caused by adulteration with Aristolochia species is frequently reported17–19; as a result of aristolochic acid toxicity. Song showed that 60% of commercial Rhodiola products are adulterants, which indicates a potential safety issue20. All of these life-threatening poisoning cases threaten the safe use of TCM. As a result, the detection of adulterant drugs is becoming a growing challenge8.

Traditional identification methods recognize materials by their morphological characteristics, and these methods primarily depend on human expertise. However, in some cases, it is extremely difficult for taxonomists to...
definitively identify plant genera, such as *Crataegus* and *Salix*. Chemical analyses, such as high-performance liquid chromatographic-mass spectrometric (HPLC-MS)\(^2\), near-infrared spectroscopy (NIRS)\(^3\), and liquid chromatography-mass spectrometry (LC-MS assays)\(^4\), can be used to detect chemical compositions to identify adulterant products. However, none of these methods alone can definitively identify closely related species that share remarkably similar morphological characteristics and chemical profiles. These techniques produce only indirect evidence of fraud and cannot definitively determine the identity of the given species. Therefore, there is an urgent need for rapid and simple identification procedures for the rapid inspection of raw herbal materials.

DNA barcoding is a new molecular diagnostic technology that was first proposed by Canadian zoologist Paul Hebert in 2003, and it identifies species by using a recognized standard, short genomic sequence\(^5\). DNA barcoding provides consistent and reliable results regardless of the age, plant part, or environmental factors of the sample\(^5\). Researchers can evaluate species information accurately by analysing DNA sequences. Other investigators have suggested that a global DNA barcode revolution would become a "big science" research programme after the human genome project\(^6\), and Miller published "the Renaissance of DNA barcode and taxonomy" in PNAS\(^7\).

This approach has been repeatedly reported in academic journals (e.g., Nature, Science) and in media outlets (e.g., National Geographic News, *The New York Times*) stating that DNA barcode technology has become a global innovation for academic research on biological taxonomy. Chen *et al.* have analysed more than 6000 plant samples belonging to 4800 species from 753 distinct genera by using the chloroplast regions *psbA-trnH, matK, rbcL, rpoC1, ycf5* and the nuclear loci ITS and ITS2. These investigators suggested that the internal transcribed spacer (ITS) fails to be amplified and sequenced in most samples and that ITS2 is the most suitable locus for DNA barcoding research, followed by *psbA-trnH* as a complementary region\(^8\). By using an ITS2 + *psbA-trnH* two-loci barcode combination, our group developed a TCM barcode platform, called the Traditional Chinese Medicine Database (TCMD)\(^9\), which contains 78,847 barcodes belonging to 23,262 medicinal species listed in the Chinese, European, Indian, Japanese, Korean and American Herbal Pharmacopoeias\(^10\). There are more than three samples per species in this database\(^10\). At present, the TCMD is the largest DNA barcode database of medicinal materials. The TCMD also contains the DNA barcoding standard operating procedure (SOP) and provides bioinformatics tools to assist in data analysis for researchers in the herbal identification industry. The TCMD can be accessed at http://www.tcmbarcode.cn/en/.

In this study, we investigated the proportions and varieties of adulterant medicine in herb markets with the aim of protecting consumers from health risks associated with herbal product substitution and contamination by using a standard DNA barcoding method. A total of 1436 raw herbal samples representing 295 medicinal species were collected from the 7 primary markets in China. The advantages and limitations of DNA barcoding for the authenticated of complex TCM materials by using the TCMD database are also discussed. Additional details are described in subsequent sections.

**Results**

**Efficiency of PCR amplification and sequencing.** Of the 1436 samples, 176 (12.26%) could not be successfully amplified and sequenced, primarily cortex and fungal medicinal species. The failure rates of cortex and fungi medicinal species were approximately 21/93 (22.6%) and 5/23 (21.7%), respectively. The unamplified species were Magnoliae Officinalis Cortex (Houpo), Periplocae Cortex (Xiangjiapi), Phellodendri Chinensis Cortex (Huangbo), Fraxini Cortex (Qinpi) and Polyporus (Zhuling). In contrast, stem and folium medicinal species were easily amplified, with failure rates of approximately 3.1% and 5.1%, respectively. There was difficulty with fungus medicinal species, such as *Fritillariae cirrhosae Bulbus* (Chuanbeimu), *Rhei Radix* (Fangfeng), *Glycyrrhizae Radix* (Gancao), and *Polygoni multiflorum Radix* (Heshouwu), and 13.9% of the cortex samples were found to be adulterant, including the *Albiziae Cortex* (Hehuanpi) samples were found to be derived from *Albizia julibrissin* (Hehuanpi), *Pseudolaricis Cortex* (Tujingpi) and *Acanthopanacis Cortex* (Wujiapi) from different markets.

Of the 295 medicinal species in this study, 198 could be amplified successfully and were validated, including species that are commonly used in TCM, such as *Fritillariae cirrhosae Bulbus* (Chuanbeimu), *Rhei Radix* and *Zhamai* (Dahuang), *Angelicae Sinensis Radix* (Danggui), *Codonopsis Radix* (Dangshen), *Saposhnikoviae Radix* (Fangfeng), *Glycyrrhizae Radix* (Gancao), and *Polygoni multiflorum Radix* (Heshouwu), and the other 97 varieties exhibited failed amplification and adulterants to some extent. The adulterants included *Ginseng Radix* and *Rhzima* (Renshen), *Rubia Parvifolia* (Maomeigen), *Dalbergiae odoriferae Lignum* (Jiangxiang), *Acori Tatarinowii Rhizoma* (Shichangpu), *Inulae Flos* (Xuanfuhua), *Lonicerae Japonicae Flos* (Jinyinhua), *Acanthopanacis Cortex* (Wujiapi) and *Bupleuri Radix* (Chaihu). The original species of *Albiziae Cortex* (Hehuanpi) was *Albizia julibrissin*, but five of the 9 *Albiziae Cortex* (Hehuanpi) samples were found to be derived from the *Cortex of Albizia kalkora* Prain (Shanhehuanpi) (Fig. 2), 3 of the 15 *Ginseng Radix* (derived from *Panax ginseng*) samples were found to be Panacis quinquefolii Radix (derived from *Panax quinquefolius*).
| Sample No. | Latin Name of Medicinal materials | Chinese Name | Latin Name of original species | Medicinal Parts | Markets | Identification Result |
|------------|-----------------------------------|--------------|--------------------------------|-----------------|---------|-----------------------|
| FHCQ076    | Abri Herba                        | Jigacao      | Abrus cantoniensis             | Herb            | CQ      | Abrus mollis          |
| S1068      | Abri Herba                        | Jigacao      | Abrus cantoniensis             | Herb            | HN      | Abrus mollis          |
| FHCQ072    | Acanthopanicis Cortex             | Wujiapi      | Acanthopanax gracilistylus     | Cortex          | CQ      | Acanthopanax giraldis |
| FHYL237    | Acanthopanicis Cortex             | Wujiapi      | Acanthopanax gracilistylus     | Cortex          | GX      | Periploca sepium      |
| S0471      | Acanthopanicis Cortex             | Wujiapi      | Acanthopanax gracilistylus     | Cortex          | BZ      | Periploca sepium      |
| S1188      | Acanthopanicis Senticosi Radix et Rhizoma Seu Caulis | Ciwujia | Acanthopanax senticosus       | Radix et Rhizome | HN      | Alangium chinense     |
| S0711      | Acanthopanicis Senticosi Radix et Rhizoma Seu Caulis | Ciwujia | Acanthopanax senticosus       | Radix et Rhizome | BZ      | Aralia sp.            |
| FHCQ065    | Acori Tatarinowii Rhizoma         | Shichangpu   | Acorus tatarinowii             | Radix et Rhizome | CQ      | Acorus sp.            |
| S0206      | Acori Tatarinowii Rhizoma         | Shichangpu   | Acorus tatarinowii             | Radix et Rhizome | BZ      | Acorus sp.            |
| FHGAG014   | Acori Tatarinowii Rhizoma         | Shichangpu   | Acorus tatarinowii             | Radix et Rhizome | AG      | Acorus calamus        |
| FHGAG234   | Acori Tatarinowii Rhizoma         | Shichangpu   | Acorus tatarinowii             | Radix et Rhizome | AG      | Acorus calamus        |
| FHGAG420   | Albiziae Flos                      | Hebuanghua   | Althea julibrissin             | Flowers         | AG      | Celastrus orbiculatus |
| FHYL134    | Albiziae Cortex                   | Hehuansi     | Althea julibrissin             | Cortex          | AG      | Althea kalkora        |
| FHYL140    | Albiziae Cortex                   | Hehuansi     | Althea julibrissin             | Cortex          | CQ      | Althea kalkora        |
| FHCQ187    | Albiziae Cortex                   | Hehuansi     | Althea julibrissin             | Cortex          | AG      | Althea kalkora        |
| AG015      | Albiziae Cortex                   | Hehuansi     | Althea julibrissin             | Cone              | AG      | Althea kalkora        |
| S0436      | Albiziae Cortex                   | Hehuansi     | Althea julibrissin             | Cone              | BZ      | Althea kalkora        |
| FHGAG163   | Alpiniae Oxyphyllae Fruits & Seeds | TiZi         | Alpinia oxyphylla              | Fruits & Seeds   | AG      | Foeniculum vulgare    |
| S0243      | Angelicae Dahuriae Radix          | Bazihi       | Angelica dahurica var.formosana | Radix et Rhizome | BZ      | Atras psichellus      |
| S0365      | Angelicae Pubescentis Radix       | Duhuo        | Angelica pubescens f. biseriata | Radix et Rhizome | BZ      | Angelica amurensis    |
| S0485      | Angelicae Pubescentis Radix       | Duhuo        | Angelica pubescens f. biseriata | Radix et Rhizome | BZ      | Levisticum officinale |
| AG165      | Arnnebiae Radix                   | Zicao        | Arnnebia euchroma              | Radix et Rhizome | AG      | Lithospermum erythrorhizon |
| FHYL175    | Bupleuri Radix                    | Chaihu       | Bupleurum chinense             | Radix et Rhizome | GX      | Bupleurum sp.         |
| S0552      | Bupleuri Radix                    | Chaihu       | Bupleurum chinense             | Radix et Rhizome | BZ      | Bupleurum sp.         |
| FHYL394    | Bupleuri Radix                    | Chaihu       | Bupleurum chinense             | Radix et Rhizome | BZ      | Bupleurum sp.         |
| S0511      | Bupleuri Radix                    | Chaihu       | Bupleurum chinense             | Radix et Rhizome | BZ      | Bupleurum sp.         |
| FHCQ015    | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | CQ      | Caesalpinia sappan    |
| AG008      | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | AG      | Caesalpinia sappan    |
| S0219      | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | BZ      | Caesalpinia sappan    |
| S0410      | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | BZ      | Caesalpinia sappan    |
| S1241      | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | BZ      | Caesalpinia sappan    |
| S1245      | Dalbergiae Odoriferae Lignum      | Jiangzhang   | Dalbergia odorfera             | Stems            | BZ      | Caesalpinia sappan    |
| FHYL083    | Erycisbes Caulis                  | Dinggongteng | Erycites schmidtii            | Stems            | GX      | Covolubalcaeae        |
| AG230      | Euphorbia Ebracteolatae Radix     | Langdu       | Euphorbia fischeriana         | Radix et Rhizome | AG      | Stellera chamaejasme  |
| S0082      | Sojae semen Praeparatum           | Dandouchi    | Glycine max                   | Fruits & Seeds   | BZ      | Phaseolus vulgaris    |
| FHYL133    | Imulae Flos                       | Xuanfuhua    | Inula japonica                | Flowers          | HN      | Inula lineariifolia   |
| S1135      | Imulae Flos                       | Xuanfuhua    | Inula japonica                | Flowers          | BZ      | Inula lineariifolia   |
| S0612      | Imulae Flos                       | Xuanfuhua    | Inula japonica                | Flowers          | BZ      | Inula lineariifolia   |
| S0757      | Imulae Flos                       | Xuanfuhua    | Inula japonica                | Flowers          | BZ      | Inula lineariifolia   |
| FHYL059    | Lonicerae Japonicae Flos          | Jinyinhua    | Loniceria japonica            | Flowers          | AG      | Loniceria macranthoides |
| AG069      | Lonicerae Japonicae Flos          | Jinyinhua    | Loniceria japonica            | Flowers          | AG      | Loniceria macranthoides |
| FHCQ151    | Moslae Herba                      | Xiangru      | Mosla chinensis               | Herb            | CQ      | Elbohizia ciliata     |
| MM042      | Ginseng Radix et Rhizoma          | Renshen      | Panax ginseng                 | Radix et Rhizome | AG      | Panax quinquiesifolium |
| MM0422-1   | Ginseng Radix et Rhizoma          | Renshen      | Panax ginseng                 | Radix et Rhizome | AG      | Panax quinquiesifolium |
| SQAG02     | Ginseng Radix et Rhizoma          | Renshen      | Panax ginseng                 | Radix et Rhizome | AG      | Panax quinquiesifolium |
| AG264      | Panacis Japonici Rhizoma          | Zhajieshen   | Panacis japonicus             | Radix et Rhizome | AG      | Panax sp.             |
| S0593      | Pseudolaricis Cortex              | Tujingpi     | Pseudolarix amabilis          | Cortex          | BZ      | Celastrus angulatus   |
| AG104      | Pseudolaricis Cortex              | Tujingpi     | Pseudolarix amabilis          | Cortex          | AG      | Celastrus angulatus   |
| FHCQ123    | Puerariae Thonsoni Radix          | Fenge        | Pueraria thonsoni             | Radix et Rhizome | CQ      | Pueraria lobata       |
| S0087      | Pulsatillae Radix                 | Baitouweng   | Pulsatilla chinensis          | Radix et Rhizome | BZ      | Platycodon grandiflorus |
| S0669      | Rhapontici Radix                  | Loula        | Rhaponticum uniflorum         | Radix et Rhizome | BZ      | Echinops latifolius   |
| B1         | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | BZ      | Cirsium japonicum    |
| B2         | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | BZ      | Cirsium japonicum    |
| B3         | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | BZ      | Cirsium japonicum    |
| B4         | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | BZ      | Cirsium japonicum    |

Continued
and 10 of the 19 Radix Rubi Parvifolii (Maomeigen) samples were Cirsii Japonici Heiba (Daji), Rosae Chinensis Flos (Yuejihua) or the root of Rubus alceaefolius. Two Lonicerae Japonicae Flos (derived from Lonicera japonica) samples were identified as Lonicerae Flos (derived from Lonicera macranthoides). Of the 4 Acanthopanacis Cortex (Wujiapi) samples, 1 was an amplification failure, 2 were identified as Periplocae Cortex (Xiangjiapi) and one sample was derived from the Cortex of Eleutherococcus giraldii Harms (Hongmaowujiapi). All 6 of the Dalbergiae odoriferae Lignum (Jiangxiang) samples were adulterant, and they were identified as Sappan Lignum (derived from

| Sample No. | Latin Name of Medicinal materials | Chinese Name | Latin Name of original species | Medicinal Parts | Markets | Identification Result |
|------------|-----------------------------------|--------------|-------------------------------|----------------|---------|-----------------------|
| CZG01      | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rubus alceaefolius    |
| CZG02      | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rubus alceaefolius    |
| AG1        | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rosa chinensis        |
| AG3        | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rosa chinensis        |
| AG4        | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rosa chinensis        |
| AG2        | Rubi Rhizoma                      | Maomeigen    | Rubus parvifolius             | Radix et Rhizome | AG      | Rosa chinensis        |
| S1297      | Spirodelae Herba                  | Fuping       | Spirodela polyrrhiza          | Herb           | HN      | Wolffia globosa       |

Table 1. Identification of adulterant medicinal plant materials. Note: BZ represent Anhui Bozhou Herb Market; AG represent Hebei Anguo Herb Market; CQ represent Chongqing Chuqimen Herb Market; GX represent Guangxi Yulin Herb Market; HN represent Henan Yuzhou Herb Market; QP represent Guangdong Qingping Herb Market; HUC represent Sichuan Hehuachi Herb Market.

![Figure 1. The adulterant rate from different samples of medicinal materials, including the radix et rhizome, fruits and seeds, herbs, flos, stems, cortex, foliums, and fungi.](image)

![Figure 2. The adulterant rate observed for 26 medicinal plants.](image)
from *Caesalpinia sappan*. In total, 53 samples were adulterants, and for 9 samples, the exact species could not be determined (Table 1).

**Survey of 7 herb markets.** The 7 herb markets investigated in this study included Guangxi Yulin (GX), Hebei Anguo (AG), Henan Yuzhou (HN), Anhui Bozhou (BZ), Chongqing Cuimeng (CQ), Guangdong Qingping (QP), and Sichuan Hehuachi (HUC). The exception of HUC, different types of adulterant medicine were found at all of the herb markets, with percentages ranging from 3.7% in AG to 13.3% in QP. Of the 176 unamplified samples, QH had the highest rate, at approximately 27.4%, and the rates for QP, HN and GX were approximately 18.9%, 16.8% and 13.5%, respectively. HUC had the lowest rate.

**Discussion**

**Advantages and limitations of DNA barcoding in the quality analysis of crude TCM materials.** DNA barcoding is a universal method that can be used to develop an international standard for product identification. At the Fourth International Barcode of Life Conference, a three-loci barcode (*matK + rbcL + psbA-trnH*) was suggested for plant identification. Chen *et al.* suggested ITS2 as a preferred barcode for medicinal plants26. Han *et al.* also showed that ITS2 is suitable for identifying medicinal samples36. In the present study, ITS2 was shown to be a very promising and effective tool for assessing adulterants in TCM markets. Of the 1260 ITS2 sequences generated, 4.2% were adulterants. Only 1 of the 7 herb markets provided authentic products with no adulterants.

The use of DNA barcoding to identify commercialized medicinal plants in southern Morocco suggests that a reference barcoding database should contain an adequate number of sequences from different locations27. Previous studies have identified the uncertainties of assigning unknown herbal products with incomplete reference barcode databases in GenBank and BOLD. One of the goals of the Herb-BOL (barcode of life) research programme was to build an herbal barcode library that covered all 1800 known medicinal species used in commercial products. Because of the importance of authenticating medicinal plant materials, it is vital to develop an exclusive, extensive herbal database35. The GenBank database ([http://www.ncbi.nlm.nih.gov/genbank/](http://www.ncbi.nlm.nih.gov/genbank/)) is possibly one of the largest sequence databases and is one of the most frequently used databases for species identification. An unknown DNA sequence can be rapidly compared to known species sequences with the BLAST program38. However, at present, many medicinal sample sequences are not adequately represented in GenBank, and in some cases investigators could only declare results at the genus level based on sequence similarity. The TCMD is a barcode database that is exclusively devoted to medicinal species, and it contains 23,262 medicinal and closely related species, including adulterants and substitutions. The TCMD covers almost all of the medicinal materials listed in herbal pharmacopoeias from around the world, including China, Europe, India, Japan, Korea and the United States. Currently, the TCMD is the largest DNA barcode database of medicinal materials in the world29. Thus, the TCMD platform is the most suitable for the rapid screening of crude medicinal materials. The establishment of the TCMD has greatly improved the resources available for medicinal species identification.

Given that some medicinal samples are heavily processed and that some artificial adulterant samples do not contain DNA, DNA barcoding is not sufficient to confirm the identity of any given sample. In the current investigation, we found that at least 50% of the medicinal samples on the market have been fumigated with sulphur to extend the storage time and prevent insect infestation and mildew. In some cases, samples treated with sulphur, such as *Lycium barbarum* and *Dioscorea opposita*, appeared very clean and bright in colour and could be sold at a high price. This factor may also affect the amplification efficiency of the sample. In addition, many herbs contain secondary compounds such as polysaccharides, pigments and others. We washed the precipitants with wash buffer three times to remove sticky residues before extraction, but some of the residues could not be removed, which could also make it difficult to extract DNA from these samples. Approximately 12.26% of the samples evaluated in this study could not be successfully amplified and sequenced.

DNA barcoding is an efficient tool for the identification of herbs and for the determination of various adulterants. However, DNA barcoding does not currently yield information regarding the concentration of active ingredients. Thus, DNA barcoding cannot be used to determine whether medicinal samples meet pharmacopoeia standards. In other words, DNA barcoding can be used to establish herbal authenticity but cannot be used to evaluate herbal quality. This drawback indicates that a combination of DNA barcoding and chemical analysis is necessary for a comprehensive quality assessment of herbal samples. HPLC has been used for the differentiation of accessions collected from different geographic regions. DNA barcoding has been used for the differentiation of inter- and intraspecific variations and to detect adulterations. Attempts have also been made by the author to match the results of DNA barcoding to the chemical analysis techniques of *Salvia* L.39.

**Building a traceable platform for traditional Chinese medicine using DNA barcoding.** In many developing countries, the introduction of herbal medicine products into the marketplace is not adequately monitored. Genuine (Daodi) herbs are usually considered to be high-quality medicinal materials that are produced in the Daodi area. However, because many genuine medicinal plants are transported to other places, their characteristics will be changed. In TCM markets, many sellers advertise that their herbs come from the genuine area, but there are no methods to evaluate genuine characteristics. Furthermore, herbal medicine contamination is higher because of the lower stringency of the rules and regulations governing the quality of these herbs in different countries40-41. In the present study, a survey of TCM markets identified approximately 4.2% of the samples as adulterants. Such adulterant incidents will only increase if measures are not taken to prevent them. Thus, it is necessary to build a traceable platform to ensure the safe use of TCM.

At present, DNA barcode technology is the best technology for providing traceability. Liu *et al.* successfully converted DNA barcoding sequences into twodimensional barcodes (2D-barcodes). In addition, our research group has developed an automated process that converts DNA barcode sequences into 1D- and 2D-barcodes.
Other information, including planting, processing and additional consumer information, can also be databased and converted into a 2D-barcode. Smartphones can be used as 2D-barcode readers so that consumers can conveniently scan samples to access information. This type of traceability system would not only help to manage TCM authentication but would also provide a valuable tool to improve TCM quality. Consumers could obtain all the information regarding a commercial TCM that was on the market, including planting, production, processing and circulation information, by scanning the 2D-barcode on the package. A workflow outlining such a system is shown in Fig. 3. In view of the above information, the establishment of a traceability system for TCM based on DNA barcode sequences is urgently needed.

The future of DNA barcoding. Traditionally, commonly used identification methods require special skills acquired through extensive experience; thus, only experts can identify taxa accurately. The current study showed that ITS2 sequences could be used to efficiently identify medicinal species. The herbal industry should adopt DNA barcoding to authenticate the raw materials used to manufacture its products.

DNA barcoding can be easily implemented and will play an increasingly important role in medicinal identification because of its ability to rapidly evaluate samples from leaves, seeds, flowers, dry materials, museum specimens, powders or medicinal materials from which DNA can be obtained. DNA barcoding and next-generation sequencing technology are powerful tools for identifying herbal ingredients in patient medicines. There are limitations to the four common methods of identification, namely, original, microscopy, morphological, and physicochemical identification. The DNA barcoding tool can provide supplementary information to improve classifications and to enable a critical examination of the precision of the four common methods used in medicinal material identification. Descriptions of "medicinal materials" in the pharmacopoeia of China with attached DNA sequences should be actively encouraged. Identification approaches that integrate DNA barcoding, morphological characters and chemical attribute information will achieve maximum efficiency for medicinal material identification. Researchers will have easy access to all the related herbal information in the database. With the development of pyrosequencing, sequencing costs have been dramatically reduced, which opens the way to the high-throughput sequencing of ITS2 sequences, facilitating a wide range of research possibilities using medicinal species. However, for some closely related species, such as the 9 unidentified samples in this study, identification will be very difficult when using universal primers, in which case a better approach would be to use the whole chloroplast genome as a super barcode.

In conclusion, the current TCM markets are unregulated. The consideration of simple and low-cost measures, such as DNA barcoding, has the potential to make a major contribution to the detection of adulterant products in TCM markets. The present work effectively demonstrates the feasibility of this approach. According to the TCMD, 4.2% of the samples we evaluated were adulterants. The TCMD provides users with easy access for sequence comparisons. The improvement of the TCMD will fulfill its important role in the authentication of medicinal ingredients, which will be beneficial to the entire Chinese herbal industry.

**Materials and Methods**

**Plant materials.** A total of 1436 raw herb samples representing 295 medicinal species were used, including 515 samples of radix et rhizoma, 451 samples of fruit and seeds, 115 samples of herbs, 98 varieties of flos, 82 stem samples, 93 cortex samples, 59 folium samples and 23 fungus samples. The samples were purchased from 7 of the primary herbal markets in China, with 163 samples from Guangxi Yulin (GX), 536 samples from Hebei Anguo (AG), 95 samples from Henan Yuzhou (HN), 402 samples from Anhui Bozhou (BZ), 146 samples from Chongqing Cuqimeng (CQ), 37 samples from Guangdong Qingping (QP) and 57 samples from Sichuan Hehuachi (HUC) (Fig. 4). Of the 295 medicinal species, 294 were listed in the Chinese Pharmacopoeia, and they accounted for approximately 96.4% (133 varieties) of the commonly used varieties in TCM (total of 138 varieties). Thus, the number of samples collected was large enough to be representative. All the specimens were deposited in the herbarium at the Institute of Medicinal Plant Development. The entire list of 1436 samples can be found in Supplementary Table S1 online. The locations of the 7 markets are shown in Fig. 4, which was created using an open source web site (http://www.dituhui.com/) with the latitude and longitude information for the 7 herb markets.
markets. The photographs were obtained from AG, which is the largest market in China, and were taken by co-author Baosheng Liao. The map and photographs were combined with Photoshop software.

**DNA extraction and polymerase chain reaction (PCR) amplification.** A 75% alcohol solution was used to clean the surfaces of the herbal material prior to DNA extraction to prevent fungal DNA contamination, and then one piece of each sample was ground into powder with a FastPrep bead mill (Retsch MM400, Germany). Total DNA was extracted with a Plant Genome DNA Kit (Tiangen Biotech Co., China), which is based on the CTAB approach. The key procedure was modified as follows. First, the powder was washed with wash buffer three times to remove sticky residues from the precipitant before extraction. Second, after the extraction buffer was added, the samples were incubated at 58 °C for 8–12 hours. Third, an equal amount of ice-cold isopropanol was used to precipitate the DNA at −20 °C in a refrigerator for at least 30 minutes. Other procedures were routinely performed as indicated in the CTAB method. The ITS2 was amplified using universal primers28. The PCR reaction mixture consisted of 1 μL (approximately 30 ng) genomic DNA, 1 × PCR buffer without MgCl2, 2.0 mM MgCl2, 0.2 mM of each dNTP, 0.1 μM of each primer (which were synthesized by Sangon Co., China), and 1.0 U of Taq DNA Polymerase (BiocolorBioScience & Technology Co., China). The PCR conditions were 40 cycles at 94 °C for 30 s, 56 °C for 30 s and 72 °C for 45 s. The entire PCR process was ended by incubating the samples at 72 °C for 10 min with a Peltier Thermal Cycler PTC0200 (Bio-Rad Lab, Inc., USA).

**Sequencing and analysis.** The PCR products were purified with a QIAquick PCR purification kit (Tiangen Biotech, Beijing, China) and were directly sequenced on an ABI 3730XL sequencer (Applied Biosystems, USA) by using the original amplification primer as the sequencing primer. The original forward and reverse sequences were assembled with a CodonCode Aligner 3.0. The assembled sequences were annotated and delimited with a hidden Markov model (HMM)-based method47, and the complete ITS2 sequences were pasted into the identification module on TCMD (http://www.tcmbarcode.cn/en/). After the query sequence was submitted, a BLASTN algorithm was activated, and its nearest neighbours to all the reference sequences were made available. When a best match to a reference sequence has been found, the identification module can provide a species-level identification and the Latin name of the best-match species will be given29.

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Figure 4. The 7 primary herb markets distributed throughout China. Note: The three colours represent the rate of genuine, adulterant and failed identification for different markets, respectively.
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Author Contributions
J.S. and S.C. conceived the study and participated in its design. J.H., X.P. and H. Y. contributed samples and performed the experiments. B.L. took the photographs and analysed the data. J.H., J.S. and S.C. drafted the manuscript. All authors have read and approved the final manuscript.

Additional Information
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