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## HOW TO CORRECTLY IDENTIFY HERBAL MATERIALS IN MARKET: A CASE STUDY BASED ON DNA BARCODES

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#### **Abstract**

**Background:** Traditional methods for identifying herbal medicines have many shortcomings. In this study, we aim to test discriminating ability of DNA barcodes and explore feasible method on evaluating identification results.

**Materials and Methods:** Materials of whole-plant medicines were sampled from herbal market. 111 samples were used for DNA sequencing of ITS2 and *trnH-psbA* regions. Assembled sequences were searched against reference database using the BLAST method. Comprehensive evaluation based on pharmacognostic investigation, macroscopic identification and identification of DNA barcodes were performed for authentication of the herbal materials.

**Results:** As a result, ITS2 had better identifying power than *trnH-psbA* in species-specific level (55.86% & 45.95%), as well as worse success rate of DNA sequencing (74.58% & 94.59%). In total, 89.19% individuals could be identified in genus level at least.

**Conclusion:** It was revealed that DNA barcoding was useful tool in identifying herbal materials. Both ITS2 and *trnH-psbA* should be incorporated into the standard regions of DNA barcodes for identifying herbal materials.

Keywords: DNA barcodes, herbal materials, ITS2, trnH-psbA, identification

#### Introduction

Traditional medicines are widely used to cure diseases and maintain health in the long history throughout many cultures and regions (Chen et al., 2010; Li et al., 2011b; Pang et al., 2013). Over the past decade, there has been a dramatic increase in the demand for these medicines and pharmaceutical products in both developing and developed countries (Heubl, 2010; Lee et al., 2008). Traditional medicines are extremely rich and diverse in China, including traditional Chinese medicine (TCM) and various ethnomedicines. They account for about 40% of all health care (Lee et al., 2008). However, a severe problem on medicinal market is that many erroneous substitutes and adulterants are traded due to their lower costs or misidentification of species with similar morphological features (Gutteridge and Burns, 2013; Heubl, 2013). There were several cases that the adulterants or substitutes caused serious intoxication and even deaths (Chan and Critchley, 1996; Ernst, 2002; Heubl, 2010; Zhao et al., 2006). Authentication of these traditional medicines and their adulterants is an essential prerequisite to ensure safety, herbal drug quality and therapeutic efficacy (Heubl, 2013). In practice, identification of herbal materials mainly relies on morphological (macroscopic and microscopic) and chemical analyses. But these methods often have obvious weakness (Heubl, 2010).

DNA barcoding, a new approach for species identification using a short, standardized DNA region, has become an important tool in distinguishing species, discovering cryptic species, protecting endangered species and identifying traditional medicine etc. (Chen et al., 2010; Hebert et al., 2003; Kress et al., 2005; Li et al., 2011a; Techen et al., 2014). In animal barcode, the mitochondrial gene cytochrome c oxidase subunit 1 (*CO1*) shows extremely great efficiency. However, the barcode is faced with serious challenge in plant due to its low substitution rates (Hollingsworth et al., 2011). Many botanists have done a lot of researches on selecting appropriate regions and their combinations (Chase et al., 2007; Chen et al., 2010; Hollingsworth et al., 2009, 2011; Kress et al., 2005, 2007; Li et al., 2011a). A two-marker combination of plastid *rbcL* and *matK* was recommended as the core plant barcode, to be supplemented with additional markers such as plastid *trmH-psbA* and nuclear ribosomal internal transcribed spacer (ITS) (Hollingsworth et al., 2009). However, Chinese experts proposed that ITS/ITS2 should be incorporated into the core barcode for seed plants (Chen et al., 2010; Li et al., 2011a; Yao et al., 2010). For herbal materials, it is more difficult to identify because their genome DNA is often seriously degraded in harvesting and prepared progress. So DNA barcodes for herbs should be easy to be amplified and sequenced. It was proposed that ITS2 served as a universal barcode for identifying plant medicine, plus *trmH-psbA* as a complementary barcode (Chen et al., 2010; Han et al., 2013; Pang et al., 2013; Sun and Chen, 2013; Yao et al., 2010). Meanwhile, establishment of open-access molecular databases has laid foundation for DNA barcoding to impart efficient identification of herbal medicine (Gutteridge and Burns, 2013).

The Third Month Fair is a renowned traditional commodity fair in Dali prefecture, Yunnan province (Liu, 2012). This fair is a grand gathering for tourism, trade, culture and sport, possessing important impact on Dali and adjacent regions. Trade of traditional medicines is an important part of this fair. Besides professional druggists, local herbalists are also important partner who usually sell herbal medicines with local characteristics (Zhang et al., 2014). Whole herb is a popular type of plant medicines. It is lack of distinguishing morphological characters generally. So this herbal material is more difficult to be identified. In the present study, we aim to test identified ability of the DNA barcodes in identifying herbal materials in market and explore feasible method on evaluating identification.

#### **Materials and Methods**

#### Medicinal materials and pharmacognostic investigation

During the Third Month Fair in 2013, 116 samples of whole herb were collected from local herbalists in medicinal market (Fig.1). For each sample, about 100g herbal materials without contamination was prepared and put into valve bag as voucher specimens and molecular materials.

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Meanwhile, pharmacognostic investigation was performed to obtain relative information of the sampled materials, including local name, producing area, collecting time and medicinal value. All the specimens were deposited in the Herbarium of Medicinal Plants and Crude Drugs of the College of Pharmacy and Chemistry, Dali University.



**Figure 1:** Herbal market in the Third Month Fair and part herbal materials in the present study. (a. local people in herbal market; b. local herbalists and his herbal materials; c. Luxiancao (QC013), *Pyrola decorata*; d. Guizhencao (QC055), *Bidens pilosa*; e. Xixiancao (QC064), *Siegesbeckia orientalis*; f. Cheqiancao (QC005), *Plantaog major*.

#### Macroscopic identification

Each medicinal material was firstly identified by macroscopic characters on basis of pharmacognostic investigation. The identification was progressed according to relative reference books, mainly including the *China Pharmacopoeia* (Chinese Pharmacopoeia Committee, 2010), the *Drug Standard of Yunnan Province* (Yang, 1996) and the *Atlas of Chinese Herbal Medicine and Ethnomedicine* (Huang, 2005) etc. Putative scientific name of original plant was determined based on the aforementioned books and rechecked using the *Flora of China* (Wu and Peter, 2012).

#### Genomic DNA extraction, amplification and sequencing

Total DNA was extracted from clean herbal materials using modified CTAB method (Doyle and Doyle, 1987). Universal primers for ITS2 and *trnH-psbA*, as well as their corresponding reaction systems and procedures were obtained from previous studies (Chen et al., 2010; Li et al., 2011a; Yao et al., 2010). Purified PCR products were sequenced in both directions with the primers used for PCR amplification on the ABI 3730XL sequencer (Applied Biosystems, USA). Only DNA sequences in accordance with corresponding standard were used for final analysis (Chen et al., 2010; Li et al., 2011a).

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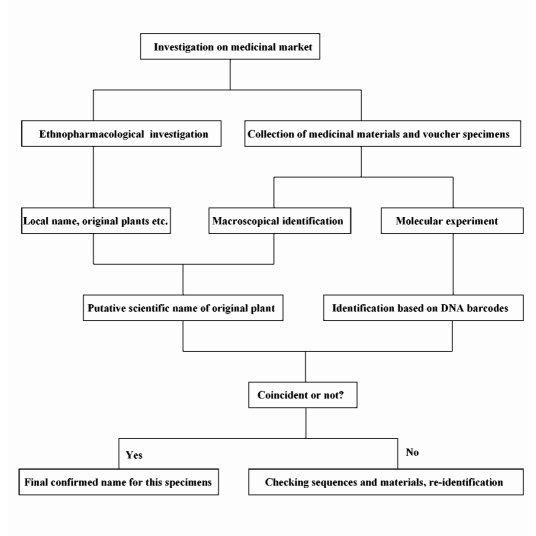


Figure 2: Graph for identifying herbal materials in market though DNA barcodes.

#### Sequencing alignment and molecular identification

Bidirectional sequences were assembled using the program SeqMan Pro 7.1.0 (DNASTAR, Lasergene). They were not aligned due to enormous variation among DNA sequences from different samples. BLAST (basic local alignment search tool), as the most successful way to identify the market samples, was performed for all individuals in the present study (de Boer et al., 2014; Kool et al., 2012). The sequence for each material as a query sequence was submitted and blasted firstly in the DNA Barcoding System for Identifying Herbal Medicine (<a href="http://www.tcmbarcode.cn/en/">http://www.tcmbarcode.cn/en/</a>) (Until April 23, 2014). The species with the nearest match, namely highest similarity was the closest species to species of the query sequence (Chen et al., 2013). In the present study, this species was regarded as possible identified species of original plant for the herbal material though DNA barcodes. But the identified result would be denied if several species were extremely close to query sequence. Meanwhile, result of molecular identification should be coincident to putative species according to macroscopic identification and pharmacognostic investigation, or else the sample would be re-identified or treated as failure identification. The samples with unconfirmed or failure identification above would be further blasted and identified in the National Center for Biotechnology Information (NCBI, <a href="http://www.ncbi.nlm.nih.gov/">http://www.ncbi.nlm.nih.gov/</a>) (Boratyn et al., 2013) (Until April 23, 2014). NCBI's web-based megablast algorithm using the default settings was used to identify the query sequences as proposed (Chen et al., 2013; Kool et al., 2012). Final identification was made manually, taking E-value, maximum identity, number of closely related species represented in the database, as well as distribution of the plants in question into consideration(de Boer et al., 2014). Confirmed scientific name of original plant for identified sample was determined based on the two DNA barcodes and macrosco

#### Results

#### Macroscopic identification and pharmacognositc investigation

In this study, 116 samples were collected from the medicinal market. Among these materials, 111 samples were used for molecular experiment and final analysis except 5 with contaminants or rot. All herbal materials had local names according to pharmacognostic investigation on herbalists. A few of names with mistake in investigation were revised. On the basis of relative reference books, putative scientific name of original plant were determined. Among the 111 samples, original plants of 88 and 15 individuals could be determined in species and genus lever, as well as 8 were

http://dx.doi.org/10.4314/ajtcam.v11i6.7 unconfirmed (Table 2).

#### DNA extraction, PCR amplification and DNA sequencing

Total DNA was extracted out successfully for all samples. For DNA barcodes, success rates of PCR amplification were 98.20% for *trnH-psbA* and 90.99% for ITS2. Dual-band in amplified product was detected in ITS2 region. Only good amplified products were used for further DNA sequencing except that with failure amplification and dual-band. Moreover, success rates of DNA sequencing were relatively low for ITS2 (84.16%) but it was much better for *trnH-psbA* (96.33%). In final, 85 samples for ITS2 (76.58%) and 105 samples for *trnH-psbA* (94.59%) obtained usable DNA sequences that were used for identification though DNA barcodes (Table 1).

Molecular experiment	Experiment result	ITS2	trnH-psbA
PCR amplification		111	111
-	Success	101	109
	Dual-band	2	No
	Failure	8	2
	Success rate	90.99%	98.20%
DNA sequencing		101	109
	Success	85	105
	Overlap pink	4	3
	Failure	14	1
	Success rate	84.16%	96.33%
Final success rate		76.58%	94.59%

Table 1: Statistical data for molecular experiment of DNA barcodes.

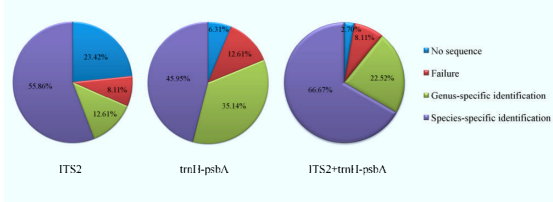


Figure 3: Identified results of ITS2, trnH-psbA and their combination.

#### Identification though DNA barcodes

For ITS2 region, 55.86% and 12.61% samples could be identified in species-specific and genus-specific level. Meanwhile, there were 23.42% individuals without usable sequences and 8.11% could not be correctly identified by ITS2 region. For *trnH-psbA*, species-specific identification was obviously lower than ITS2 (45.95%) but genus-specific identification was relatively high (35.14%). However, individuals without usable sequences for *trnH-psbA* region (6.31%) were much fewer than ITS2 (23.42%). On the basis of coalition analysis of the two DNA barcodes, 98 samples (89.19%) could be identified in genus level at least. All the statistical data were shown in Fig. 3 and Table 2.

Among the 111 samples, original plants of 8 were unconfirmed according to macroscopic identification and pharmacognostic investigation. Most of them could be correctly identified in species-specific (4/8) or genus-specific (2/8) level. Meanwhile, identification based on DNA barcodes successfully revealed adulterants (14/111) in sampled herbal materials. Furthermore, 12 substitutes were also found by DNA barcodes in this study.

There were 98 samples that were successfully identified at genus or species level. Their original plants were relatively rich in Asteraceae (18) and Lamiaceae (14). Some herbal materials were from original plant that had massive relative species. They were extremely difficult to be correctly identified regardless of macroscopic or molecular identification, such as *Adiantum*, *Artemisia* etc.

#### Discussion

Trade of herbal materials is an important part of the Third Month Fair. There were 362 plant medicines at least that were correctly identified and recorded according to incomplete statistics (Zhang et al., 2014). However, massive substitutes and adulterants of genuine herbal materials also arise in herbal market. Whole herb is often difficult to be identified using macroscopic identification because they are usually badly wrinkled or fragmentized. Microscopical and physic-chemical identifications are popular methods as well, but they need relatively high professional quality for authenticator and their discriminatory ability is limited. On the contrary, identification though DNA barcodes, possessing obvious merits compared with the above

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methods becomes a better choice to identify herbal materials (Heubl, 2010). DNA barcoding opens up a unique avenue for the identification of organisms (Li et al., 2011a). It provides a powerful tool to complement chemical analyses for authentication of Chinese medicinal plants and to ensure that herbal materials are not contaminated with ineffective or potentially harmful substitutes or adulterants (Buriani et al., 2012; Heubl, 2010). Herbal materials are generally difficult for molecular identification because their genome DNA usually degraded seriously due to processing treatment (Coghlan et al., 2012). So DNA barcodes for identifying herbal materials should have great usability and variation. Previous data suggested that ITS2 represented the most suitable region for DNA barcoding applications so it was proposed as a novel universal barcode for herbal materials (Chen et al., 2010, 2013; Yao et al., 2010). Until now, the barcode has showed excellent identified power on medicinal materials (Pang et al., 2013; Sun and Chen, 2013). Meanwhile, *trnH-psbA* region, possessing great usability and reliability for species authentication, is also listed as a complementary barcode to ITS2 (Chen et al., 2010, 2013).

In this study, both ITS2 and *trnH-psbA* regions were used to identify herbal materials but they showed obviously difference in usability and identifying ability. Success rates of PCR amplification and DNA sequencing for ITS2 were extremely lower than that for *trnH-psbA* region (Table 1). Only 85 samples got usable DNA sequences for ITS (76.58%); meanwhile, 105 samples were successfully sequenced for *trnH-psbA* (94.59%). Usability of ITS2 here was not such ideal as proposed (Chen et al., 2010; Pang et al., 2013; Yao et al., 2010). But these studies were performed using medicinal plants in which better DNA quality was deposited. There were a few similar studies for identifying herbal materials using the BLAST method of the two barcodes (Kool et al., 2012; Sun and Chen, 2013). So no suitable referenced data could be used here. On the contrary, ITS2 region showed better discriminatory ability than *trnH-psbA* in species-specific identification (55.86% & 45.95%); whereas, they were converse in genus-specific level (12.61% & 35.14%). It revealed that ITS2 possessed better discriminatory ability than *trnH-psbA* region without consideration of its low success rate of DNA sequencing. The two regions possessed obvious advantage and disadvantage each other. In total, 89.19% of all the herbal materials could be correctly identified in genus-specific level at least. It showed that combination of the two DNA barcodes could obviously improve identifying power compared to any single barcode. So we propose that both ITS2 and *trnH-psbA* should be listed as standard regions of DNA barcodes for herbal materials.

In this study, DNA barcodes also showed excellent power in identifying counterfeits and adulterants of certified reference materials. There were 14 counterfeits and 12 adulterants which were correctly identified from sampled materials. For example, "Baihuasheshecao" (QC002) was a common local medicine possessing anti-inflammatory, anti-tumor and antiviral values (Huang, 2005). Its original plant is *Hedyotis diffusa*, but the sample was identified to be counterfeit, namely *Arenaria serpyllifolia*. Moreover, some unknown herbal materials were correctly identified on basis of the DNA barcodes. Original plant of "Maticao" (QC024) was not confirmed according to macroscopic identification and pharmacognostic investigation which was successfully identified by DNA barcodes, namely *Centella asiatica*. So the method is reliable tool for identifying herbal materials, their substitutes and adulterants, as well as unknown medicines (Techen et al., 2014).

Although DNA barcode is beneficial to identify herbal materials it still needs assistance from traditional methods, including herbal information from herbalists and macroscopic identification etc. In fact, two aspects of studies are equally important. On the one hand, pharmacognostic investigation or communication with druggists can afford rich information for herbal materials. But the information may be incorrect. So further macroscopic identification using reference book, such as the *Chinese Pharmacopoeia* is necessary that also provides formal information. On the other hand, identification though DNA barcodes give confirmed result or clue. Basic method of identification though DNA barcodes is BLAST searches for query sequences in open-access database, such as the DNA Barcoding System for Identifying Herbal Medicines (Chen et al., 2013). This is a specific platform for identifying traditional medicines by DNA barcodes regions which affords ITS2 and *trnH-psbA* for medicinal plants. The database is very useful for traditional medicines from China and can give identified results quickly. But it is probably inapplicable to overseas and folk medicines. Moreover, NCBI provides much more comprehensive identification but it includes some DNA sequences with misidentification (Bridge et al., 2003; Federhen, 2012). Meanwhile, intra-species genetic distance may be higher than inter-species genetic distance especially in some complex and widespread species. So species of the nearest match is not necessarily correct identification (de Boer et al., 2014; Heubl, 2010, 2013). Identification should be made manually taking E-value, maximum identify, number of closely related species represented in the database, as well as distribution of the plant in question into consideration.

There are still many problems to be solved for identifying herbal materials in market though DNA barcodes. Firstly, plant medicines from fern were difficult to be correctly identified, such as "Zhuzongcao" (QC002). DNA sequences of these groups were very absent and they were apt to be identified in error. These samples were generally identified in genus-specific level. Secondly, some genus is composed of multitudinous species and many species are used as medicines, such as *Artemisa* L.(Heubl, 2013). It is also difficult to identify herbal materials from these genera. Thirdly, some folk or local medicines could not be identified using DNA barcodes, especially using the DNA Barcoding System for Identifying Herbal Medicines. So we propose that relative organization and researchers should further supply DNA sequences for special groups and improving identification of submitted sequences.

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Table 2: Herbal materials used for identification though DNA barcodes in this study and their identified information (I).

Sample	Local name of	Putative scientific name	Identified result by	Identified result by	Confirmed scientific	Family name
ID	traditional medicine	of original plant	ITS2	trnH-psbA	name of original plant	
QC002	Zhuzongcao	Adiantum capillus-veneris L.	No sequence	Adiantum L.	Adiantum L.	Adiantaceae
QC003 **	Baihuasheshecao	Hedyotis diffusa Willd.	Arenaria serpyllifolia L.	Arenaria serpyllifolia L.	Arenaria serpyllifolia L.	Caryophyllaceae
	Qianzhenwanxiancao	Stellaria yunnanensis	Wahlenbergia	Failure	Wahlenbergia	Campanulaceae
QC004 **	Quanzarenwananeae	Franch.	marginata (Thunb.) A.DC	Turide	marginata (Thunb.) A.DC	Campanaraceae
QC005 *	Cheqiancao	Plantago asiatica L.	Plantago major L.	Plantago major L.	Plantago major L.	Plantaginaceae
	Biandaxiuqiu	Hemiphragma	Hemiphragma	Failure	Hemiphragma	Scrophulariaceae
QC007		heterophyllum wall.	heterophyllum wall.		heterophyllum wall.	
QC008	Tougucao	Gaultheria leucocarpa Bl.	Gaultheria leucocarpa Bl.	Gaultheria Kalm ex L.	Gaultheria leucocarpa Bl.	Ericaceae
QC009	Jinxiancao	Lysimachia christinae Hance	Lysimachia L.	Lysimachia L.	Lysimachia L.	Primulaceae
QC010	Shuizhuzong	Adiantum L.	No sequence	Adiantum L.	Adiantum L.	Adiantaceae
	Guanyincao	Reineckia carnea Kunth	No sequence	Reineckia carnea	Reineckia carnea	Liliaceae
QC011	Ž		•	Kunth	Kunth	
	Huixincao	Rhodobryum giganteum	Rhodobryum Schimp	Rhodobryum	Rhodobryum	Bryaceae
QC012		Par.		giganteum Par.	giganteum Par.	
0.001.0	Luxiancao	Pyrola decorate H. Andr.	Pyrola decorate H.	Pyrola L.	Pyrola decorate H.	Pyrolaceae
QC013	г	D	Andr.	n	Andr.	D : 11
QC014	Feixincao	Botrychium lanuginosum Wall.	No sequence	Botrychium Sw.	Botrychium Sw.	Botrychiaceae
QC015	Shiwei	Pyrrosia petiolosa (Christ) Ching	No sequence	Pyrrosia petiolosa (Christ) Ching	Pyrrosia petiolosa (Christ) Ching	Polypodiaceae
-	Yimucao	Leonurus japonicas	Leonurus japonicas	Leonurus japonicas	Leonurus japonicas	Lamiaceae
QC016		Houtt.	Houtt.	Houtt.	Houtt.	
QC017	Huangjingzi	Vitex negundo L.	Vitex negundo L.	Vitex negundo L.	Vitex negundo L.	Verbenaceae
0.0010	Bohe	Mentha Canadensis L.	Mentha Canadensis L.	Mentha Canadensis	Mentha Canadensis L.	Lamiaceae
QC018	7.00 and ar - !!	Unacafiamed	No seemens	L.	Tuniatus	Lilianna
QC019	Zuogushengjingcao	Unconfirmed	No sequence	Tupistra wattii (C.B.Clarke) Hook.	Tupistra wattii (C.B.Clarke) Hook.	Liliaceae
QC019	Banzhilian	Scutellaria barbata D.	Scutellaria barbata D.	Scutellaria barbata	Scutellaria barbata D.	Lamiaceae
QC020	Danzinnan	Don	Don	D. Don	Don	Laimaceae
QC020	Shijiaocao	Boenninghausenia	Failure	Failure	Failure	_
QC021	Sinjiuocuo	sessilicarpa Levl.	T diffare	Tulluic	Tullulo	
QC022	Bianxu	Polygonum aviculare L.	Polygonum L.	No sequence	Polygonum L.	Polygonaceae
-	Sanxuedan	Unconfirmed	Peperomia blanda	Failure	Peperomia blanda	Piperaceae
QC023			(Jacquin) Kunth		(Jacquin) Kunth	_
	Maticao (Jinxiancao)	Unconfirmed	Centella asiatica (L.)	Centella asiatica	Centella asiatica (L.)	Apiaceae
QC024			Urban	(L.) Urban	Urban	
QC025 **	Muguajisheng	Scurrula parasitica L.	Failure	Taxillus chinensis	Taxillus chinensis (DC)	Loranthaceae
ጥጥ				(DC) Danser	Danser	

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Table 2: Herbal materials used for identification though DNA barcodes in this study and their identified information (II).

Sample ID	Local name of traditional medicine	Putative scientific name of original plant	Identified result by ITS2	Identified result by trnH-psbA	Confirmed scientific name of original plant	Family name
QC026	Luticao	Pyrola decorate Andr.	Pyrola decorate Andr.	Pyrola L.	Pyrola decorate Andr.	Pyrolaceae
QC027	Xiakucao	Prunella vulgaris L.	Prunella vulgaris L.	Prunella vulgaris L.	Prunella vulgaris L.	Lamiaceae
QC028	Longdancao	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentianaceae
QC029 **	Muguajisheng	Scurrula parasitica L.	Failure	Taxillus chinensis (DC) Danser	Taxillus chinensis (DC) Danser	Loranthaceae
	Pugongying	Taraxacum mongolicum	Taraxacum mongolicum	Taraxacum mongolicum HanMazz.	Taraxacum mongolicum	Asteraceae
QC030	Yinchenhao	HanMazz. Artemisia	HanMazz.  Artemisia capillaris	Artemisia capillaris	HanMazz.  Artemisia capillaris	Asteraceae
QC031		capillaris Thunb.	Thunb.	Thunb.	Thunb.	
QC032 *	Aiye	Artemisia argyi Levl. et Van.	Artemisia lavandulaefolia DC. Prodr.	Artemisia L.	Artemisia lavandulaefolia DC. Prodr.	Asteraceae
QC032	Xiakucao	Prunella vulgaris L.	Prunella vulgaris L.	Prunella vulgaris L.	Prunella vulgaris L.	Lamiaceae
QC034	Shiwei	Pyrrosia Mirbel	No sequence	Pyrrosia petiolosa (Christ) Ching	Pyrrosia petiolosa (Christ) Ching	Polypodiaceae
QC035	Yinyanghuo Jingjie	Epimedium L. Nepeta tenuifolia	Epimedium L. Nepeta tenuifolia	Epimedium L. Nepeta tenuifolia Benth.	Epimedium L. Nepeta tenuifolia	Berberidaceae Lamiaceae
QC036		Benth.	Benth.		Benth.	
00027**	Yinyanghuo	Epimedium L.	Astilbe rivularis BuchHam. ex	Astilbe rivularis BuchHam.exD.Don	Astilbe rivularis BuchHam. ex	Saxifragaceae
QC037**	Yimucao	Leonurus	D.Don Leonurus japonicus	Leonurus japonicus	D.Don Leonurus japonicus	Lamiaceae
QC038	Yexiahua	japonicus Houtt. Ainsliaea	Houtt.  Ainsliaea pertyoides	Houtt. Faliure	Houtt.  Ainsliaea pertyoides	Asteraceae
QC039	Rendongteng	pertyoides Franch. Lonicera japonica	Franch. Lonicera	Lonicera L.	Franch.  Lonicera	Caprifoliaceae
QC040 *		Thunb.	macranthoides HandMazz.		<i>macranthoides</i> HandMazz.	
QC041	Dacheqian Fengweicao	Plantago major L. Pteris nervosa	Plantago major L. No sequence	Plantago major L. Pteris L.	Plantago major L. Pteris L.	Asteraceae Pteridaceae
QC042 QC043	Maweihuanglianye Juanbai	Thunb. Thalictrum L. Selaginella tamariscina	Thalictrum L. Selaginella pulvinata Maxim.	Thalictrum L. Selaginella pulvinata Maxim.	Thalictrum L. Selaginella pulvinata Maxim.	Ranunculaceae Selaginellaceae
QC044 *	Daogouci	Spring Rubus delavayi	Rubus L.	Rubus L.	Rubus L.	Rosaceae
QC045	Litoucao	Franch.  Viola inconspicua	Viola philippica Cav.	Viola philippica Cav.	Viola inconspicua B1.	Violaceae
QC046	Xianhecao	Bl. Agrimonia pilosa	Agrimonia pilosa	Agrimonia L.	Agrimonia pilosa	Rosaceae
QC047		Ledeb.	Ledeb.		Ledeb.	
QC048	Jinwaer(Waercao)	Carpesium divaricatum Sieb. et Zucc.	Carpesium divaricatum Sieb. et Zucc.	Carpesium divaricatum Sieb. et Zucc.	Carpesium divaricatum Sieb. et Zucc.	Asteraceae

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Table 2: Herbal materials used for identification though DNA barcodes in this study and their identified information (III).

Sample ID	Local name of traditional medicine	Putative scientific name of original plant	Identified result by ITS2	Identified result by trnH-psbA	Confirmed scientific name of original plant	Family name
QC049	Tiexianjue	Adiantum L.	No sequence	Cheilanthes Sw.	Cheilanthes Sw.	Sinopteridaceae
QC050	Baihao	Artemisia sieversiana	Artemisia argyi Levl.	Artemisia L.	Artemisia argyi Levl.	Asteraceae
*		Ehrh. ex Willd.	et Van.		et Van.	
QC051	Xiangyejisheng	Unconfirmed	No sequence	Failure	Failure	-
	Jinyinhua	Lonicera japonica	Lonicera	Lonicera L.	Lonicera	Caprifoliaceae
QC052		Thunb.	macranthoides		macranthoides	
*			HandMazz.		HandMazz.	
QC053	Cujiangcao	Oxalis corniculata L.	Oxalis corniculata L.	Oxalis corniculata L.	Oxalis corniculata L.	Oxalidaceae
QC054	Xiaofengweicao	Asplenium L.	No sequence	Asplenium L.	Asplenium L.	Aspleniaceae
QC055	Guizhencao	Bidens pilosa L.	Bidens pilosa L.	Bidens L.	Bidens pilosa L.	Asteraceae
QC056	Liangmianzhen	Solanum torvum Sw.	No sequence	Solanum torvum Sw.	Solanum torvum Sw.	Solanaceae
QC057	Heihanlian	Eclipta prostrate L.	Eclipta prostrate L.	Eclipta prostrate L.	Eclipta prostrate L.	Asteraceae
	Laoguancao	Geranium nepalense	Geranium nepalense	Geranium nepalense	Geranium nepalense	Geraniaceae
QC058		Sw.	Sw.	Sw.	Sw.	
QC059	Kudingding	Unconfirmed	Solanum nigrum L.	Solanum nigrum L.	Solanum nigrum L.	Solanaceae
QC060	Wuzhuajinlong	Cayratia japonica	Tetrastigma (Miq.)	Tetrastigma (Miq.)	Tetrastigma (Miq.)	Vitaceae
**		(Thunb.) Gagnep.	Planch.	Planch.	Planch.	
	Pahao	Elsholtzia rugulosa	Elsholtzia Willd.	Elsholtzia Willd.	Elsholtzia Willd.	Lamiaceae
QC061		Hemsl.				
QC062	Zhuzongcao	Adiantum L.	No sequence	Adiantum L.	Adiantum L.	Adiantaceae
	Jiejiecao	Equisetum	No sequence	No sequence	No sequence	-
QC063		ramosissimum Desf.				
	Xixiancao	Siegesbeckia	No sequence	Siegesbeckia	Siegesbeckia	Asteraceae
QC064		orientalis L.		orientalis L.	orientalis L.	
	Digandou	Rorippa indica (L.)	Rorippa indica (L.)	Failure	Rorippa indica (L.)	Brassicaceae
QC065		Hiern	Hiern		Hiern	
QC066	Dengxincao	Juncus effuses L.	No sequence	No sequence	No sequence	-
	Chaihu	Bupleurum L.	Bupleurum chinense	Bupleurum L.	Bupleurum chinense	Apiaceae
QC067			DC. Prodr.		DC. Prodr.	
	Xixin	Asarum himalaicum	Asarum himalaicum	No sequence	Asarum himalaicum	Aristolochiaceae
QC068		Hook. et Thoms.	Hook. et Thoms		Hook. et Thoms	
	Yinchenhao	Artemisia capillaris	Artemisia capillaris	Artemisia capillaris	Artemisia capillaris	Asteraceae
QC069		Thunb.	Thunb.	Thunb.	Thunb.	
QC070	Xuelian	Saussurea DC.	Saussurea medusa Maxim.	Saussurea DC.	Saussurea medusa Maxim.	Asteraceae
-	Jinyinhua	Lonicera japonica	Lonicera	Lonicera	Lonicera	Caprifoliaceae
QC071	•	Thunb.	macranthoides	macranthoides	macranthoides	-
*			HandMazz.	HandMazz.	HandMazz.	

# Zhang et al., Afr J Tradit Complement Altern Med. (2014) 11(6):66-76 <a href="http://dx.doi.org/10.4314/ajtcam.v11i6.7">http://dx.doi.org/10.4314/ajtcam.v11i6.7</a> Table 2 Herbal materials used for identification though DNA barcodes in this study and their identified information (IV).

Sample	Local name of	Putative scientific	Identified result by	Identified result by	Confirmed scientific	Family name
ID	traditional medicine	name of original plant	ITS2	trnH-psbA	name of original plant	
	Yuxingcao	Houttuynia cordata	Houttuynia cordata	Houttuynia cordata	Houttuynia cordata	Saururaceae
QC072	Tumigeuo	Thunb.	Thunb.	Thunb.	Thunb.	Suararaceae
	Chaihu (Wild)	Bupleurum	Bupleurum L.	Bupleurum L.	Bupleurum L	Apiaceae
		marginatum Wall. ex				
QC073		DC.				~
QC075	Jinyinhua	Lonicera japonica Thunb.	Lonicera macranthoides	Lonicera L.	Lonicera macranthoides	Caprifoliaceae
*		i iiuiio.	HandMazz.		HandMazz.	
	Shihu	Dendrobium nobile	No sequence	No sequence	No sequence	_
QC076		Lindl.	- 10 0-4			
QC077	Qinghao	Artemisia annua L.	Artemisia L.	Artemisia L.	Artemisia L.	Asteraceae
QC079	Celan	Eupatorium fortune	Polygonum chinense	Polygonum chinense	Polygonum chinense	Polygonaceae
**	Y' 1 ' 1	Turcz.	L.	L.	L.	
QC080	Jinzhongyinchen	Siphonostegia chinensis Benth.	No sequence	Failure	Failure	-
QC080 QC081	Zihuadiding	Viola philippica Cav.	Viola philippica Cav.	Viola philippica Cav.	Viola philippica Cav.	Violaceae
QC081	Xiangye	Laurus nobilis L.	Failure	Laurus nobilis L.	Laurus nobilis L.	Lauraceae
QC083	Nanxixin	Asarum forbesii	Asarum himalaicum	No sequence	Asarum himalaicum	Aristolochiaceae
*	(Tuxixin)	Maxim.	Hook. et Thoms.	•	Hook. et Thoms.	
QC084	Xiaofeixincao	Unconfirmed	No sequence	Botrychium Sw.	Botrychium Sw.	Botrychiaceae
QC085	Shexucao	Ophioglossum L.	No sequence	Ophioglossum L.	Ophioglossum L.	Ophioglossaceae
QC086	Dengzhanhua	Erigeron breviscapus HandMazz.	Erigeron breviscapus HandMazz.	Erigeron L.	Erigeron breviscapus HandMazz.	Asteraceae
QC080	Jiaogulan	Gynostemma	No sequence	Gynostemma	Gynostemma	Cucurbitaceae
	Juoguran	pentaphyllum Makino	110 sequence	pentaphyllum	pentaphyllum Makino	Cacaronaceae
QC087		I I		Makino	r · · · · r · · · · · · · · · · · · · ·	
	Banzhilian	Scutellaria barbata	Scutellaria barbata D.	Scutellaria barbata	Scutellaria barbata D.	Lamiaceae
QC088	~.	D. Don	Don	D. Don	Don	
00000	Sheyancao	Saussurea	Saussurea DC.	Saussurea DC.	Saussurea DC.	Asteraceae
QC089	Xiaoshancha	romuleifolia Franch. Elsholtzia bodinieri	Failure	Elsholtzia Willd.	Elsholtzia Willd.	Lamiaceae
QC090	Alaoshancha	Vaniot bouilleri	ranuic	Eishotizia Willa.	Eisnouzia Willa.	Lamaccac
Quoyu	Sanxuedan	Peperomia blanda	No sequence	Failure	Failure	-
QC091		(Jacquin) Kunth	•			
	Guichuixiao	Leycesteria Formosa	Failure	Failure	Failure	-
QC092	*	Wall.	<i>a</i> , , , , , , , , , , , , , , , , , , ,	a	a	
QC093 **	Lvticao	Caltha palustris L.	Centella asiatica (L.) Urban	Centella asiatica (L.) Urban	Centella asiatica (L.) Urban	Apiaceae
QC094	Cebaiye	Platycladus orientalis	No sequence	Lycopodium	Lycopodium	Lycopodiaceae
**	Coarye	(L.) Franco	140 sequence	<i>japonicum</i> Thunb.	<i>japonicum</i> Thunb.	Lycopouraceae
	Aiye	Artemisia argyi Levl.	Artemisia argyi Levl.	Artemisia L.	Artemisia argyi Levl.	Asteraceae
QC095		et Van.	et Van.		et Van.	

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Table 2: Herbal materials used for identification though DNA barcodes in this study and their identified information (V).

Sample ID	Local name of traditional medicine	Putative scientific name of original plant	Identified result by ITS2	Identified result by trnH-psbA	Confirmed scientific name of original plant	Family name
QC096 *	Choulingdan	Laggera pterodonta (DC.) Benth.	Laggera pterodonta (DC.) Benth.	Laggera pterodonta (DC.) Benth.	Laggera pterodonta (DC.) Benth.	Asteraceae
0.0007	Xianglingcao	Crotalaria ferruginea	Crotalaria L.	Crotalaria L.	Crotalaria L.	Fabaceae
QC097	Longdancao	Grah. ex Benth.  Gentiana rigescens	Gentiana rigescens	Gentiana rigescens	Gentiana rigescens	Gentianaceae
QC098	Longdaneao	Franch.	Franch.	Franch.	Franch.	Gentianaceae
QC099	Longdancao	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentiana rigescens Franch.	Gentianaceae
QC100 **	Sifanghao	Elsholtzia blanda Benth.	Salvia plebeia R. Br.	Salvia plebeia R. Br.	Salvia plebeia R. Br.	Lamiaceae
QC101	Dafeixincao	Unconfirmed	No sequence	Botrychium Sw.	Botrychium Sw.	Botrychiaceae
QC102	Tianjihuang	Unconfirmed	Failure	No sequence	Failure	-
	Gangbangui	Polygonum	Polygonum	Polygonum L.	Polygonum	Polygonaceae
QC103		perfoliatum L.	perfoliatum L.		perfoliatum L.	
00101	Baitouweng	Pulsatilla chinensis	Carpesium	Carpesium L.	Carpesium	Asteraceae
QC104 **		(Bunge) Regel	divaricatum Sieb. et Zucc.		divaricatum Sieb. et Zucc.	
QC106 **	Xuelimei	Gentiana rhodantha	Metagentiana	Failure	Metagentiana	Gentianaceae
	V:	Franch.	rhodantha T.N. Ho	O.:	rhodantha T.N. Ho	T!
QC107 **	Xiangru	Elsholtzia ciliate (Thunb.) Hyland	Origanum vulgare L.	Origanum vulgare L.	Origanum vulgare L.	Lamiaceae
QC108	Dashiwei	Pyrrosia Mirbel	No sequence	Pyrrosia Mirbel	Pyrrosia Mirbel	Polypodiaceae
QC109 **	Xiangrucao	Elsholtzia ciliate (Thunb.) Hyland	Origanum vulgare L.	Origanum vulgare L.	Origanum vulgare L.	Lamiaceae
	Xianglingcao	Crotalaria ferruginea	Failure	Failure	Failure	_
QC110	88	Grah. ex Benth.				
	Zisu	Perilla frutescens (L.)	Perilla frutescens (L.)	Perilla frutescens	Perilla frutescens (L.)	Lamiaceae
QC111		Britt.	Britt.	(L.) Britt.	Britt.	
	Qingyedan	Swertia mileensis	Swertia macrosperma	Swertia	Swertia macrosperma	Gentianaceae
QC112 *		T.N.Ho et W. L. Shi	C.B. Clarke	macrosperma C.B.	C.B. Clarke	
	Huashicao	Dunnagia Mirhal	No soquence	Clarke Failure	Failure	
QC113 QC114	Huasnicao Hanzhuzongcao	Pyrrosia Mirbel. Adiantum L.	No sequence Failure	Failure Failure	Failure Failure	-
QC114 QC115	Fanxieye	Senna Miller	Senna Miller	Senna Miller	Senna Miller	- Fabaceae
QC116	Qumai	Dianthus superbus L.	Dianthus chinensis L.	Dianthus chinensis	Dianthus chinensis L.	Caryophyllaceae
*	£	= super em Di		L.	_ :	j op.i.j incode

Note: \* substitute of herbal material; \*\* adulterant.

#### Conclusions

In the present study, DNA barcodes showed great identified power for whole herb. ITS2 possessed better discriminatory ability than *trnH-psbA* region but its success rates of PCR amplification and DNA sequencing were obviously lower. Both ITS2 and *trnH-psbA* should be listed as standard regions of DNA barcodes for herbal medicines. The DNA Barcoding System for Identifying Herbal Medicine is an efficient identified platform for Chinese traditional medicines. But DNA sequences of the folk medicines should be further supplemented and improved. Moreover, reference databases, such as NCBI, must construct error-corrected mechanism that eliminates DNA sequences with mistaken identification (Heubl, 2013). Although molecular identification often fails to assign individuals to species it is a helpful tool in providing clues for identifying herbal materials that lack morphological features for species identification.

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#### References

- 1. Boratyn, G.M., Camacho, C., Cooper, P.S., Coulouris, G., Fong, A., Ma, N., Madden, T.L., Matten, W.T., McGinnis, S.D. and Merezhuk, Y. (2013). BLAST: a more efficient report with usability improvements. Nucl. Acids. Res. 41, 29-33.
- 2. Bridge, P.D., Roberts, P.J., Spooner, B.M. and Panchal, G. (2003). On the unreliability of published DNA sequences. New Phytol. 160, 43-48.
- 3. Buriani, A., Garcia-Bermejo, M.L., Bosisio, E., Xu, Q., Li, H., Dong, X., Simmonds, M.S., Carrara, M., Tejedor, N. and Lucio-Cazana, J. (2012). Omic techniques in systems biology approaches to traditional Chinese medicine research: present and future. J. Ethnopharm. 140, 535-544.
- 4. Chan, T.Y. and Critchley, J.A. (1996). Usage and adverse effects of Chinese herbal medicines. Hum. Exp. Toxicol. 15, 5-12.
- 5. Chase, M.W., Cowan, R.S. and Hollingsworth, P.M. (2007). A proposal for a standardised protocol to barcode all land plants. Taxon 56, 295-299.
- 6. Chen, S.L., Yao, H., Han, J.P., Xin, T.Y. and Pang, X.H. (2013). Principles for molecular identification of traditional Chinese materia medica using DNA barcoding. China J. Chin. Mater. Med. 38, 141-148.
- 7. Chen, S.L., Yao, H., Han, J.P., Liu, C., Song, J.Y., Shi, L.C., Zhu, Y.J., Ma, X.Y., Gao, T., Pang, X.H., Luo, K., Li, Y. and Li, X.W. (2010). Validation of the ITS2 region as a novel DNA barcode for identifying medicinal plant species. PLoS ONE 5, e8613.
- 8. Coghlan, M.L., Haile, J., Houston, J., Murray, D.C., White, N.E., Moolhuijzen, P., Bellgard, M.I. and Bunce, M. (2012). Deep sequencing of plant and animal DNA contained within traditional Chinese medicines reveals legality issues and health safety concerns. PLoS Genet. 8, e1002657.
- 9. Chinse Pharmacopoeia Committee (2010). Chinese Pharmacopoeia (1st part). China Medical Science and Technology Press, Beijing.
- 10. de Boer, H.J., Ouarghidi, A., Martin, G., Abbad, A., Kool, A. (2014). DNA Barcoding reveals limited accuracy of identifications based on folk taxonomy. PLoS ONE 9, e84291.
- 11. Doyle J.J and Doyle, J.L. (1987). A rapid DNA isolation procedure from small quantities of fresh leaf tissue. Phytochem. 19, 11-15.
- 12. Ernst, E. (2002). Adulteration of Chinese herbal medicines with synthetic drugs: a systematic review. J. Inter. Med. 252, 107-113.
- 13. Federhen, S. (2012). The NCBI taxonomy database. Nucl. Acids Res. 40, 136-143.
- 14. Gutteridge, A. and Burns, M. (2013). The application of DNA molecular approaches for the identification of herbal medicinal products. J. Assoc. Public. Anal. 41, 53-66.
- 15. Han, J.P., Zhu, Y.J., Chen, X.C., Liao, B.S., Yao, H., Song, J.Y., Chen, S.L. and Meng, F.Y. (2013). The short ITS2 sequence serves as an efficient taxonomic sequence tag in comparison with the full-length ITS. Biomed. Resear. Inter. 2013, 741476.
- 16. Hebert, P.D.N., Cywinska, A., Ball, S.L. and deWaard, J.R. (2003). Biological identifications through DNA barcodes. Proceed. Roy. Soc. B 270, 313-327.
- 17. Heubl, G. (2010). New aspects of DNA-based authentication of Chinese medicinal plants by molecular biological techniques. Planta Med. 76, 1963-1974.
- 18. Heubl, G. (2013). DNA-based authentication of TCM-plants: current progress and future perspectives. Springer Vienna, Kassel, Germany.
- 19. Hollingsworth, P.M., Forrest, L.L., Spouge, J.L., Hajibabaei, M., Ratnasingham, S., van der Bank, M., Chase, M.W., Cowan, R.S., Erickson, D.L. and Fazekas, A.J. (2009). A DNA barcode for land plants. Proc. Natl. Acad. Sci. USA 106, 12794-12797.
- 20. Hollingsworth, P.M., Graham, S.W. and Little, D.P. (2011). Choosing and using a plant DNA barcode. PLoS ONE 6, e19254.
- 21. Huang, L.Q. (2005). Atlas of Chinese herbal medicine and ethnomedicine. Peking University Medical Press, Beijing.
- 22. Kool, A., de Boer, H.J., Krüger, Å., Rydberg, A., Abbad, A., Björk, L. and Martin, G. (2012). Molecular identification of commercialized medicinal plants in southern Morocco. PLoS ONE 7, e39459.
- 23. Kress, W.J. and Erickson, D.L. (2007). A two-locus global DNA barcode for land plants: the coding rbcL gene complements the non-coding trnHpsbA spacer regionhe. PLoS ONE 6, e508.
- 24. Kress, W.J., Wurdack, K.J., Zimmer, E.A., Weigh, L.A., Janzen and D.H. (2005). Use of DNA barcodes to identify flowering plants. Proc. Natl. Acad. Sci. USA 102, 8369-8374.
- 25. Lee, S.W., Xiao, C.J. and Pei, S.J. (2008). Ethnobotanical survey of medicinal plants at periodic markets of Honghe Prefecture in Yunnan Province, SW China. J. Ethnopharm. 117, 362-377.
- 26. Li, D.Z., Gao, L.M., Li, H.T., Wang, H., Ge, X.J., Liu, J.Q., Chen, Z.D., Zhou, S.L., Chen, S.L. and Yang, J.B. (2011). Comparative analysis of a large dataset indicates that internal transcribed spacer (ITS) should be incorporated into the core barcode for seed plants. Proc. Natl. Acad. Sci. USA 108, 19641-19646.
- 27. Li, M., Cao, H., But, P.P.H. and Shaw, P.C. (2011). Identification of herbal medicinal materials using DNA barcodes. J. Syst. Evol. 49, 271-283.
- 28. Liu, B. (2012). Studies on impact factors of the Third Month Fair changes in Dali Bai prefecture (D). Guizhou University of Finance and Economics, Guiyang.
- 29. Pang, X.H., Shi, L.C., Song, J.Y., Chen, X.C. and Chen, S.L. (2013). Use of the potential DNA barcode ITS2 to identify herbal materials. J. Nat. Med. 67, 571-575.
- 30. Sun, Z. and Chen, S. (2013). Identification of cortex herbs using the DNA barcode nrITS2. J. Nat. Med. 67, 296-302.
- 31. Techen, N., Parveen, I., Pan, Z. and Khan, I.A. (2014). DNA barcoding of medicinal plant material for identification. Curr. Opin. Biotechnol. 25, 103-110.
- 32. Wu, Z.Y. and Peter, R.H. (2012). Flora of China. Science Press, Beijing; Missouri Botanical Garden Press, St. Louis.
- 33. Yang, C.S. (1996). Drug standard of Yunnan province. Yunnan University Press, Kunming.
- 34. Yao, H., Song, J.Y., Liu, C., Luo, K., han, J.P., Li, Y., Pang, X.H., Xu, H.X., Zhu, Y.J., Xiao, P.G. and Chen, S.L. (2010). Use of ITS2 region as the universal DNA barcode for plants and animals. PLoS ONE 5, e13102.
- 35. Zhang, D.Q., Duan, L.Z. and Zhou, N. (2014). Market survey on traditional medicine of the Third Month Fair in Dali perfecture in Yunnan province, southwest China. Afr. J. Tradit. Complement. Altern. Med. 11, 377-401.
- 36. Zhao, Z.Z., Hu, Y.N., Liang, Z.T., Yuen, J.P.S., Jiang, Z.H. and Leung, K.S.Y. (2006). Authentication is fundamental for standardization of Chinese medicines. Planta Med. 72, 865-874.