

Molecular Phylogeny of the *Lactuca* Alliance (Cichorieae Subtribe Lactucinae, Asteraceae) with Focus on Their Chinese Centre of Diversity Detects Potential Events of Reticulation and Chloroplast Capture

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Abstract

The first comprehensive molecular phylogenetic reconstruction of the Cichorieae subtribe Lactucinae is provided. Sequences for two datasets, one of the nuclear rDNA ITS region, the other of five concatenated non-coding chloroplast DNA markers including the *petD* region and the *psbA-trnH*, 5'trnL(UAA)-trnF, rpl32-trnL(UAG) and trnQ(UUG)-5'rps16 spacers, were, with few exceptions, newly generated for 130 samples of 78 species. The sampling spans the entire subtribe Lactucinae while focusing on its Chinese centre of diversity; more than 3/4 of the Chinese Lactucinae species are represented. The nuclear and plastid phylogenies inferred from the two independent datasets show various hard topological incongruences. They concern the internal topology of major lineages, in one case the placement of taxa in major lineages, the relationships between major lineages and even the circumscription of the subtribe, indicating potential events of ancient as well as of more recent reticulation and chloroplast capture in the evolution of the subtribe. The core of the subtribe is clearly monophyletic, consisting of the six lineages, Cicerbita, Cicerbita II, Lactuca, Melanoseris, Notoseris and Paraprenanthes. The Faberia lineage and the monospecific Prenanthes purpurea lineage are part of a monophyletic subtribe Lactucinae only in the nuclear or plastid phylogeny, respectively. Morphological and karyological support for their placement is considered. In the light of the molecular phylogenetic reconstruction and of additional morphological data, the conflicting taxonomies of the Chinese Lactuca alliance are discussed and it is concluded that the major lineages revealed are best treated at generic rank. An improved species level taxonomy of the Chinese Lactucinae is outlined; new synonymies and some new combinations are provided.

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Introduction

The Lactuca Alliance

Lettuce (Lactuca sativa L.) is the economically most important crop of the tribe Cichorieae, and Lactuca is one of its widest known genera. With almost all members of that tribe, Lactuca shares the combined presence of latex and homogamous capitula with usually ligulate 5-toothed flowers. Lactuca is also the namegiving member of one of the larger groups of the tribe, which is treated today as the subtribe Lactucinae [1]. In its revised circumscription the Lactucinae comprise about 230 species, distributed in Europe, Africa, Asia and North America [1], with a preference of montane habitats. Many of them are mesic tall forbs, many others are perennial herbs of other kinds, among them the only scandent herbs present in the Cichorieae, or rosette herbs and acaulescent herbs, and more rarely they are xeric subshrubs and annual herbs. This subtribe constitutes the youngest branch in the larger of the two core groups of the Cichorieae, its divergence is estimated to

have taken place c. 15–4 Ma ago during the Middle Miocene to Early Pliocene [2–3].

The taxonomy of no other alliance of the tribe has faced so many controversies over the last 200 years than that of *Lactuca* and its presumed allies. This pertains to the circumscription and systematic position of the *Lactuca* alliance within the tribe as well as, and even much more so, to the generic classification of its members. In the 19th century, the *Lactuca* alliance, although sometimes recognised as a separate subtribe [4], was mostly included in the subtribe Crepidinae, as was done also by Hoffmann [5], whose treatment became influential and the basis for most of the 20th century flora treatments. Also in the first two important 20th century classifications of the Cichorieae, by Stebbins [6] and Jeffrey [7], the Lactuca alliance was treated as a subgroup of the Crepidinae or of a corresponding entity: the first author treated it as the *Prenanthes-Lactuca* line of subtribe Crepidinae, the second as the *Prenanthes* series of the *Crepis* group.

Only towards the end of the 20th century, the Lactuca alliance was recognised as a separate subtribe Lactucinae by Bremer [8], after his morphological phylogenetic analysis of the tribe had revealed the Crepidinae to be polyphyletic. Bremer therefore divided the Crepidinae into the three subtribes Lactucinae, Crepidinae s.str. and Sonchinae, which were largely maintained by Lack [9]. Based on a nrITS phylogeny of the Cichorieae, which remarkably well agrees with the results inferred from chloroplast DNA restriction site variation [10], Kilian & al. [1] maintained these three subtribes of Bremer among the 11 subtribes they recognised in the tribe, but narrowed down the circumscription of subtribe Lactucinae compared to Bremer [8] and Lack [9]. Kilian & al. [1] excluded from subtribe Lactucinae the genera Prenanthes s.l., which was characterised as a dust-bin of various unrelated elements by Kilian & Gemeinholzer [11], and Faberia, as well as Nabalus and Syncalathium, the last two having been recognised as members of subtribe Crepidinae s.str. The exclusion from subtribe Lactucinae of all elements of the polyphyletic genus Nabalus, which is represented in China, depending on the species concept, by four species (under *Prenanthes*) [12] or only one (plus one additionally included species) [13] and of all but one species of Syncalathium has been corroborated recently by Zhang & al. [2,14].

Subtribe Lactucinae in its Chinese Centre of Diversity

The subtribe has two centres of current diversity, one in the Mediterranean-SW Asian region, the other in China and the adjacent Himalayan region. The diversity of the subtribe in its Mediterranean-SW Asian centre came into the focus of systematic research around the middle of the 19th century and led to the description of many new species and two new genera, Cephalor-thynchus Boiss. in 1844 and Steptorhamphus Bunge in 1852. A first comprehensive treatment of the members of the Lactucinae in this centre was provided by Boissier [15] (p. 795ff, as parts of subtribe "Crepideae"). Noteworthy among the more recent publications is in particular the taxonomic revision of the Lactuca alliance in the Iranian Highlands and neighbouring regions by Tuisl [16].

In contrast, the actual extent of the subtribe's diversity in its Sino-Himalayan centre remained unveiled much longer, apart from the Himalayan portion, which was covered rather early by Clarke [17] and Hooker [18], with the most recent updates by Mamgain & Rao [19] and Grierson & Long [20]. Although many species of the subtribe in the large territory of China were discovered and described already in the late 19th and early 20th century, and some of them were included in the revision of Cicerbita sensu lato by Beauverd [21], the subtribe in China became subject of comprehensive studies only towards the end of the 20th century. Pioneer works were done almost exclusively by Shih [12,22–25], who described the new genera Chaetoseris C. Shih, Faberiopsis C. Shih & Y. L. Chen, Notoseris C. Shih, Paraprenanthes C. C. Chang ex C. Shih, *Pterocypsela* C. Shih, *Stenoseris* C. Shih to accommodate the diversity of the subtribe encountered. Shih subsequently also provided the first comprehensive floristic treatment of the entire tribe Cichorieae in China [12]. The subtribal classification applied by Shih largely conforms to that of Stebbins [6], but with corrected subtribal nomenclature. Shih's [12] Lactucinae (corresponding to the Crepidinae s.l. of Stebbins 1953) span the four subtribes Crepidinae s.str., Hieraciinae, Lactucinae s.str. and Hyoseridinae as recognised in the current classification by Kilian & al. [1].

Recently, a reappraisal of the systematics of the *Lactuca* alliance in China, supported by our then still initial nrITS phylogeny of the subtribe including representatives of most Chinese groups, was provided in the frame of the English "Flora of China" [13]. The most striking difference to the treatment by Shih [12] concerns the generic classification: whereas the species of the Lactucinae sensu

Kilian & al. [1] were classified by Shih [12] in altogether 12 genera (Cephalorrhynchus, Chaetoseris, Cicerbita, Lactuca, Lagedium, Mulgedium, Notoseris, Paraprenanthes, Prenanthes, Pterocypsela, Scariola, Stenoseris), they were placed in only five genera (Cicerbita, Lactuca, Melanoseris, Notoseris and Paraprenanthes) by Shih & Kilian [13]. Such different generic classification of the Lactuca alliance is symptomatic for the entire history of the systematics of this alliance. No stability in generic classification has been reached over more than 200 years, because morphological features fail to provide unanimous support for any classification proposed.

Hitherto many Chinese Lactucinae species were only known from herbarium material but never studied in the wild. The first author of the present paper, in contrast, has succeeded to study, collect and sample most Chinese species of the subtribe in the wild, in addition to herbarium studies. Consequently, our initial, sparse molecular sampling of Chinese taxa for nrITS available during the preparation of the "Flora of China" account, now has grown to include the vast majority of the species of the *Lactuca* alliance in China and the nuclear dataset has been complemented by a chloroplast dataset.

The aims of the present paper are (1) to provide the first molecular phylogeny of the Lactucinae which, although focusing on the Chinese centre of diversity, spans the entire subtribe; (2) to detect potential events of reticulation in the evolution of the subtribe by comparing corresponding nuclear and plastid datasets; (3) to test the robustness of the different taxonomies of the Chinese Lactuca alliance in the light of evolution as inferred from the nuclear and plastid trees; (4) to improve the taxonomy of the Chinese Lactucinae based on the molecular phylogenetic reconstruction and morphological studies of living plants and herbarium material including types.

Materials and Methods

Plant Material

The authors have studied herbarium material from the herbaria A, B, CAS, CDBI, E, G, GH, K, KUN, MO, NY, PE and SZ (herbarium codes following Thiers [26]) as well as from the personal herbaria of Ralf Hand (Berlin, Germany), Georg & Sabine Miehe (Marburg, Germany) and Michael Ristow (Potsdam, Germany), of almost all species of the subtribe known from China and adjacent areas, including the types, and the first author extensively studied and collected most Chinese species also in the wild (collection deposited at KUN with some duplicates at B). Besides the permissions for the nature reserves in the Chinese provinces of Chongqing, Sichuan, Xizang and Yunnan by the corresponding Provincial Forestry Departments, no specific permissions were required for material collection; the locations are not privately-owned and none of the species collected in the field are endangered or protected.

Sampling Strategy

Our sampling for the molecular analyses aimed at a dense representation of the subtribe Lactucinae in China. This has been achieved largely so, with the only exception of a few species in North China with Central Asian relation, of which no material could be gathered for this study but which will be included in our global phylogeny of the subtribe (unpublished data). Sequences of one nuclear and five plastid markers were obtained for a total of 130 samples of 78 species. Except for 9, all of the 767 individual marker sequences involved were newly generated for this study. Among the 126 ingroup samples, there are 119 samples of Lactucinae species of China, representing 66 species and 76.7% of the total 86 species recognised by Shih [12], or 55 species and

77.5% of the total species recognised by Shih & Kilian [13], respectively. The corresponding information on the material, including the vouchers preserved, is listed in Appendix S1. Our taxon sampling includes the species providing the types of all generic names established in the subtribe that are relevant to the Lactucinae in China as based on our global phylogeny of the subtribe (unpublished data); these are the types of Cephalorrhynchus Boiss. (C. glandulosus Boiss. \equiv C. hispidus (DC.) Boiss.), Chaetoseris C. Shih (C. lyriformis C. Shih), Cicerbita Wallr. (C. alpina (L.) Wallr.), Faberia Hemsl. (F. sinensis Hemsl.), Lactucella Nazarova (L. undulata (Ledeb.) Nazarova), Lagedium Soják (L. sibiricum (L.) Soják), Melanoseris Decne. (M. lessertiana (DC.) Decne.), Mulgedium Cass. (M. runcinatum Cass. = M. tataricum (L.) DC.), Notoseris C. Shih (N. psilolepis C. Shih), Paraprenanthes C. C. Chang ex C. Shih (P. sororia (Miq.) C. Shih), Prenanthes L. (P. purpurea L.), Parasyncalathium J. W. Zhang & al. (P. souliei (Franch.) J. W. Zhang & al.), Pterocypsela C. Shih (P. indica (L.) C. Shih), Stenoseris C. Shih (S. graciliflora (Wall. ex DC.) C. Shih), Scariola F. W. Schmidt (S. viminea (L.) F. W. Schmidt) and Steptorhamphus Bunge (S. tuberosus (Jacq.) Grossh.) [27]. For many species several individuals were sampled to cover the morphological variation observed, and, wherever possible, samples were gathered from, or as close as possible to, the type locality.

As outgroup, we selected four taxa of the subtribes Crepidinae (*Crepis* and *Soroseris*), Hyoseridinae (*Launaea*) and Hypochaeridinae (*Leontodon*), which represent the decreasingly related other subtribes of the same core group of the Cichorieae according to the molecular analyses by Kilian & al. [1] and Tremetsberger & al. [3]. *Launaea sarmentosa* (subtribe Hyoseridinae) was used to root the trees.

DNA Isolation, Amplification and Sequencing

Genomic DNA was extracted from c. 20 mg of silica-dried leaf tissue or recently collected specimens, either using a modified CTAB methods [28], or the DNeasy kit (Qiagen GmbH, Germany) or Plant Kit Rev. 03 (Macherey-Nagel GmbH & Co. KG, Germany), following the manufacturer's protocols. The DNA amplifications were performed using T1 or T3 Thermocyclers (Biometra, Göttingen, Germany). The amplification reactions with a total volume of 25 µl were of one of the following two compositions: (A) 2 µl DNA template with a concentration of c.15 ng, 1 μ l of each primer (5 pm/ μ l), 1.5 μ l Mg²⁺ (13.9 pm/ μ l), 2.5 µl dNTP mix (2 pm/µl), 2.5 µl×10 Taq reaction Buffer (Chenlü, Kunming, China), 1 µl BSA (bovine serum albumin, 10 ng/μl), 0.3 μl Taq DNA polymerase (2.5 U/μl) (Chenlü, Kunming, China), H₂O; (B) 1 µl DNA template of 20 ng/µl, 1 μl of each primer (10 pm/μl), 1.5 μl MgCl₂ (1.25 mM), 2.5 μl dNTP mix (1.25 pm/μl), 2.5 μl 10x peqLab Taq. Buffer S, 2.5 μl Betain (1.25 mM) [or: 1.5 µl BSA (1.25 mM)], 0.15 µl peqLab HOT Taq. Polymerase (5 units/ μ l), H₂O.

One nuclear and five non-coding chloroplast regions were used as markers. The nuclear ribosomal Internal Transcribed Spacer (nrITS) region (ITS1, 5.8S rDNA, ITS2) was amplified using either the primer combinations ITS4/ITS5 [29] or ITSA/ITSB [30]. Amplification conditions were as follows: an initial denaturation step at 95°C for 3 min, followed by 29 cycles of denaturation at 95°C for 30 s, annealing at 53°C for 30 s, and extension at 72°C for 45 s, then a final extension step at 72°C for 8 min.

The chloroplast markers were amplified using the following primers: (1) the *petD* intron and *petB-petD* spacer were co-amplified with the universal primers PIpetB1411F/PIpetD738R [31]; (2) the *psbA-tmH* spacer with the universal primers psbAF/trnHR [32]; (3) the 5' trnL(UAA)-trnF spacer with the universal primers trnC/trnF [33]; (4) the *rpl32-trnL(UAG)* spacer with the primers rpl32-F/

trnL(UAG) [34] and (5) the *tmQ(UUG)-5'rps16* spacer with the primers trnQ(UUG)/rps16x1 [34]. The PCR amplification conditions were identical for all five chloroplast markers: an initial denaturation step at 80°C for 5 min, followed by 29 cycles consisting of denaturation at 94°C for 45 s, annealing at 52°C for 45 s, extension at 65°C for 50 s, and a final extension step at 65°C for 7 min.

Amplification products and negative controls were visualised in a 1 or 1.2% NEEO agarose electrophorese gel and purified for sequencing using the QIAquick PCR purification Kit (BioTeke Corporation, Beijing, China or Qiagen GmbH, Germany) following the manufacturer's instructions. The concentrations of the purified PCR products were measured with a NanoDrop spectrophotometer (ND-1000, PeqLab, Erlangen, Germany). The purified products were directly sequenced on an ABI 3730XL automated DNA sequencer (Applied Biosystems, Foster City, California, USA) or sequenced via StarSeq (Mainz, Germany) with the same primers as used for amplification.

Sequence Alignment and Coding of Length Mutational Events

The boundaries of the nrITS region (ITS1, 5.8S rDNA, ITS2) and the *petD* marker (*petD* intron and *petB-petD* spacer) were defined according to Goertzen & al. [35] and Borsch & al. [36], respectively. The boundaries of the other markers were taken as indicated in the complete chloroplast genome sequence of *Lactuca sativa* (EMBL/Genbank/DDBJ DQ383816) by Timme & al. [37].

The ITS sequences were aligned manually in PhyDE version 0.9971 [38], according to the Cichorieae part of the Asteraceae alignment by Goertzen & al. [35], which was based on their secondary structure analyses. The plastid sequences were first automatically aligned using Muscle [39], then adjusted manually to a motif-based alignment in PhyDE [38] following the criteria outlined by Kelchner [40], Borsch & al. [41] and Löhne & Borsch [31]. Regions of uncertain homology were excluded from the analysis and inversions were re-inverted (as documented in Appendix S2) prior to the phylogenetic reconstruction.

Indels (as documented in Appendix S3) were coded as informative characters according to the Simple Indel Coding (SIC) method [42] as implemented in the program SeqState version 1.40 [43]. SIC performs about as good as the Modified Complex Indel Coding (MCIC) [44] but has the advantage that the SIC matrix can also be easily analysed with Bayesian Inference.

Additive polymorphic sites (APS) in the nrITS sequences, indicating potential introgressive hybridisation, were detected following the criteria outlined by Fuertes Aguilar & Nieto Feliner & al. [45].

Phylogenetic Reconstruction

Incongruence Length Difference (ILD) test [46] implemented in PAUP* version 4.0b10 [47] as the Partition Homogeneity Test, was performed to assess the congruence between the nuclear and plastid data sets. For this test, which calculates the ILD first for the original partitions and then for a series of randomized partitions of the same size, the following parameters were used: heuristic search of 10 000 replicates, each with 100 random addition searches, maxtrees set to 1 and one tree held each step. As significance threshold for congruence or homogeneity of the partitions a P value of >0.01 is considered as appropriate [48].

Phylogenetic relationships were reconstructed using Maximum Parsimony (MP) and Bayesian Inference (BI). Maximum Parsimony analyses were performed using the Parsimony Ratchet [49] with PRAP [50] in combination with PAUP* version 4.0b10 [47].

Standard ratchet settings were used: 200 ratchet iterations with 25% of the positions randomly upweighted (weight = 2) during each replicate and 10 random addition cycles. The generated command files also including the nexus data matrix were run in PAUP* version 4.0b10 [47] using heuristic search with the following parameters: all characters have equal weight, gaps are treated as 'missing', simple addition of sequences, TBR branching swapping, maxtrees setting to 100 and auto-increased by 100, one non-binary starting tree arbitrarily dichotomized before branch swapping, only one tree saved. A majority rule consensus tree was calculated from the most parsimonious trees received. Jackknife (IK) support values for the nodes found by the MP analysis were calculated in PAUP*version4.0b10 applying the optimal jackknife parameters according to Farris & al. [51] and Müller [52]: 10 000 jackknife replicates were performed using the TBR branch swapping algorithm with 36.788% of characters deleted and one tree held during each replicate.

Bayesian Inference analyses were performed using MrBayes 3.2 [53]. Optimal nucleotide substitutions models were searched separately for each of the three partitions of the nrITS dataset (i.e. ITS1, 5.8S, ITS2) and each of the five plastid markers with MrModeltest 2.3 [54], following the Akaike Information Criterion (AIC). The optimal model chosen for ITS1 and ITS 2 was GTR+I+G, for 5.8S SYM+I, for the petD region GTR+I, and for the other four plastid markers GTR+G. A binary (restriction site) model was implemented for the coded indels. The datasets were partitioned in MrBayes 3.2 into three (nuclear) or five (plastid) DNA markers, respectively, and one partition for the coded indels. All analyses in MrBayes 3.2 were performed with four simultaneous runs of Metropolis-coupled Markov Chains Monte Carlo (MCMCMC), each with four parallel Markov chains. Each chain was performed for 2 million generations and, starting with a random tree, one tree was saved every 100th generation. For other parameters the default settings of the program were left unchanged. A conservative burn-in of 0.2 (i.e. discarding the first 20% of the trees) was applied after graphically checking chain convergence using the program AWTY [55]. The remaining trees were used to generate a majority rule consensus tree.

TreeGraph 2 [56] was used to assess the tree topologies and to visualise the trees with node supports.

Results

Molecular Datasets and Phylogenetic Analyses

Nuclear ribosomal ITS region. The ITS region varied from 592 to 644 nt in our 130 (126 ingroup +4 outgroup) samples. Of a total of 667 characters in the aligned data set, 261 were parsimony informative. Simple Indel Coding increased the total number of characters to 734 and the number of parsimony informative characters to 301. With 39.1% (41.0% including coded indels) parsimony informative sites it has the highest phylogenetic performance of all markers used, but has the lowest consistency index and retention index of all individual marker trees (Table 1).

The Maximum Parsimony (MP) search resulted in 70 most parsimonious trees (L = 1204, CI = 0.485, RI = 0.840, RC = 0.408, see Table 1). The 50% majority rule MP consensus tree was essentially congruent in topology with the Bayesian Inference (BI) 50% majority rule consensus tree, apart from an incongruence in one subclade of the *Lactuca* lineage, where in the BI tree the *L. sativa-L. serriola* clade is sister to the *Scariola* and *Lagedium-Mulgedium* clades, while in the MP tree the *Lagedium-Mulgedium* clade is sister to the other two. We give here only the BI phylogram (Fig. 1), with

the MP Jackknife support (JK) values above and the BI posterior probability (PP) values below the branches.

Non-coding chloroplast regions. The plastid matrix was of the same sample size and composition as the ITS region matrix. The length of the individual plastid markers ranged from 171 (with a unique large deletion in *Chaetoseris macrantha*) to 421 nt in *psbA-tmH*, to 929–998 nt in *tmQ*^(UUG)-5' *rps16*. The length of the five combined plastid markers ranged from 3784–4028 nt. The full data are provided in Table 1.

Areas with uncertain homology classified as "hotspots" of sequence mutation according to Borsch & al. [41], mostly length-variable poly A/T-stretches, were excluded from the analyses. One exon (petD) and one hotspot were excluded from the petD region, five hotspots from psbA-tmH, one exon (tmL) and one hotspot from 5'tmL(^{UAA)}-tmF, eight hotspots from pl32-tmL(^{UAG)} and three hotspots from tmQ(^{UUG)}-5'rps16 (see Appendix S2). The length of the five combined plastid markers after exclusion of the hotspots ranged from 3619 to 3884 nt (see Table 1).

The final matrix of the *rpl32-tmL*^(UAG) region comprised 154 parsimony informative characters without and 191 parsimony informative characters including the coded indels, having the highest phylogenetic performance among the five cp markers used (Table 1). It is followed by the *tmQ*^(UUG)-5'rps16 region, with 111 and including coded indels 129 parsimony informative characters. The smaller *psbA-tmH* region has a percentage of informative sites comparable to the *rpl32-tmL*^(UAG) region, but excessive variation (even within species) rendered the alignment and homology confirmation partly difficult. 5'tmL^(UAA)-tmF had the lowest phylogenetic performance with 56 and including coded indels 66 parsimony informative characters. The final concatenated plastid matrix comprised 450 and including coded indels 545 parsimony informative characters.

MP analyses were performed for both the individual cp markers and the concatenated plastid data set. The tree statistics are given in Table 1. MP analysis of the concatenated matrix resulted in 48 most parsimonious trees with L = 1342, CI = 0.847, RI = 0.950, RC = 0.805 (see Table 1). The resulting 50% MP consensus tree is congruent with the corresponding BI tree, apart from (a) two cases where smaller crown clades recognised in the MP tree collapsed in the BI tree, and (b) an incongruence in the relationship within the outgroup, where Faberia clustered in the BI tree with the two members of subtribe Crepidinae (i.e. Crepis and Soroseris), following Leontodon (Hypochaeridinae) as the nearest sister to the subtribe Lactucinae (incl. P. purpurea), while in the MP tree Faberia clustered only with Soroseris, in the closest position to Lactucinae, followed by Leontodon and Crepis as the successive sisters. We give here the BI phylogram (Fig. 2), with the MP Jackknife support (JK) values above and the BI posterior probability (PP) values below the branches.

Phylogenetic Relationships

ITS phylogeny. Maximum Parsimony (MP) and Bayesian Inference (BI) analyses based on the ITS matrix both depict seven major lineages within a well supported (JK = 97.5, PP = 0.99) subtribe Lactucinae, which all have high statistical support, whereas deeper node have low or lack statistical support. Clade 1 (JK = 99.9, PP = 1) comprises the genus Faberia in the circumscription of Shih & Kilian [13], thus including Faberiopsis and Prenanthes faberi; this Faberia clade is sister (JK = 56, PP = 1) to the remainder of the subtribe. Clade 2 (JK = 84, PP = 1) is restricted to and unites the non-Chinese species providing the types of Cephalorrhynchus and Cicerbita; it is sister to a large clade A (JK < 50, PP = 0.95) including clades 3–7 of the subtribe. The large clade A in turn comprises the clades 3 and B. Clade 3 (JK = 94.6,

Table 1. Sequence and tree statistics of the six individual markers and the concatenated plastid matrix.

Data matrix	Length range total/ HS ¹ excluded (nt)	No. total char. ² / No. total char. ³ (nt)	No. inform. sites ⁴ / No. inform. sites ³ (nt)	No. MPTS ³	TL ³	CI ³	RI ³	RC ³
ITS region	592–644	667/734	261(39.1%)/301(41.0%)	70	1204	0.485	0.840	0.408
petD region	887-922/876-906	928/948	67(7.2%)/80(8.4%)	22	171	0.877	0.972	0.853
psbA-trnH	171-421/131-382	464/497	62(13.4%)/79(15.9%)	1474	182	0.824	0.926	0.763
trnL-F	786-841/737-792	825/857	56(6.8%)/66(7.7%)	28	178	0.899	0.958	0.861
trnQ-rps16	929-998/928-997	1174/1221	111(9.5%)/129(10.6%)	131	339	0.861	0.951	0.819
rpl32-trnL	830-939/807-894	1139/1223	154(13.5%)/191(15.6%)	19	456	0.844	0.953	0.804
combined cpDNA	3784-4028/3619-3884	4530/4746	450(9.9%)/545(11.5%)	48	1342	0.847	0.950	0.805

¹hotspots (and exons), see Table S1:

²number of total character;

³with indel codina:

⁴number of informative sites.

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PP=1) included three species placed by Shih & Kilian [13] in Cicerbita. Clade B forms with low support (JK<50, PP = 0.62) the polytomous backbone of the Lactucinae, including clades 4-7. **Clade 4** (JK = 94.2, PP = 1) represents *Lactuca*, the type genus of the subtribe, among which the former Lactuca segregates Pterocypsela, Steptorhamphus, Mulgedium and Lactucella are nested. **Clade 5** (IK = 89.1, PP = 1) includes the types and most other species of the genera *Chaetoseris* and *Stenoseris*, among which the species providing the type of the old generic name Melanoseris is nested, and also the recently erected genus Parasyncalathium. Clade **6** (JK = 98.9, PP = 1) comprises the Chinese near-endemic genus Notoseris, but excluding two of its members in the sense of Shih [12] or Shih & Kilian [13], which cluster in clade 7 with Paraprenanthes. **Clade 7** (JK = 91.2, PP = 1), finally, includes the Chinese endemic genus Paraprenanthes plus a few more species not considered by Shih [12] and Shih & Kilian [13] as members of that genus.

Plastid phylogeny. MP and BI analyses of the combined plastid data set fully agree with respect to the phylogenetic relationships between and in the circumscription of the major lineages. They yielded six major lineages with mostly high statistical support, which are not all identical with those in the ITS phylogeny. Their relationships to each other also received high statistical support. The Faberia clade (JK = 100, PP = 1) is identical in circumscription to the corresponding clade in the ITS phylogeny, but here placed among the outgroup taxa clustered with Soroseris (JK = 99.9, PP = 1) and then Crepis (JK < 50, PP = 0.81), which are members of the subtribe Crepidinae. The Lactucinae ingroup has high support (JK = 98.9, PP = 1), comprising clades 1-6. **Clade 1**, only comprising *Prenanthes purpurea*, is sister (IK = 100, PP = 1) to the remainder of the ingroup. **Clade 2** (JK = 99.9, PP = 1) is congruent to clade 2 of the ITS tree and is sister to a large **clade A** (IK = 99.8, PP = 1), which includes the remaining major lineages 3-6. Clades 3-5 are parts of a clade B (JK = 99.9, PP = 1), to which clade 6 is sister. **Clade 3** (JK = 100, PP = 1) is congruent to clade 3 of the ITS tree and sister to **clade** \mathbf{C} (JK = 99.9, PP = 1), which comprises clades 4 and 5. Clade 4 (JK = 68.8, PP = 0.98), comprising *Lactuca*, is congruent in circumscription but less so in internal topology with clade 4 of the ITS tree. Clade 5 (JK = 81.7, PP = 1) is congruent in circumscription to clade 5, including Melanoseris, Chaetoseris, Parasyncalathium and Stenoseris, of the ITS tree, but has a somewhat different internal topology. Clade 6 (JK = 100, PP = 1) finally, is congruent in circumscription to clades 6 Notoseris plus 7

Paraprenanthes, of the ITS tree, merging the taxa of these two clades in a different internal topology.

Incongruences between Nuclear and Plastid Phylogenies

The ILD test detected incongruence with high significance (P=0.001) between the entire nuclear and plastid data sets as well as, in the calculation for the single clades, for the *Notoseris*, *Paraprenanthes* and *Melanoseris* clades. Therefore no analyses of a combined data set have been performed. While the ILD test is known to be overly sensitive in indicating conflicts between datasets [47], and alone therefore no sufficient proof for incongruence, its result in our case is fully corroborated by the high statistical branch support for the incongruent tree topologies (see MP Jackknife support values above and the BI posterior probability values below the branches in Fig. 1 and 2).

Incongruences between the two phylogenies with good to high branch support concern (1) the circumscription of the Lactucinae, (2) the relationships between major lineages, (3) assignment of taxa to major lineages, and (4) the internal topology of major lineages.

- (1) The circumscription of the subtribe Lactucinae is incongruent between the ITS and plastid trees: (a) the Faberia clade is sister to the remainder of the subtribe in the ITS tree (JK = 56, PP = 1, and JK = 97.5, PP = 0.99 for the Lactucinae including Faberia) but nested within the outgroup in the plastid tree (JK = 99.1, PP = 1 for the sister group relationship with Soroseris); (b) Prenanthes purpurea is nested in the outgroup in the ITS tree (JK = 97.5, PP = 0.99 for the ingroup without P. purpurea) but forms the first diverging branch of the Lactucinae in the plastid tree (JK = 98.9, PP = 1 for the sister group relationship with the core Lactucinae). Disregarding these two lineages, the Lactucinae are monophyletic in both phylogenies.
- (2) The most obvious incongruence in the relationships between the major lineages is that the *Notoseris* lineage (clade 6) and the *Paraprenanthes* lineage (clade 7) of the ITS phylogeny (JK = 98.9, PP = 1 for clade 6 and JK = 91.2, PP = 1 for clade 7) are represented in the plastid phylogeny by a single clade 6 (JK = 100, PP = 1) of different internal topology. In contrast, the topological incongruences in the relationships of these lineages to the other major lineages as well as in the relationships among these other major lineages are without statistical support, because the most major lineages are found

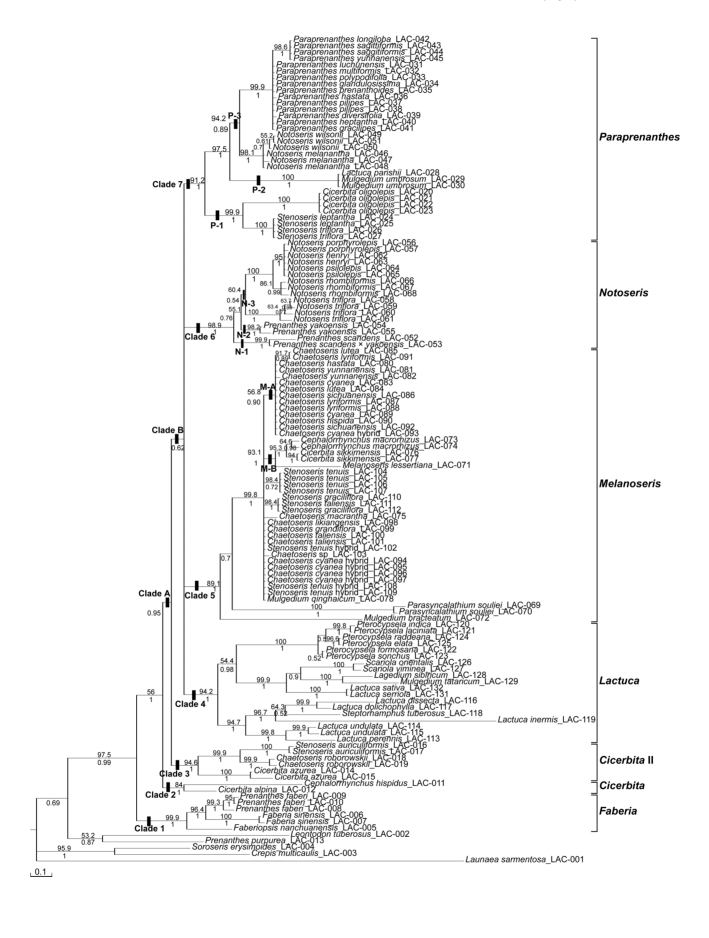


Figure 1. Bayesian phylogram (majority rule tree) of subtribe Lactucinae based on nrITS dataset including coded indels. Posterior probabilities (PP) are given below the branches, the jackknife support values (JK) of the corresponding Maximum Parsimony majority rule consensus tree above the branches. Reference point for the names of Chinese taxa is in general the morphology-based taxonomy of Shih (1997), whereas the clade names at the vertical bars on the right show our revised generic classification outlined in more detail and with the relevant synonymies in the Taxonomic conclusions.

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along the polytomous backbone in the core of the subtribe in the ITS phylogeny.

- (3) In the single case of Mulgedium bracteatum (≡ Melanoseris bracteata), the assignment of a species to the major lineages is incongruent between the ITS and the plastid phylogeny. In our ITS tree focusing on Chinese Lactucinae this species is included in the Melanoseris clade with good statistical support (JK = 89.1, PP = 1) but in a fairly isolated position as the first diverging branch of that clade. In our plastid tree, in contrast, this species is nested in the Lactuca clade with moderate support (JK = 68.8, PP = 0.98), clustering therein with SW to E Asian members of Lactuca in a polytomous clade.
- (4) Incongruences in the internal topology occur (a) in the Notoseris and Paraprenanthes clades (clades 6 and 7 in the ITS phylogeny, parts of the single clade 6 in the plastid phylogeny), (b) in the Lactuca clade (clade 4 in both the ITS and plastid phylogeny), and (c) in the Melanoseris clade (clade 5 in both the ITS and plastid phylogeny). These are addressed in more detail in the Discussion.

Discussion

Our phylogenetic reconstruction of the Lactucinae by molecular techniques is based on the most extensive sampling published for the subtribe to date. Our sampling, although focusing on the Chinese centre of diversity, spans the entire subtribe, including not only all genera present in China but also non-Chinese species providing the types of relevant generic names in the subtribe. We provide the first comprehensive phylogeny of this taxonomically difficult and controversial group and use this together with morphological data as basis for a revised generic classification of its members in the Chinese centre of diversity.

Possible Causes of Incongruence between the Nuclear and Plastid Phylogenies

Technical causes, such as insufficient taxon sampling, long-branch attraction, sequencing errors, for the statistically well supported and thus "hard" topological incongruences [57] between our nuclear and plastid phylogenies, appear excludable in the light of our dense sampling, frequently with more than one sample per species, and the similar topologies obtained from both MP and BI analyses. Causes for these incongruences are judged with confidence therefore as essentially biological.

The nrITS sequences of our dataset appeared reliable (no pseudogenes) but we cannot exclude the possibility of divergent alleles among the multiple ITS copies within a nucleus [58]. In a few cases, additive polymorphism [45] seems in fact present among sequences of closely related taxa (see Table 2) and supports the hypothesis that nuclear introgression has taken place.

The sequences of the exclusively maternally inherited and thus non-recombining chloroplast genome come along with another drawback. This is the relatively high potential for interspecific cytoplasmic (chloroplast) gene flow, or chloroplast capture, also in absence of any nuclear gene flow, due to introgressive hybridisation [59–62] or even due to horizontal gene flow between sexually incompatible species [63]. Chloroplast capture is known from the

Cichorieae even at intergeneric level [64] and is with or without incomplete lineage sorting [65–66] an important cause for incongruence between nuclear and plastid phylogenies in general.

Putative cases of ancient reticulation and chloroplast **capture.** The *Notoseris and Paraprenanthes* lineages, which form the well supported clades 6 and 7 along the polytomous backbone of the larger part of the subtribe in our nuclear phylogeny (Fig. 1), in contrast form the single joined clade 6 in the plastid phylogeny (Fig. 2). Notably, the first basally diverging branches of both lineages in the nuclear tree (N-1+ N-2 and P-1) appear as subclades N-1/2 and P-1 in the basal polytomy of the common clade in the plastid tree, while the core clades of both lineages in the nuclear tree appear as subclades of a second, later diverging polytomy in the plastid tree (compare Fig. 1 and 2). Only a few chromosome counts are known from species of the core clades of the two lineages, all indicating them to be diploids with 2n = 18[67]. The only plausible explanation for this incongruence appears to us the assumption of an event of intergeneric reticulation with chloroplast capture already between ancestors of the current lineages. Early divergence of the basally branching subclades, along with geographical isolation and ecological separation through flowering time, may have led them accumulate sufficient chloroplast gene variation to be well distinguished from the remainder. The inner polytomous topology of both core clades of Paraprenanthes and Notoseris sensu Shih [12], in combination with their morphological homogeneity in each clade, may probably be ascribed to recent rapid radiation in a similar distributional area and ecological niche. The *Paraprenanthes umbrosa* subclade (P-2, represented in the trees by Lactuca parishii and Mulgedium umbrosum), is sister to the core Paraprenanthes clade P-3 in the ITS tree, but sister to the polytomous mixed Paraprenanthes-Notoseris core clade (including N-3 and P-3a+P-3b with different internal topology) in the plastid tree. This topology makes it likely that between the ancestors of the two core clades N-3 and P-3 further events of reticulation and cytoplasmic introgression may have taken place. With respect to the generic classification, we consider the nuclear phylogeny, which places Notoseris and Paraprenanthes in separate lineages, a better estimate of the taxon phylogeny because it is more in line with morphology.

A second putative case of ancient reticulation and chloroplast capture is exemplified by the entire genus *Faberia*, which appears in different subtribal placements in both trees (see under *Faberia* lineage., below). *Faberia* is alloploid with 2n = 34 [68–69], cytoplasmic gene flow was thus evidently accompanied by nuclear gene flow.

A third putative case of ancient reticulation constitutes the diploid *Prenanthes purpurea*. From morphological and cytological evidence it appears in this case very unlikely that the ITS tree represents the actual species phylogeny, whereas much more so that the plastid tree does (see under *Prenanthes purpurea* lineage, below).

Putative cases of introgressive hybridisation between extant species. A rather clear example for incongruence indicating reticulation and cytoplasmic gene flow among extant species concerns the scandent species Notoseris scandens and N. yakoensis (see Fig. 3B–C; as Prenanthes scandens and P. yakoensis in the trees) in the Notoseris clade of the ITS tree and the joint Notoseris

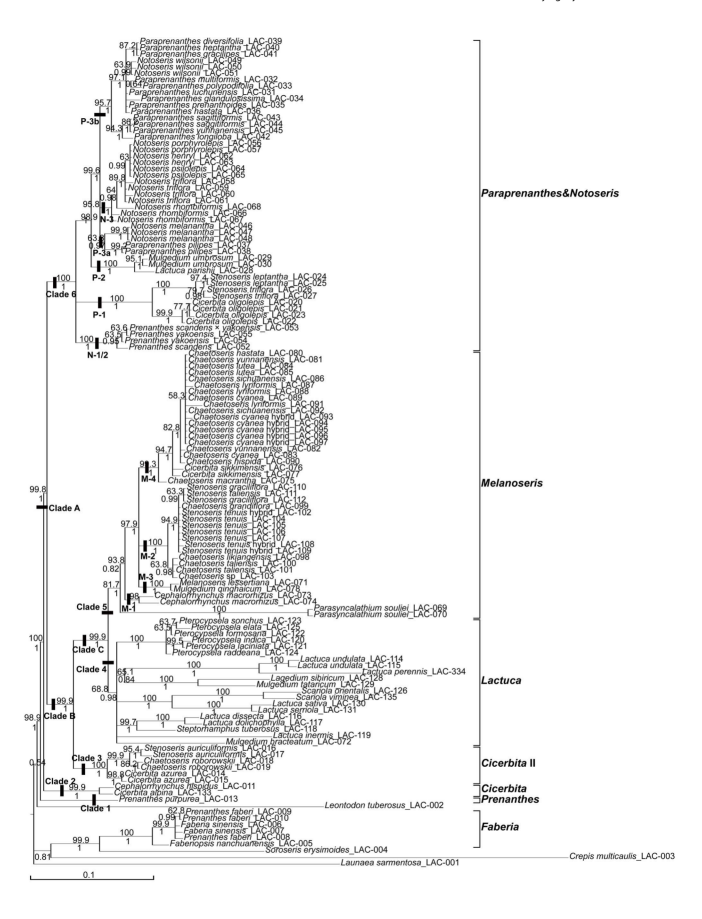


Figure 2. Bayesian phylogram (majority rule tree) of subtribe Lactucinae based on plastid dataset with coded indels. Posterior probabilities (PP) are given below the branches, the jackknife support values (JK) of the corresponding Maximum Parsimony majority rule consensus tree above the branches. Reference point for the names of Chinese taxa is in general the morphology-based taxonomy of Shih (1997), whereas the clade names at the vertical bars on the right show our revised generic classification outlined in more detail and with the relevant synonymies in the Taxonomic conclusions.

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Paraprenanthes clade in the plastid tree. The two species form a clade of their own in the ITS tree and consecutive sister clades in the plastid tree, but the morphologically intermediate accession designated as Prenanthes scandens \times yakoensis clusters with P. scandens in the ITS tree (JK = 99.9, PP = 1), whereas with P. yakoensis in the plastid tree (JK = 63.6, PP = 0.95). Additive polymorphism [45] is present in the ITS sequence of the accession P. scandens \times yakoensis (Table 2) and the putative hybrid population is morphologically clearly intermediate in the number and length of the inner phyllaries, the flower number per capitulum and the anthertube length. Since divergent ITS paralogues merged in a genome after a hybridisation event become homogenised by concerted evolution, the occurrence of a number of additive polymorphic sites (APS) supports a rather recent (as opposed to an ancient)

introgression event. Based on these evidences, we hypothesise the formation of a natural hybrid population between *P. yakoensis* and *P. scandens*, with the former as its male parent and the latter as female parent, involving both plastid and nuclear introgression. The two scandent species typically grow at edges of montane forests to tall forb communities, e.g. along rivers, but found new habitats along roads through montane forests, which eventually helped formerly isolated populations of the two species to meet.

A second case is *Paraprenanthes melanantha* (Fig. 3E; as *Notoseris melanantha* in the trees). This species clusters with the morphologically closely allied *P. wilsonii* (Fig. 3D; $\equiv \mathcal{N}$ wilsonii) with strong support (JK = 98.1, PP = 1) in the ITS tree, but with the widely distributed *P. sororia* (represented in the tree by its glandular hairy form that was treated as *P. pilipes* by Shih [12]), with lesser support

Table 2. Additive Polymorphic Sites (APS*) in the nrITS region sequences in four exemplar cases of putative introgressive reticulation.

Sample name in the tree	Pos	itior	ns of	Add	litive	Polym	orphi	c Site:	s (APS) in th	e nrlT	S reg	ion se	quend	:e		
1. Prenanthes scandens × yakoensis	40	41	50	53	57	73	82	127	129	155	199	202	210	456	525	603	62
Prenanthes scandens_LAC-052	T	Α	C	Т	Α	C	T	Α	G	T	C	T	T	C	T	T	Т
Prenanthes scandens × yakoensis_LAC-053	Υ	W	Υ	Υ	R	Υ	Υ	R	R	Υ	М	Υ	K	Υ	Υ	Υ	Υ
Prenanthes yakoensis_LAC-054	C	Т	Т	C	G	Т	C	Α	Α	C	Α	C	G	Т	C	Т	C
Prenanthes yakoensis_LAC-055	C	Т	Т	C	G	T	C	R	Α	C	Α	C	G	T	C	C	C
2. Stenoseris tenuis hybrid	26	53	64	86	125	199	202	231	236	443	446	450	528	534	576	579	62
Stenoseris tenuis_LAC-104	C	Т	G	C	Т	Т	Т	T	Т	C	Т	Α	Т	C	C	C	Т
Stenoseris tenuis_LAC-105	C	Т	G	C	Т	Т	Т	Т	Т	C	Т	Α	Т	C	C	C	Т
Stenoseris tenuis hybrid_LAC-108	Υ	Υ	R	Υ	Υ	Υ	Υ	Υ	Υ	М	Υ	R	Υ	Υ	Υ	Υ	Υ
Stenoseris tenuis hybrid_LAC-109	Υ	Υ	R	Υ	Υ	Υ	Υ	Υ	Υ	М	Υ	R	Υ	Υ	Υ	Υ	Υ
Chaetoseris cyanea_LAC-083	Т	C	Α	Т	C	C	C	C	C	Α	C	G	C	Т	T	Т	C
Chaetoseris lyriformis_LAC-088	Т	C	Α	Т	C	C	C	C	C	Α	C	G	C	Т	Т	Т	C
3. Chaetoseris cyanea hybrid	14	26	64	86	123	443	446	450	554	565	579						
Chaetoseris taliensis_LAC-100	G	C	G	C	Α	C	Т	Α	C	Α	C						
Chaetoseris taliensis_LAC-101	G	C	G	C	Α	C	T	Α	C	Α	C						
Chaetoseris cyanea hybrid_LAC-094	K	Υ	R	Υ	М	М	Υ	R	Υ	М	Υ						
Chaetoseris cyanea hybrid_LAC-095	K	Υ	R	Υ	М	М	Υ	R	Υ	М	Υ						
Chaetoseris cyanea hybrid_LAC-096	K	Υ	R	Υ	М	М	Υ	R	Υ	М	Υ						
Chaetoseris cyanea hybrid_LAC-097	K	Υ	R	Υ	М	М	Υ	R	Υ	М	Υ						
Chaetoseris cyanea_LAC-083	Т	Т	Α	Т	C	Α	C	G	T	C	Т						
Chaetoseris lyriformis_LAC-088	Т	Т	Α	Т	C	Α	C	G	T	C	T						
4. Notoseris melanantha	26	53	55	82	120	164	195	200	462	596							
Paraprenanthes pilipes_LAC-037	C	C	Т	C	G	G	G	G	Т	G							
Paraprenanthes pilipes_LAC-038	C	C	Т	C	G	G	G	G	Т	G							
Notoseris melanantha_LAC-046	Υ	Υ	W	Υ	R	R	R	R	K	R							
Notoseris melanantha_LAC-047	Υ	Υ	W	Υ	R	R	R	R	K	R							
Notoseris wilsonii_LAC-049	Т	C	Α	C	G	G	G	G	Т	G							
Notoseris wilsonii_LAC-051	Т	С	Α	С	G	G	G	G	Т	G							

*An APS is recorded when at least one of the bases involved in a polymorphic site occurs separately at the same position in samples of putative parents. doi:10.1371/journal.pone.0082692.t002



Figure 3. Selected species of Notoseris (A–C) and Paraprenanthes (D–F) in situ. A, Notoseris henryi (Sichuan, 9 Sep. 2013, photo by H. J. Dong; voucher: H. J. Dong & al. 870 (KUN)), B. N. scandens (Yunnan, 11 Nov. 2011, photo by Y. Tang; voucher: Z. H. Wang, L. Chen & Y. Tang 457 (KUN)), C. N. yakoensis (Yunnan, 11 Nov. 2011, photo by Y. Tang; voucher: Z. H. Wang, L. Chen & Y. Tang 458 (KUN)), D. Paraprenanthes wilsonii (Sichuan, 25 Jun. 2011, photo by Z. H. Wang; voucher: Z. H. Wang & L. Chen 344 (B, KUN)), E, P. melanantha (Sichuan, 2 Aug. 2011, photo by Z. H. Wang; voucher: Z. H. Wang & L. Chen 489 (B, KUN)), F, P. oligolepis (Yunnan, 22 Sep. 2011, photo by G. X. Hu; voucher: H. J. Dong & al. 416 (KUN)). doi:10.1371/journal.pone.0082692.q003

[JK = 63.8, PP = 0.97) in our plastid tree. Additive polymorphism [45] in the ITS sequence of *P. melanantha* compared to *P. wilsonii* and *P. sororia* (Table 2) supports that cytoplasmic gene flow was accompanied in this case also by nuclear introgression. This could indicate that *P. melanantha* is hybridogenous with *P. wilsonii* as paternal and *P. sororia* as maternal parent. However, besides clearly additive polymorphic sites, we notice also polymorphic sites in *P. melanantha* that are not additive with respect to *P. sororia* and *P. wilsonii*. Moreover, and in contrast to the preceding case, morphologically, *P. melanantha* is not intermediate between the putative parental species but close to the paternal one, as both share an involucre with only 5 inner phyllaries (8 in *P. sororia*) and anther tubes without appendages >3 mm (not exceeding 1.6 mm in *P. sororia*). Presumably, in this case a more complex pattern of

reticulation might have taken place and further studies are necessary to shed some light on it.

Other cases are addressed under the *Lactuca (Pterocypsela sonchus* and *P. elata)* and *Melanoseris (M. bracteata, M. graciliflora* and *M. tenuis, M. cyanea* group) lineages below.

Monophyly and Circumscription of Subtribe Lactucinae

Considering the joint evidence produced by the nuclear and the plastid phylogeny, subtribe Lactucinae is monophyletic only if the *Faberia* and *Prenanthes purpurea* lineages are disregarded. Otherwise its circumscription as a monophyletic entity depends on whether the nuclear or chloroplast phylogeny is followed.

Faberia lineage. The genus Faberia, endemic to SW China, was included in Prenanthes and treated as a member of subtribe Lactucinae by Bremer [8] and Lack [9], but excluded by Kilian & al. [1] from the Lactucinae, in absence of DNA sequence data for morphological grounds only. Later Shih & Kilian [13] included Faberia (merged again with its former segregate Faberiopsis) in subtribe Lactucinae, based on our initial phylogenetic analysis of nrITS sequences. The position of Faberia in the nuclear tree, where it is placed with moderate statistical support (JK = 56, PP = 1) as sister to all other members of subtribe Lactucinae, and were the Lactucinae including Faberia received high support (JK = 97.5, PP = 0.99), is incongruent with its position in our plastid tree, where it is nested in the outgroup, with relative low support (JK<50, PP=0.81) among the Crepidinae and within them as sister to the single *Soroseris* sample included (JK = 100, PP = 1). Liu & al. [68] have shown that Faberia has the chromosome number of 2n = 34, which is unusual in the Cichorieae and indicates an alloploid origin of the genus from parents with x = 8 and x = 9. Its incongruent positions in the nuclear and plastid trees make a reticulation with a maternal ancestor of the genus from the Crepidinae and a paternal ancestor from the Lactucinae the most likely scenario, but a plastid phylogeny with a much more extensive sampling would be necessary to asses its potential maternal ancestor. Whether the nuclear or chloroplast phylogeny provide the better phylogenetic estimate for the genus is difficult to assess, because morphology is little decisive in this case. The assumed sudden and rapid diversification of tribe Cichorieae in its evolutionary history [1,3], might be an explanation that clear synapomorphies are frequently missing for the major lineages recognised as subtribes [8]. This applies especially to subtribes Lactucinae and Crepidinae, and certainly is the major reason for their late recognition as separate lineages. Bremer [8] identified for the Crepidinae an involucre distinctly differentiated between inner and outer phyllary series (typically so in e.g. Youngia and Ixeris) as a possible synapomorphy. In fact, in the Lactucinae often the outer phyllary series grade into the inner ones (e.g. often so in Lactuca and Melanoseris), but Notoseris and Paraprenanthes, e.g., have distinctly separated inner and outer phyllary series, as present also in Faberia. Morphological reasons for both placements of Faberia can be found according to our current knowledge. For classification purposes, we follow, for the time being, the nuclear DNA phylogeny and hence treat Faberia as a member of subtribe Lactucinae.

The revised circumscription of *Faberia* as a genus of seven species endemic to China given by Shih & Kilian [13], with reinclusion of the former segregate *Faberiopsis* and inclusion of *Prenanthes faberi*, is fully corroborated by both our nuclear and plastid phylogenies.

Prenanthes purpurea lineage. Prenanthes purpurea L., a chiefly European species, provides the type of the generic name Prenanthes. Kilian & Gemeinholzer [11] and Kilian & al. [1] stated that this genus should probably be considered as monospecific, because the many other species formerly included seem unrelated to P. purpurea. This hold true also for the seven Chinese species maintained as members of Prenanthes by Shih [12,22]: four of them (P. angustiloba, P. leptantha, P. macrophylla and P. tatarinowii) were found to belong actually to subtribe Crepidinae [2,13–14]; among the three remaining species, one, P. faberi, is nested in our analyses in Faberia, and two, P. scandens and P. yakoensis, in the Notoseris clade of the ITS tree or the Notoseris-Paraprenanthes clade of the plastid tree, respectively (Fig. 1–2).

In the nrITS trees published, *Prenanthes purpurea* is placed far distant from the Lactucinae [70] and clusters instead with the subtribe Hypochaeridinae [1,71] as in our ITS tree. This placement is meanwhile supported by ITS sequences of four

different accessions but is surprising because P. purpurea and the Hypochaeridinae are morphologically entirely unrelated: P. purpurea has cyanic flowers (instead of always yellow or, rarely, white flowers in the Hypochaeridinae), pendent (instead of usually erect) flowering capitula, a pappus of scabrid (instead of almost always stiffly fimbriately plumose) bristles. They also do not agree cytologically. P. purpurea is diploid with x = 9 [72], while the basic chromosome numbers in the Hypochaeridinae range from x = 3 to x = 7 with a single exception of x = 11 [71].

In contrast to the ITS phylogeny, *Prenanthes purpurea* has a basally branching position in our plastid phylogeny, being sister with full support (JK = 100, PP = 1) to all other genera of the Lactucinae except *Faberia*. The same has been indicated, but without statistical support, in a previous *matK* tree [1]. Both from morphology and cytology, *P. purpurea* would in fact best fit into subtribe Lactucinae. Pending further studies to elucidate the causes for the incongruent molecular results, it would be appropriate either to include it, with reservations, in the Lactucinae, or else to leave *Prenanthes* unassigned to a subtribe, instead of placing it into the Hypochaeridinae.

Core lactucinae. Our analyses, which include (a) all major lineages of the subtribe Lactucinae, (b) all species groups present in China, and (c) also the species providing the types of the relevant generic names established in the subtribe, revealed congruently in the nuclear and plastid phylogenies a core of the subtribe comprising six (five in the plastid phylogeny) major lineages, of which five (four) are present in its Chinese centre of diversity (Fig. 1–2): (1) the *Cicerbita* lineage, (2) the *Cicerbita* II lineage, (3) the *Lactuca* lineage, (4) the *Melanoseris* lineage, (5) the *Notoseris* lineage, and (6) the *Paraprenanthes* lineage, the last two revealed as a single clade in the plastid phylogeny.

Relationships of the major lineages within the core Lactucinae can be inferred from our analyses with some caution only, because of the lacking resolution for the deeper nodes in the ITS tree. Good support, however, is received for the sister group relationship of the Cicerbita lineage to the remainder of the core Lactucinae in both phylogenies (JK \leq 50, PP = 0.95 in the ITS tree; JK = 99.8, PP = 1 in the plastid tree, see Fig. 1–2). The relationship of the Cicerbita II lineage is incongruent in both datasets: in the ITS phylogeny it is sister with low support in the BI tree (JK <50, PP=0.62, relationship unresolved in the MP tree) to the unresolved remainder of the core Lactucinae, whereas in the plastid phylogeny it is sister to a clade comprising the Lactuca and Melanoseris lineages with almost full support (JK = 99.9, PP = 1). Considering the weak support through the ITS dataset, this incongruence should be regarded as soft and rather the sister group relationship of the Cicerbita II lineage to the Lactuca + Melanoseris lineages favoured as hypothesis. Inferred from the plastid tree, the Notoseris and Paraprenanthes lineages may be regarded as sisters, which are in turn sister to the Cicerbita II + Lactuca + Melanoseris lineages.

Hence, the following hypothesis on the evolution of the subtribe Lactucinae may be proposed: the ancestors of the mesic European-SW Asian *Cicerbita* lineage have, on the one hand, migrated towards eastern Asia giving rise to the mesic *Notoseris* and *Paraprenanthes* lineages and, on the other hand, migrated north- and northeastwards across Eurasia to North America as well as south- and southeastwards into Africa and S Asia, giving rise to the mesic to xeric *Cicerbita* II, *Lactuca* and *Melanoseris* lineages.

Molecular clock calculations estimate the age of the most recent common ancestor of subtribe Lactucinae, as the youngest branch of the core group of tribe Cichorieae (clades 4 and 5 according to Kilian & al. [1] and [3]), to be c. 15–4 Ma [2–3,14], thus spanning the Middle Miocene to Early Pliocene. This period is characterised

by significant tectonic events, such as the uplift of the Qinghai-Xizang Plateau in Asia, the southern part of which reached its present elevation by c. 15 Ma [73] with larger impact on climate and vegetation.

Phylogeny of the Major Lineages of the Core Lactucinae

Cicerbita lineage. The Cicerbita lineage, in our study represented by the type species of the generic names Cicerbita and Cephalorrhynchus (both species with a chromosome number of 2n = 18 [72]), constitutes the oldest diverging branch of the core Lactucinae. Since Cephalorrhynchus is part of this lineage, it can be treated as congeneric with *Cicerbita*. None of the Chinese members of the subtribe included in our study is part of this clade. Altogether twelve species have been classified in the two genera by Shih [12] or in *Cicerbita* by Shih & Kilian [13], respectively. Four of them, from N China, are not included in the present study (compare Shih & Kilian pp214-215 [13]), but the eight species included are all nested either in the Cicerbita II (CII) clade, the Melanoseris (M) clade or the Paraprenanthes clade (P); these are: Cephalorrhynchus albiflorus, C. macrorhizus and C. saxatilis (M), Cicerbita azurea (CII), C. sikkimensis (M) and C. oligolepis (P) of Shih [12], and Cicerbita auriculiformis, C. azurea and C. roborowskii (CII) of Shih & Kilian [13].

Cicerbita, established as early as 1822 by Wallroth, appeared vaguely defined right from the beginning, including eight, partly very different species, and soon came in competition with Mulgedium, established for a similar heterogenous assemblage of species by Cassini in 1824, which then displaced the name Cicerbita during the 19th century. Through the revision by Beauverd [21], where the name Cicerbita was taken up again, it received its widest circumscription in the history of Lactucinae systematics, diagnosed solely by a pappus composed of an outer row of minute hairs and an inner row of bristles. Later, this feature was characterised by Stebbins [74] as similar useless for generic delimitation as the presence or absence of an achene beak, because it separates species that are closely allied beyond any doubt. It was, however, still employed, e.g. by Tuisl [16] to delimit the genera Cephalorrhynchus, Cicerbita and Steptorhamphus with an outer row of minute hairs from Lactuca, Mulgedium and Scariola without such an outer row (see below). Stebbins [74], in an initial attempt to redefine Cicerbita, in contrast established the narrowest circumscription of the genus, containing only three species, C. alpina, C. pancicii (Vis.) Beauverd and C. abietina (Boiss.) Stebbins, that all have columnar achenes with 5 equal main ribs, coarse pappus hairs and a C. alpina habit. A revised concept of the genus will be provided by Kilian & al. (unpublished data).

Cicerbita II lineage. Based on our initial ITS phylogeny with largely unresolved relationships of the major lineages, Shih & Kilian [13] assigned an assemblage of seven, mainly N Chinese species, comprising one species with certain affinity and three very little known species with assumed affinity to Chaetoseris roborowskii (≡ Cicerbita roborowskii), plus Cicerbita azurea and C. tianschanica, tentatively to Cicerbita. It is clear from our analyses, which represents three species of this assemblage (the species pair Cicerbita auriculiformis and C. roborowskii, plus C. azurea), that they constitute a separate lineage clearly distant from Cicerbita. Whether the remaining species of that assemblage share this positions, has still to be seen. Study of the type material of Chaetoseris rhombiformis, treated as a member of Melanoseris by Shih & Kilian [13], revealed that it is actually conspecific with C. roborowskii. The phylogeny of this predominantly Central Asian lineage, as well as its circumscription, nomenclature and classification will be treated in a consecutive paper on the global phylogeny and systematics of subtribe Lactucinae (unpublished data).

Lactuca lineage. Lactuca is not only the name-giving genus of the Lactucinae, its circumscription and delimitation is also crucial for the generic classification of the subtribe. Its circumscription varied extraordinarily in the history of the systematics of the Lactuca alliance. An extremely broad concept of Lactuca was introduced by Bentham [75] and maintained by Hoffmann [5], not only spanning most of the known diversity of the entire present-day subtribe but even including genera and species today placed into subtribes Crepidinae and Hyoseridinae. Very narrow concepts, in contrast, were advocated, in particular, by Tuisl [16] and Shih [23–24], who generically separated a number of elements from the core of Lactuca. Moderately wide concepts were established by Stebbins [74,76–77] and Ferákova [78].

The genus has never been revised in its entirety, and all four last mentioned authors only dealt with regional subsets of the genus. Because of its economic importance, many studies and also the first molecular studies [70,79-80] focused on the lettuce "gene pool" [81], which constitutes the core of Lactuca. Koopman & al. [70] provides the only molecular phylogeny of the genus available to date and is based on nrITS1. The results of their analysis are corroborated by our phylogeny based on the entire nrITS region and a small but representative sampling of Lactuca. Three well supported major clades are revealed: (1) One (JK = 99.9, PP = 1) comprises the lettuce, Lactuca sativa, which provides the type of the generic name, as well as its primary, secondary and tertiary gene pool [70]. Their distribution is centred in Europe, the Mediterranean and SW Asia and all are diploids with 2n = 18. This clade includes the type species of the segregates Scariola (S. viminea = Lactuca viminea), Mulgedium (M. runcinatum = Lactuca tatarica) and Lagedium (L. sibiricum \equiv Lactuca sibirica). (2) The second (JK = 100, PP = 1) clade comprises the E Asian *Lactuca indica* and its relatives, which were generically separated from Lactuca by Shih [23] as Pterocypsela. This clade is the dominant representative of the genus in E Asia and replaces the first clade there. Its entire species are likewise diploid with 2n = 18. Both clades together form a clade with less statistical support (JK = 54.4 PP = 98) than the individual clades have themselves. (3) The third clade (JK = 94.7, PP = 1) in turn is sister to the former two clades and has the highest number of species, of which only few are represented in our study. In contrast to the first two clades, it comprises subclades with chromosome numbers of 2n = 18, 2n = 16 and 2n = 34, the last one apparently by alloploidisation. Its members have a pappus with an outer ring of minute hairs or not, while all members of the first two clades uniformly lack such an outer ring. It includes the type species of the segregates Steptorhamphus (S. tuberosus = Lactuca tuberosa) and Lactucella (L. undulata \equiv Lactuca undulata), the Asian L. dissecta and L. dolichophylla, both also present in China, the widespread African L. inermis Forssk. (= L. capensis), L. perennis L. and, not shown here, other European, Mediterranean and SW Asian species as well as the group of native North American species with a chromosome number of 2n = 34 ([72]; unpublished

Our plastid tree, which is the first one with a selection of *Lactuca* species published, reveals a polytomy of six clades. Differences to the topology of the ITS tree are: (a) the *L. sibirica-L. tatarica* subclade clusters with the *L. perennis* clade, although with weak support (JK = 65.1, PP = 0.84), but not with the *L. sativa* clade. This is less consistent with the hybridisation experiments reviewed by Koopman & al. [70], which place *L. tatarica* into the secondary lettuce gene pool because it produces fertile hybrids when somatically hybridised with *L. sativa*, but place *L. perennis* outside the lettuce gene pool because it is not crossable with *L. sativa* primary gene pool species. (b) *L. inermis* is not nested in the *Steptorhamphus tuberosus-L. dissecta* clade but constitutes a branch of

its own. (c) As noted already above, *Mulgedium bracteatum* of the *Melanoseris* lineage in the ITS tree is nested here in the *Lactuca* lineage as a further separate branch.

While the ITS tree is inconclusive with respect to the relationship of the *Lactuca* lineage with other major lineages in the core Lactucinae, the plastid tree indicates a highly supported sister group relationship (JK = 99.9, PP = 1) between the *Lactuca* and the *Melanoseris* lineages. The two lineages themselves only receive moderate support, *Lactuca* (JK = 68.8, PP = 0.98) still less than *Melanoseris* (JK = 81.7, PP = 1). Exclusion of *M. bracteatum* from the analysis does not affect the statistical support of either lineage and since *M. bracteatum* is a diploid species (2n = 16 [72]), the reason for its incongruent position may perhaps be chloroplast capture through introgressive hybridisation. A sister group relationship with *Melanoseris* is also supported by morphology, where differences between the two lineages are particularly difficult to define.

Lactuca is a suitable example to elucidate the shortcomings of the previous classification attempts in the Lactuca alliance with the molecular phylogenetic results. Although it is evident that the achene as dispersal unit faces a particularly strong exposure to selection pressure and corresponding morphological changes affecting their functionality [82], a very static, sometimes even typological, understanding of achene features has often enough pervaded the taxonomy of the *Lactuca* alliance. Absence of a true achene beak and a weakly compressed achene body were the main features for the separation of Mulgedium (L. tatarica and L. sibirica [16], somewhat altered concept by Shih [23]) or Lagedium (including L. tatarica and L. sibirica [83] or L. sibirica only [23]); the combination of a compressed achene body, winged lateral ribs and a beak justified the separation of the E Asian Lactuca lineage as Pterocypsela [23], and the apomorphy of two rod-like, pendent basal appendages at the long-beaked achene apex justified separation of L. undulata as monotypic genus Lactucella [84]. A relict of 19th classification, where schematically pappus features were in use for classification at generic and higher ranks, is the use in the Lactuca alliance of the absence of an outer row of minute hairs in the pappus to delimit Lactuca from Steptorhamphus as well as from, in particular, Cephalorrhynchus and Cicerbita [5,16,21,23,25,78]. The Steptorhamphus tuberosus-L. dissecta clade is an example, where even a single, well supported clade, revealed both in the nuclear and plastid phylogenies, unites members having a pappus with (S. tuberosus) and without (L. dissecta, L. dolichophylla) an outer ring of minute hairs. The segregation of the L. viminea-L. orientalis species group as Scariola for the low number of 4 or 5 flowers per capitulum along with white stems and adnately decurrent leaves, in contrast, appears morphologically much more plausible, yet is equally unsubstantiated in the light of the molecular phylogenetic results. All these former segregates are deeply nested in Lactuca according to both the nuclear and plastid phylogenies.

Among the E Asian Lactuca indica species group, different species concepts, which depend on the evaluation of conspicuous leaf shape differences found, have been applied recently and consequently different numbers of species recognised. Whereas Shih [12,23] recognised seven species (under Pterocypsela), Shih & Kilian [13] reduced them to only four, considering the otherwise similar plants with entire-leafy and pinnately lobed leaves only as extremes of infraspecific ranges of variation. The latter authors therefore sunk L. elata (with entire leaves) into L. raddeana (with lyrately or pinnately lobed leaves), L. laciniata (with strongly pinnately lobed leaves) into L. indica (with mostly entire leaves) and L. sonchus (with entire leaves) into L. formosana (with strongly pinnately lobed leaves). Using the narrower species concepts in our analyses, which includes all species of the group but L. triangularis,

both phylogenies link with high support *L. laciniata* and *L. indica* (JK = 99.8, PP = 1 in ITS tree; JK = 99.5, PP = 1 in plastid tree). The ITS phylogeny also links with high support *L. elata* and *L. raddeana* (JK = 96.6, PP = 1), only *L. sonchus* and *L. formosana* are linked with weak support (JK <50, PP = 0.52). The plastid phylogeny in contrast links *L. elata* with *L. sonchus* and *L. formosana* with weak support (JK = 63.5, PP = 1). These results in combination with the low amount of sequence variation involved among the six *Pterocypsela* samples (12 variable sites, 11 informative in the nuclear data set; 10 and 6 informative in the plastid data set) can be seen as an additional support for the hypothesis of wide ranges of infraspecific leaf shape variation and consequently wider species concepts at least in the first case, while the other cases deserve further studies because of and also with respect to the ambiguous position of *L. elata*.

Melanoseris lineage. The genus Melanoseris (for exemplar species see Fig. 4) was established by Decaisne in 1843 to include two species from the Himalayas, which are now treated as a single species, M. lessertiana. It was considered to differ from Cicerbita (then under the name Mulgedium) because of its beaked achenes and from Lactuca, because of its pappus with an outer series of minute hairs. Edgeworth [85] added a few more Himalayan species, which we confirm to belong to this lineage, but afterwards the use of the name Melanoseris was abandoned. The name was only recently revived by Shih & Kilian [13] for this lineage, based on our initial ITS phylogeny, through which it became evident that the types of the newly established genera Chaetoseris and Stenoseris by Shih [25] are part of one lineage with M. lessertiana, which provides the type of the name Melanoseris and was treated by Shih [12] under Mulgedium. Shih's genera Chaetoseris and Stenoseris are, moreover, shown in our analyses to be actually bi- and triphyletic, respectively (Fig. 1-2). Apart from the bulk of the Chaetoseris species nested in the Melanoseris lineage, one species, C. roborowskii (including also C. rhombiformis), is nested in the Cicerbita II lineage. Chaetoseris was circumscribed and delimited from Lactuca and Cicerbita by the combination of beaked achenes, an achene corpus with broad, thickened lateral ribs and a pappus with an outer series of minute hairs [25]. Shih's six Stenoseris species are distributed among the Melanoseris lineage (S. graciliflora, S. taliensis, S. tenuis), the Paraprenanthes lineage (S. leptantha, S. triflora) and the Cicerbita II lineage (S. auriculiformis). Stenoseris was circumscribed by the combination of narrowly cylindrical, 3-flowered capitula, an achene corpus with broad, thickened lateral ribs and a pappus with an outer series of minute hairs [25]. All features used to circumscribe the two genera are clearly shown to be homoplastic. It is therefore not surprising that, compared to Shih [12,25], the Melanoseris lineage, moreover, includes all species of Cephalorrhynchus (distinguished by Shih through the achene corpus lacking thick, broadened lateral ribs), one of Cicerbita (C. sikkimensis) and three species of Mulgedium (distinguished by Shih through the pappus lacking an outer row of minute hairs, M. bracteatum, M. lessertianum and M. monocephalum).

Melanoseris constitutes a large, well supported lineage (JK = 89.1, PP = 1 in the ITS phylogeny, JK = 81.7, PP = 1 in the plastid phylogeny). Most species, in particular all its Sino-Himalayan members, are diploid with 2n = 16 ([72]; under Chaetoseris and Stenoseris [86]), otherwise a number of species also has 2n = 18 (unpublished data). Besides the Sino-Himalayan species, Melanoseris also includes S, SW and Middle Asian as well as African species (unpublished data), but our present sampling is restricted chiefly to the species occurring in China. Mulgedium bracteatum, which appears in the ITS tree of the global sampling (unpublished data, there also with a second sample) within a further basally branching clade of SW and Middle Asian species, therefore takes



Figure 4. Selected species of *Melanoseris* in situ. A and C, *Melanoseris* atropurpurea (Yunnan, 9 Sep. 2009, photo by Z. J. Yin; voucher: *Z. J. Yin* & al. 1970 (KUN)), B. M. likiangensis (Sichuan, 23 Aug. 2012, photo by N. Kilian; voucher: N. Kilian & al. 10808 (B, KUN), D. M. cyanea (Yunnan, 22 Sep. 2011, photo by G. X. Hu; voucher: H. J. Dong & al. 446 (KUN), E, M. tenuis (Yunnan, 10 Sep. 2009, photo by Z. J. Yin; voucher: Z. J. Yin & al. 1969 (KUN)), F, M. graciliflora (Sichuan, 19 Aug. 2012, photo by N. Kilian; voucher: N. Kilian & al. 10509 (B, KUN). doi:10.1371/journal.pone.0082692.g004

an isolated, basally branching position in the present ITS tree. The strikingly incongruent position in the plastid phylogeny as a member of the *Lactuca* lineage deserves further investigation. From the morphological evidence we consider the nuclear phylogeny as the better estimate for the species phylogeny.

The next following branch, congruently revealed by the nuclear and plastid phylogeny, is sister with robust support (JK = 99.8, PP = 1 in the ITS phylogeny; JK = 93.8, PP = 0.82 in the plastid phylogeny) to all other Sino-Himalayan species of the lineage and consists of *Parasyncalathium souliei* only. Originally described as *Lactuca souliei* in 1895, the attractive bright blue-flowered acaulescent alpine species was placed together with habitually and ecologically strikingly similar species in *Lactuca* sect. *Aggregatae*, which later became the separate genus *Syncalathium*. Stebbins

(pp47–50 [87]) inferred from achene morphology, Zhang & al. [88] from karyology, and Kilian & al. (pp348–350 [1]) and Zhang & al. [2,14] from molecular phylogeny, all provided evidence that *L. souliei* is entirely unrelated to the other species of *Syncalathium* and that their overall similarity is hence a result of convergent evolution, presumably in response to the environmental changes following the uplift of the Qinghai-Xizang Plateau. Kilian & al. [1] recognised the species as a member of subtribe Lactucinae rather than of Crepidinae, to which *Syncalathium* belongs to, and Shih & Kilian [13] later placed it into *Melanoseris*, while Zhang & al. [14], arguing with its peculiar morphology, accommodated it in their newly established genus *Parasyncalathium*. Our analyses presented here do not provide unambiguous support for either classification.

For the time being, we prefer to maintain its inclusion in *Melanoseris*.

The bulk of the Sino-Himalayan species all appear in a large polytomy in the ITS phylogeny, with only two subclades that comprise samples of more than one species (Fig. 1: M-A and M-B). The plastid phylogeny provides higher resolution for the lineage and shows four major subclades with well support (Fig. 2: M-1 to M-4). None of the subclades that comprise samples of more than one species, however, is fully congruent with either subclade revealed in the ITS tree.

- Melanoseris cyanea group: The larger of the two subclades of the ITS tree (clade M-A) includes all but four samples (as Chaetoseris cyanea hybrid_LAC094-097 in the tree) that belong to the M. cyanea group of clade M-4 in the plastid phylogeny. The core of the group congruently revealed in both phylogenies contains a number of taxa, morphologically clearly allied to M. cyanea (Fig. 4D). Morphological variation within this group of robust tall forbs in particular regards indumentum, leaf shape, size of capitula and number of flowers per capitulum, flower colour, and length of the anther tube. Delimitation of taxa is very problematic due to a lack of clear morphological discontinuities. These may, however, be the results of previous areal changes with subsequent events of hybridisation and introgression, processes that are apparently still ongoing. Notably, the aforementioned four sympatric samples (as Chaetoseris cyanea hybrid_LAC094-097 in the tree) of the M. cyanea group, which fall into the large polytomy of the Sino-Himalayan species in the ITS phylogeny, have a number of additive polymorphic sites in their ITS sequences (Table 2), indicating the occurrence of still divergent, nonhomogenised ITS paralogues likely as a result of nuclear gene flow, and this finding corresponds to the presence of intermediate morphological characters states, because of which these samples do not match either of the species distinguished and were therefore designated as putative hybrids. Even the already widened species concepts by Shih & Kilian [13], compared to Shih [12], do not work when confronted with the variation actually encountered in the field across the distribution area of the M. cyanea group in China. The lacking molecular resolution within this group thus corresponds well to the lack of morphological discontinuities and makes further taxonomic adjustments necessary (see Taxonomic conclusion).
- (2) Melanoseris macrorhiza group: In the ITS phylogeny M. macrorhiza (= Cephalorrhynchus macrorhizus in Shih 1997) clusters together with M. violifolia (= Cicerbita sikkimensis in Shih 1997) and M. lessertiana (providing the type of Melanoseris) in a well supported (JK = 93.1, PP = 1) clade (Fig. 1: clade M-B). In the plastid phylogeny, in contrast, this clade does not exist at all but the three species occur in three different clades (Fig. 2: M-1, M-3, M-4). M. lessertiana instead forms a clade with full support (JK = 100, PP = 1) together with M. qinghaica (\equiv Mulgedium qinghaicum [89]). M. qinghaica actually represents Mulgedium lessertianum in the sense of Shih [12] and the Chinese populations of Melanoseris lessertiana in the sense of Shih & Kilian [13], and replaces entirely the latter species in China. In the ITS tree M. qinghaica forms a separate branch within the large polytomy of the Sino-Himalayan species. Morphologically M. lessertiana and M. qinghaica have apparent close affinities to each other and are mainly distinguished by the distinctly longer achene beak and very short anthertube of M. qinghaica. It thus appears that the plastid phylogeny in this respect is more in line with morphology. Inferred from

- morphology, however, all four aforementioned species are considered to be more closely related to each other, as is revealed in the ITS tree for three of them. They are all rather low growing herbs usually without a dominant main stem.
- Melanoseris graciliflora group: The morphologically closely allied, few-flowered species pair M. graciliflora (Fig. 4F; Stenoseris graciliflora [12]) and M. tenuis (Fig. 4E; Stenoseris tenuis [12]) is nested in the plastid phylogeny (Fig. 2: clade M-2) in a clade together with several species having capitula with many to numerous (M. atropurpurea, Fig. 4A+C) flowers and usually clasping stem leaves. All are robust tall forbs with cyanic flowers. In the ITS tree the members of this clade all form separate branches in the large polytomy except for the multiple samples of M. graciliflora and M. tenuis. Morphology makes this clade in the plastid tree neither obvious nor unlikely, at least if we accept also more drastic changes in the flower number per capitula as a common trend in character evolution, what we certainly have to do. We may hence accept the inferred relationship as a hypothesis for further studies, but also taking into consideration that relationships in Sino-Himalayan Melanoseris may be blurred by events of hybridisation and introgression. A number of well detected additive polymorphic sites in the ITS sequences of Stenoseris tenuis hybrid_LAC-108 and 109 (Table 2), plus the intermediate morphological characters (especiall the number of inner phyllaries), in combination with the first author's observation in the field that some typical plants of M. cyanea, M. tenuis and M. atropurpurea co-occurred in the same habitat, all indicates introgressive hybridisation between populations of these taxa, which accounts for the incongruent positions of these two hybrid individuals in the ITS and plastid phylogenies.

Melanoseris is not only the largest lineage of Lactucinae in China, but we have experienced it also taxonomically as particularly difficult. It comprises, on the one hand, elements that are morphologically so diverse that their affinities let alone relationships are far from obvious, on the other hand elements that constitute rather uniform groups in which the differences are predominantly gradual rather than clear-cut or of qualitative nature, and delimitations thus are often difficult to establish. Our molecular phylogenetic analyses provide the first indications that hybridisation and reticulate evolution could be one cause of this situation.

Notoseris lineage. The genus Notoseris (for exemplar species see Fig. 3A-C) was established by Shih [22] to accommodate a number of tall forb species endemic to SW China, which share a combination of morphological features that set them apart from both the genera Prenanthes and Nabalus. These features were: nodding 3-5-flowered capitula; slender cylindrical involucres with purplish red phyllaries; purplish red flowers; purplish red, fusiform, unbeaked, somewhat compressed achenes (with 5 main ribs and 2 secondary ribs in between); pappus without an outer row of minute hairs. In his revised treatment, Shih [12] accepted 11 (plus two doubtful) species of Notoseris. Shih & Kilian [13] reduced this number to seven by changing several species concepts and transferring one species to Paraprenanthes. As a conclusion from our then initial ITS phylogeny, Shih & Kilian [13] added the two scandent species (Fig. 3B-C), formerly treated as Prenanthes scandens and P. yakoensis to Notoseris, which extended the original circumscription of the genus to capitula with up to 12 flowers and also non-purplish red but pale brown achenes. Their inclusion is corroborated by the present extended analysis. The scandent Notoseris species are not related to the scandent species of the subtribe in Africa and Indonesia, evolution of the scandent habit in subtribe Lactucinae thus has apparently occurred independently three times from montane tall forb ancestors (unpublished data).

Our phylogenetic analysis revealed that *Notoseris* in the revised sense of Shih & Kilian [13] is still not monophyletic. Two species, *N. melanantha* and *N. wilsonii*, are nested instead in the *Paraprenanthes* clade of the ITS phylogeny, or in the *Paraprenanthes* subclades of the joined *Notoseris-Paraprenanthes* clade of the plastid tree, and have thus to be excluded from *Notoseris* and transferred to the genus *Paraprenanthes* as *P. melanantha* and *P. wilsonii* (Fig. 3D–E; see Taxonomic conclusions). Possible causes of the topological incongruences were discussed above.

Additional evidences gathered in the present study from the taxonomic revision of all types and extensive studies of the species in the field, which support the molecular results, urge us to a further revision of the species concepts compared to both Shih [12] and Shih & Kilian [13]. It became obvious that discontinuities, of leaf features especially, inferred from the herbarium material of these tall forbs by Shih [12,22] and used for the delimitation of species, frequently break down when variation is studied in the field. As herbarium specimen preservation of tall forbs (often exceeding 2 m in height) was in the past usually done highly selectively, even intraindividual variation of leaf shapes from the base to the top of the main axis was rarely documented to a sufficient extent, while leaf shape played an important role in the taxonomic treatments by Shih [12,22]. Consequently, four other species compared to the last treatment by Shih & Kilian [13] are sunken in the synonymy here, leaving *Notoseris* with a total of six species only. So far known, all species are diploids with 2n = 18[67]. The genus has its centre of diversity in SW China, where all six species occur. Four of them are endemic to China, the other two species also touch neighbouring countries.

Paraprenanthes lineage. The genus Paraprenanthes (for exemplar species see Fig. 3D-F) was formally established by Shih [24], based on an earlier proposal by C. C. Chang, segregating species from Lactuca that are morphologically allied to L. sororia, which he designated as the type of the name Paraprenanthes. These species are usually tall forbs, they have usually nodding capitula with 3 (in the revised circumscription established here, 4 according to Shih [12])-15 cyanic flowers, slender cylindrical involucres, fusiform, somewhat compressed unbeaked dark brown to blackish achenes with 5 main ribs and 2 rather similar secondary ribs in between, and a pappus without an outer row of minute hairs. Formerly 11 species were distinguished by Shih [24], most of which newly described, the number increased to 15 finally [12]. Shih's circumscription of the genus was maintained by Shih & Kilian [13], apart from the transfer of one species from Notoseris and the addition of a second one, following Sennikov [90], from Mulgedium, but somewhat wider species concepts were established, reducing the species number to 12.

Inferred from our analysis, the recent additions to the genus by Sennikov [90] and Shih & Kilian [13] are corroborated, but as hitherto circumscribed, *Paraprenanthes* is clearly paraphyletic. One group of species previously placed in *Notoseris (N. melanantha/wilsonii* group) and a second group of two species (*Cicerbita oligolepis/Stenoseris triflora* group) formerly placed in *Stenoseris* and *Cicerbita* [12,25] or *Melanoseris* [13], respectively, must also be transferred to *Paraprenanthes* according to the evidence from both the nuclear and the chloroplast phylogeny. Although *Notoseris* and *Paraprenanthes* form a joined clade in the latter (see discussion, above), both groups clearly cluster with the respective *Paraprenanthes* subclades. The consequences for the morphological circumscription of the genus are, however, less significant, owing to the anyway shallow morphological divisions between the major lineages of the subtribe, and mainly concern the achenes, which can also be

shortly beaked and pale brown. The case of putative introgressive hybridisation involving *Paraprenanthes melanantha* (as *Notoseris melanantha* in the trees), *P. wilsonii* (as *N. wilsonii* in the trees) and *P. sororia*, is discussed above.

Similar to the situation in Notoseris, the core of Paraprenanthes forms a polytomy in the ITS tree with the terminal taxa in most cases found individually on short or very short branches, reflecting the few character state differences in this marker sequence, whereas somewhat more resolution is provided by the plastid tree. The molecular evidence is in good accordance with the phenetic evidence, in so far as (a) speciation among the core of Paraprenanthes has not yet, in most cases, led to more conspicuous discontinuities, and (b) that even the wider species concepts applied by Shih & Kilian [13] compared to Shih [12] are still too narrow for quite similar reasons as stated for *Notoseris*. Supported by the taxonomic revision of all types, extensive studies of the species in the field, our revised taxonomy of *Paraprenanthes* halves the number of its species recognised by Shih & Kilian [13] to six. Adding the species newly to be transferred to this genus, we now recognise 10 species in Paraprenanthes, eight of which are endemic to China while two, P. sororia and P. umbrosa, extend to Vietnam and Japan, and Myanmar(?) and Thailand, respectively. So far known, all species are diploids with 2n = 18 [67]. The single exception of a chromosome count of 2n = 16 by Deng & al. [86] for Stenoseris leptantha, which is a synonym of Paraprenanthes triflora, vouchered by the specimen Nie 1159 (KUN!), actually represents Melanoseris tenuis.

Taxonomic Conclusions

Concluding from our molecular and morphological analyses, the latter also including the study of the type material of the names involved, we outline here a new classification of the genera *Notoseris, Paraprenanthes* and *Melanoseris* in their Chinese centre of diversity. It revises the recent classification of these genera by Shih & Kilian [13]. Full synonymies and further data are available through the Cichorieae Portal [27]. Monographic treatments of these genera are in preparation and will be the subject of consecutive publications.

1. Notoseris. C. Shih in Acta Phytotax. Sin. 25: 196. 1987. – Type: *Notoseris psilolepis* C. Shih [= *N. macilenta*].

6 species, all in China, 3 endemic (=*).

Distribution: China (Chongqing, Guangdong, Guangxi, Guizhou, Hubei, Hunan, Jiangxi, Sichuan, Taiwan, Xizang, Yunnan) and E Himalaya region.

- (1) **Notoseris yakoensis** (Jeffrey) N. Kilian in Wu & al., Fl. China 20–21: 231. 2011 ≡ *Prenanthes yakoensis* Jeffrey in Notes Roy. Bot. Gard. Edinburgh 5: 203. 1912.
 - = Prenanthes volubilis Merr.
- (2) **Notoseris scandens** (Hook. f.) N. Kilian in Wu & al., Fl. China 20-21: 231. $2011 \equiv Prenanthes scandens$ Hook. f. in Bentham & Hooker, Gen. Pl. 2: 527. 1873.
- (*3) **Notoseris triflora** (Hemsl.) C. Shih in Acta Phytotax. Sin. 25: 202. 1987 \equiv *Lactuca triflora* Hemsl. in J. Linn. Soc., Bot. 23: 485. 1888.
- (4) **Notoseris khasiana** (C. B. Clarke) N. Kilian in Wu & al., Fl. China 20–21: 233. 2011 ≡ *Prenanthes khasiana* C. B. Clarke, Comp. Ind.: 273. 1876.
 - = Notoseris rhombiformis C. Shih, **syn. nov.**
- (*5) **Notoseris macilenta** (Vaniot & H. Lév.) N. Kilian in Wu & al., Fl. China 20–21: 231. 2011 ≡ *Prenanthes macilenta* Vaniot & H. Lév. in Bull. Soc. Bot. France 53: 550. 1906.
 - = Notoseris psilolepis C. Shih
 - = Notoseris formosana (Kitam.) C. Shih
 - = Notoseris nanchuanensis C. Shih, **syn. nov.**

- = Notoseris guizhouensis C. Shih, **syn. nov.**
- = Notoseris yunnanensis C. Shih, syn. nov.
- (*6) **Notoseris henryi** (Dunn) C. Shih in Acta Phytotax. Sin. 25: 202. 1987 ≡ *Prenanthes henryi* Dunn in J. Linn. Soc., Bot. 35: 514. 1903.
 - = Notoseris porphyrolepis C. Shih, syn. nov.

Excluded species:

Notoseris melanantha (Franch.) C. Shih in Acta Phytotax. Sin. 25: 198. 1987 ≡ **Paraprenanthes melanantha** (Franch.) Z. H. Wang

Notoseris wilsonii (C. C. Chang) C. Shih in Acta Phytotax. Sin. 25: 202. 1987 ≡ **Paraprenanthes wilsonii** (C. C. Chang) Z. H. Wang

- **2. Paraprenanthes.** C. C. Chang ex C. Shih in Acta Phytotax. Sin. 26: 418. 1988. Type: *Paraprenanthes sororia* (Miq.) C. Shih.
- = Lactuca sect. Prenanthesiae Franch. in J. Bot. (Morot) 9: 291. 1895. Lectotype (**here designated**): Lactuca melanantha Franch. 10 species, all in China, 8 endemic (=*).

Distribution: China (Anhui, Chongqing, Fujian, Guangdong, Guangxi, Guizhou, Hainan, Hubei, Hunan, Jiangsu, Jiangxi, Shanxi, Sichuan, Taiwan, Xizang, Yunnan, Zhejiang), the E Himalayan region, Myanmar, Thailand, Vietnam and Japan.

- (*1) *Paraprenanthes oligolepis* (C. C. Chang ex C. Shih) Z. H. Wang, **comb. nov.** ≡ *Cicerbita oligolepis* C. C. Chang ex C. Shih in Acta Phytotax. Sin. 29: 398. 1991 ≡ *Melanoseris oligolepis* (C. C. Chang ex C. Shih) N. Kilian, **syn. nov.**
- (*2) Paraprenanthes triflora (C. C. Chang & C. Shih) Z. H. Wang & N. Kilian, comb. nov. ≡ Stenoseris triflora C. C. Chang & C. Shih in Acta Phytotax. Sin. 29: 413. 1991 ≡ Melanoseris triflora (C. C. Chang & C. Shih) N. Kilian, syn. nov.
- = Stenoseris leptantha C. Shih ≡ Melanoseris leptantha (C. Shih) N. Kilian, syn. nov.
- (3) **Paraprenanthes umbrosa** (Dunn) Sennikov in Bot. Zhurn. 82(5): 111. 1997 ≡ *Lactuca umbrosa* Dunn in J. Linn. Soc., Bot. 35: 513. 1903 ≡ *Mulgedium umbrosum* (Dunn) C. Shih
- =? Lactuca parishii Craib in Kew Bull. 1911: 403. 1911, syn. nov.
- (4) **Paraprenanthes sororia** (Miq.) C. Shih in Acta Phytotax. Sin. 26: 422. 1988 ≡ *Lactuca sororia* Miq. in Ann. Mus. Bot. Lugduno-Batavi 2: 189. 1866
 - = Paraprenanthes pilipes (Migo) C. Shih
 - = Paraprenanthes multiformis C. Shih, **syn. nov.**
- (*5) **Paraprenanthes diversifolia** (Vaniot) N. Kilian in Wu & al., Fl. China 20–21: 229. 2011 ≡ *Lactuca diversifolia* Vaniot in Bull. Acad. Int.Geogr. Bot. 12: 245. 1903
 - = Paraprenanthes sylvicola C. Shih
 - = Paraprenanthes heptantha C. Shih & D. J. Liou, syn. nov.
 - = Paraprenanthes gracilipes C. Shih
- (*6) **Paraprenanthes yunnanensis** (Franch.) C. Shih in Acta Phytotax. Sin. 26: 421. 1988 \equiv Lactuca yunnanensis Franch. in J. Bot. (Morot) 9: 264. 1895
 - = Paraprenanthes sagittiformis C. Shih
 - = Paraprenanthes longiloba Y. Ling & C. Shih, syn. nov.
 - = Paraprenanthes auriculiformis C. Shih, **syn. nov.**
- (*7) **Paraprenanthes prenanthoides** (Hemsl.) C. Shih in Acta Phytotax. Sin. 26: 423. 1988 ≡ *Crepis prenanthoides* Hemsl. in J. Linn. Soc., Bot. 23: 477. 1888
 - = Paraprenanthes glandulosissima (C. C. Chang) C. Shih, **syn. nov.**
 - = Paraprenanthes polypodiifolia (Franch.) C. Shih, **syn. nov.**
 - = Paraprenanthes thirionnii (H. Lév.) C. Shih
 - = Paraprenanthes luchunensis C. Shih, syn. nov.

- (*8) **Paraprenanthes meridionalis** (C. Shih) Sennikov in Bot. Zhurn. 82(5): 111. 1997 ≡ *Mulgedium meridionale* C. Shihin Acta Phytotax. Sin. 26: 392. 1988
 - = Paraprenanthes hastata C. Shih, syn. nov.
- (*9) **Paraprenanthes melanantha** (Franch.) Z. H. Wang, **comb. nov.** ≡ *Lactuca melanantha* Franch. in J. Bot. (Morot) 9: 291. 1895 ≡ *Notoseris melanantha* (Franch.) C. Shih

Note: This species was misinterpreted by Shih & Kilian [13], where it was treated, under *Notoseris*, in a wide sense, merged with actually unrelated other species that have similar pinnately divided leaves. In the sense of its type, in contrast, it is a species endemic to Sichuan and Chongqing, well characterised by the combination of (a) a strikingly narrow, paniculiform, densely glandular synflorescence, (b) leaves with a large triangular to ovate or rhombic, basally cordate to cuneate terminal segment and 0–3(–6) pairs of lateral segments on a winged rachis, and (c) achenes with a \pm truncate to attenuate apex.

- (*10) **Paraprenanthes wilsonii** (C. C. Chang) Z. H. Wang, **comb. nov.** ≡ *Prenanthes wilsonii* C. C. Chang in Bull. Fan Mem. Inst. Biol., Bot. 5: 322. 1934 ≡ *Notoseris wilsonii* (C. C. Chang) C. Shih
 - = Notoseris gracilipes C. Shih
- = Paraprenanthes dolichophylla (C. Shih) N. Kilian & Z. H. Wang in Wu & al., Fl. China 20–21: 229. 2011, **syn. nov.** = Notoseris dolichophylla C. Shih

Note: Paraprenanthes dolichophylla is apparently very closely related to *P. wilsonii* and pending further assessment, is tentatively considered as conspecific here.

- **3. Melanoseris.** Decne. in Jacquemont, Voy. Inde 4: 101. 1843. Lectotype (designated by Pfeiffer, Nomencl. Bot. 2: 259. 1874): *Melanoseris lessertiana* (DC.) Decne.
- = Chaetoseris C. Shih in Acta Phytotax. Sin. 29: 398. 1991. Type: Chaetoseris lyriformis C. Shih [= Melanoseris cyanea s.l.]
- = Stenoseris C. Shih in Acta Phytotax. Sin. 29: 411. 1991. − Type: Stenoseris graciliflora (DC.) C. Shih [≡ Melanoseris graciliflora]
- = Parasyncalathium J. W. Zhang & al. in Taxon 60: 1680. 2011. − Type: Parasyncalathium souliei (Franch.) J. W. Zhang & al. [≡ Melanoseris souliei]

Some 70 species in total, 17 species in China, 9 endemic (=*). Distribution: China (Chongqing, Guizhou, Sichuan, Xizang, Yunnan); Himalayas and adjacent areas, SW and Central Asia, sub-Saharian Africa.

Notes: In the Himalayan territories the following seven species of *Melanoseris* are distributed but not known to occur in China: M. brunoniana (Wall. ex DC.) N. Kilian & Z. H. Wang, comb. nov. Prenanthes brunoniana Wall. ex DC., Prodr. 7(1): 195. 1838; M. decipiens (Hook. f. & Thomson ex C. B. Clarke) N. Kilian & Z. H. Wang, **comb. nov.** \equiv *Lactuca decipiens* Hook. f. & Thomson ex C. B. Clarke, Compos. Ind.: 266. 1876; M. filicina (Stebbins) N. Kilian, **comb. nov.** ≡ *Lactuca filicina* Duthie ex Stebbins in Indian Forest Rec., Bot. 1: 241. 1939; M. kashmiriana (Mamgain & R. R. Rao) N. Kilian, **comb. nov.** \equiv *Lactuca kashmiriana* Mamgain & R. R. Rao in J. Bombay Nat. Hist. Soc. 83: 406-408. 1986; M. lahulensis (Mamgain & R. R. Rao) N. Kilian, comb. nov. = Lactuca lahulensis Mamgain & R. R. Rao in Bull. Bot. Surv. India 27: 120-122. 1987; M. polyclada (Boiss.) Akhani, N. Kilian & Sennikov, **comb. nov.** ≡ *Zollikoferia polyclada* Boiss., Fl. Orient. 3: 827. 1875; M. rapunculoides (DC.) Edgeworth.

- (1) *Melanoseris bracteata* (C. B. Clarke) N. Kilian in Wu & al., Fl. China 20–21: 225. 2011 ≡ *Lactuca bracteata* C. B. Clarke, Compos. Ind.: 270. 1876 ≡ *Mulgedium bracteatum* (C. B. Clarke) C. Shih
- (*2) **Melanoseris souliei** (Franch.) N. Kilian in Wu & al., Fl. China 20–21: 225. 2011 ≡ *Lactuca souliei* Franch. in J. Bot. (Morot)

- 9: 257. 1895 ≡ Syncalathium souliei (Franch.) Y. Ling ≡ Parasyncalathium souliei (Franch.) J. W. Zhang & al.
 - = Syncalathium orbiculariforme C. Shih
- (3) **Melanoseris qinghaica** (S. W. Liu & T. N. Ho) N. Kilian & Z. H. Wang, **comb. nov.** ≡ *Mulgedium qinghaicum* S. W. Liu & T. N. Ho in Acta Phytotax. Sin. 39: 556. 2001

Note: Tentatively included by Shih & Kilian [13] in a rather widely circumscribed *Melanoseris lessertiana*, our analyses since have revealed that all reports of *M. lessertiana* from China are actually referable to *M. qinghaica*, which is mainly distinguished by the distinctly longer achene beak and a very short anthertube. *M. lessertiana* is restricted to the Himalayas.

- (4) *Melanoseris cyanea* (D. Don) Edgew. in Trans. Linn. Soc. London 20: 81. 1846 ≡ *Sonchus cyaneus* D. Don, Prodr. Fl. Nepal. 164. 1825 ≡ *Chaetoseris cyanea* (D. Don) C. Shih
- = Melanoseris beesiana (Diels) N. Kilian, **syn. nov.** ≡ Chaetoseris beesiana (Diels) C. Shih
- = Chaetoseris hastata (DC.) C. Shih \equiv Melanoseris hastata (DC.) Edgew.
 - = Chaetoseris hispida C. Shih
 - = Chaetoseris lyriformis C. Shih
- = Melanoseris sichuanensis (C. Shih) N. Kilian, **syn. nov.** ≡ Chaetoseris sichuanensis C. Shih

Tentatively included:

Chaetoseris lutea (Hand.-Mazz.) C. Shih in Acta Phytotax. Sin. 29: 409. 1991 ≡ Cicerbita cyanea var. lutea Hand.-Mazz., Symb. Sin. 7: 1180. 1936.

Melanoseris yunnanensis (C. Shih) N. Kilian & Z. H. Wang in Wu & al., Fl. China 20-21: 219. $2011 \equiv Chaetoseris yunnanensis C. Shih [= Chaetoseris teniana (Beauverd) C. Shih <math>\equiv$ Cicerbita cyanea var. teniana Beauverd]

Melanoseris pectiniformis (C. Shih) N. Kilian & J. W. Zhang in Wu & al., Fl. China 20–21: 222. 2011 ≡ Chaetoseris pectiniformis C. Shih

Note: *Melanoseris cyanea* is a widespread species and polymorphic especially with respect to indumentum features. The wider concept of the species, compared to Shih [12], used by Shih & Kilian [13] is still too narrow: (1) The delimitation towards *M. beesiana* (= Chaetoseris lyriformis C. Shih) as well as towards *M. sichuanensis* breaks, when considering besides leaf shape features also relevant capitula and flower features. (2) The status of the yellow-flowered plants and populations treated by Shih & Kilian [13] under *Melanoseris yunnanensis* (= Chaetoseris lutea = C. teniana) is still not fully clear, their very close relationship to *M. cyanea* is proven, however, by the molecular analysis. (3) The status and assignment of *M. pectiniformis* are still not beyond doubt.

- (*5) *Melanoseris ciliata* (C. Shih) N. Kilian in Wu & al., Fl. China 20–21: 219. 2011 ≡ *Chaetoseris ciliata* C. Shih in Acta Phytotax. Sin. 29: 403. 1991
- (*6) *Melanoseris macrocephala* (C. Shih) N. Kilian & J. W. Zhang in Wu & al., Fl. China 20–21: 221. 2011 ≡ *Chaetoseris macrocephala* C. Shih in Acta Phytotax. Sin. 29: 404. 1991
- (7) *Melanoseris macrorhiza* (Royle) N. Kilian in Wu & al., Fl. China 20–21: 224. 2011 ≡ *Mulgedium macrorhizum* Royle, Ill. Bot. Himal. Mts. 1: 251. 1835 ≡ *Cephalorrhynchus macrorhizus* (Royle) Tuisl
 - = Cephalorrhynchus albiflorus C. Shih
- (8) **Melanoseris violifolia** (Decne.) N. Kilian in Wu & al., Fl. China 20–21: 225. 2011 ≡ *Prenanthes violifolia* Decne. in Jacquemont, Voy. Inde 4. 1843
 - = Cicerbita sikkimensis (Hook. f.) C. Shih
- (9) *Melanoseris macrantha* (C. B. Clarke) N. Kilian & J. W. Zhang in Wu & al., Fl. China 20–21: 219. 2011 ≡ *Lactuca macrantha* C. B. Clarke, Compos. Ind.: 267. 1876 ≡ *Chaetoseris macrantha* (C. B. Clarke) C. Shih

- (*10) **Melanoseris likiangensis** (Franch.) N. Kilian & Z. H. Wang in Wu & al., Fl. China 20–21: 222. 2011 ≡ *Lactuca likiangensis* Franch. in J. Bot. (Morot) 9: 259. 1895 ≡ *Chaetoseris likiangensis* (Franch.) C. Shih
- (*11) *Melanoseris bonatii* (Beauverd) Z. H. Wang, **comb. nov.** ≡ *Cicerbita bonatii* Beauverd in Bull. Soc. Bot. Genève 2: 126. 1910 ≡ *Chaetoseris bonatii* (Beauverd) C. Shih
- (12) **Melanoseris atropurpurea** (Franch.) N. Kilian & Z. H. Wang in Wu & al., Fl. China 20-21: 221. $2011 \equiv Lactuca$ atropurpurea Franch. in J. Bot. (Morot) 9: 260. $1895 \equiv Chaetoseris$ grandiflora (Franch.) C. Shih, nom. illeg.
- = Melanoseris taliensis (C. Shih) N. Kilian & Z. H. Wang, **syn. nov.** ≡ Chaetoseris taliensis C. Shih
- (*13) *Melanoseris leiolepis* (C. Shih) N. Kilian & J. W. Zhang in Wu & al., Fl. China 20–21: 222. 2011 ≡ *Chaetoseris leiolepis* C. Shih in Acta Phytotax. Sin. 29: 402. 1991
- (*14) *Melanoseris dolichophylla* (C. Shih) Z. H. Wang, **comb. nov.** ≡ *Chaetoseris dolichophylla* C. Shih in Acta Phytotax. Sin. 29: 401. 1991

Note: Included in the synonymy of *Melanoseris atropurpurea* by Shih & Kilian [13], herbarium work by the first author revealed it to be a separate species, consistently distinguished by the absence of a main stem, long rosette leaves and subscapose stems with 1–2 capitula only.

(*15) *Melanoseris monocephala* (C. C. Chang) Z. H. Wang, **comb. nov.** ≡ *Lactuca monocephala* C. C. Chang in Contr. Biol. Lab. Sci. Soc. China, Bot. Ser. 9: 132. 1934 ≡ *Mulgedium monocephalum* (C. C. Chang) C. Shih

Note: This fairly rare species was, with doubts, considered by Shih & Kilian [13] as conspecific with *Melanoseris lessertiana*, it is, however, unrelated and clearly distinct.

- (16) **Melanoseris graciliflora** (DC.) N. Kilian in Wu & al., Fl. China 20–21: 223. 2011 ≡ *Lactuca graciliflora* DC., Prodr. 7: 139. 1838 ≡ *Stenoseris graciliflora* (DC.) C. Shih
 - = Stenoseris taliensis (Franch.) C. Shih
- (*17) *Melanoseris tenuis* (C. Shih) N. Kilian in Wu & al., Fl. China 20–21: 223. 2011 ≡ *Stenoseris tenuis* C. Shih in Acta Phytotax. Sin. 29: 412. 1991

Excluded species:

Melanoseris oligolepis (C. C. Chang ex C. Shih) N. Kilian in Wu & al., Fl. China 20–21: 224. 2011 ≡ **Paraprenanthes oligolepis** (C. C. Chang ex C. Shih) Z. H. Wang

Melanoseris triflora (C. C. Chang & C. Shih) N. Kilian in Wu & al., Fl. China 20–21: 223. 2011 ≡ **Paraprenanthes triflora** (Chang & C. Shih) Z. H. Wang & N. Kilian

Note: See under Paraprenanthes.

Melanoseris rhombiformis (C. Shih) N. Kilian & Z. H. Wang in Wu & al., Fl. China 20–21: 219. 2011 ≡ Chaetoseris rhombiformis C. Shih. = "Cicerbita" roborowskii

Note: Analysis of the type of the name *Chaetoseris rhombiformis* by the first author made it evident that this yellow-flowered species is actually referable to "*Cicerbita*" roborowskii. This appears surprising because the latter species has always been considered to be blue-flowered (with occasional white forms). However, yellow-flowered individuals that are clearly conspecific with *C. roborowskii*, as inferred from both morphological and molecular analysis, have been collected from Sichuan (*Kilian & al. 10809* at B, KUN; see also images in [27] under that species).

4. Species of uncertain status and placement. Melanoseris hirsuta (C. Shih) N. Kilian in Wu & al., Fl. China 20–21: 220. 2011 ≡ Chaetoseris hirsuta C. Shih ≡ Lactuca hirsuta Franch. 1895 [non Nutt. 1818]

Melanoseris henryi (Dunn) N. Kilian in Wu & al., Fl. China 20–21: 221. 2011 ≡ Lactuca henryi Dunn

Lactuca scandens C. C. Chang in Contr. Biol. Lab. Sci. Soc. China, Bot. Ser. 9: 133, 1934

Supporting Information

Appendix S1 Plant material used. The data are arranged in the following order: accepted taxon name in bold and synonyms used in the phylograms (Fig. 1–2) in square brackets; unique sample identifier also used in the phylograms and, in square brackets where applicable, unit ID in the GGBN data portal [91] of stored DNA sample; abbreviated voucher data (country, locality, collecting date, collectors and collecting number, herbarium code according to Thiers [26]), full data can be obtained from the specimen labels; EMBL/Genbank/DDBJ accession numbers in the following sequence: ITS, petD, psbA-tmH, 5'tmL^(UAA)-tmF, rpl32-tmL^(UAG), tmQ^(UUG)-5'rps16. In the few cases, where already published sequences were used, only the EMBL/Genbank/DDBJ accession number preceded by an asterisk is given. (PDF)

Appendix S2 Positions of mutational hotspots (= HS) and exons in the individual chloroplast marker sequences excluded from phylogenetic analysis. The position within each marker sequence is calculated without gap; a dash denotes the absence of this sequence portion in the corresponding samples. (PDF)

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Appendix S3 Indels coded in the phylogenetic analysis.

For each marker, position, length [nt] and description of the coded indels are given according to the sequences alignment matrix. (PDF)

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Author Contributions

Conceived and designed the experiments: HP NK. Analyzed the data: ZHW HP NK. Contributed reagents/materials/analysis tools: ZHW HP NK. Wrote the paper: ZHW NK. Generated the data: ZHW NK.

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Appendix S1. Plant material used. The data are arranged in the following order: accepted taxon name in bold and synonyms used in the phylograms (Fig. 1–2) in square brackets; unique sample identifier also used in the phylograms and, in square brackets where applicable, unit ID in the GGBN data portal [91] of stored DNA sample; abbreviated voucher data (country, locality, collecting date, collectors and collecting number, herbarium code according to Thiers [26]), full data can be obtained from the specimen labels; EMBL/Genbank/DDBJ accession numbers in the following sequence: ITS, *petD*, *psbA-trnH*, *5'trnL* (UAA)-trnF, rpl32-trnL (UAG), trnQ (UUG)-5'rps16. In the few cases, where already published sequences were used, only the EMBL/Genbank/DDBJ accession number preceded by an asterisk is given.

Outgroups: Crepis multicaulis Ledeb.: LAC-003: Russia, S Siberia, Altay Mts, 22 Jun 2002, E. von Raab-Straube 020302 (B), KF485539, KF485665, KF485794, KF486050, KF485922, KF486178. — Launaea sarmentosa (Willd.) Kuntze: LAC-001: Sri Lanka, S coast, Dikwella, 25 Dec 2000, N. Kilian 7001 (B), KF485537, KF485663, KF485792, KF486048, KF485920, KF486176. — Leontodon tuberosus L.: LAC-002 [DB4947]: Greece, Insel Rhodos, Salakos, 23 Mar 2009, D. Lauterbach, T. Böhmer & A. Rolf RH2-182 (B), KF485538, KF485664, KF485793, KF486049, KF485921, KF486177. — Soroseris erysimoides (Hand.-Mazz.) C. Shih: LAC-004: China, Sichuan, Ganzi, 2 Aug 2009, Y. S. Chen & Z. H. Wang 9270 (KUN), KF485540, KF485666, KF485795, KF486051, KF485923, KF486179.

Faberia faberi (Hemsley) N. Kilian [= *Prenanthes faberi* Hemsley]: LAC-008: China, Chongqing, Nanchuan, 18 Sep 2010, *Z. H. Wang & L. Chen 245* (KUN), KF485544, KF485670, KF485799, KF486055, KF485927, KF486183; LAC-009: China, Yunnan, Qiaojia, 21 Jul 2009, *Y. S. Chen & Z. H. Wang 9034* (KUN), KF485545, KF485671, KF485800, KF486056, KF485928, KF486184; LAC-010: China, Guizhou, Hezhang, 1 Sep 2012, *Z. H. Wang & L. Chen 485* (KUN), KF485546, KF485672, KF485801, KF486057, KF485929, KF486185. *F. nanchuanensis* C. Shih [= *Faberiopsis nanchuanensis* (C. Shih) C. Shih & Y. L. Chen]: LAC-005: China, Chongqing, Nanchuan, 14 Jul 2010, *Z. H. Wang & L. Chen 131* (KUN), KF485541, KF485667, KF485796, KF486052, KF485924, KF486180. *F. sinensis* Hemsley: LAC-006 [DB3264]: China, Sichuan, Emei, 6 Jul 1997, *C. H. Li 590* (MO 04513521), KF485542, KF485668, KF485797, KF486053, KF485925, KF486181; LAC-007: China, Sichuan, Emei, 8 Jul 2010, *Z. H. Wang & L. Chen 63* (KUN), KF485543, KF485669, KF485798, KF486054, KF485926, KF486182. — *Prenanthes purpurea* L.: LAC-013: cult. Bot. Garten Berlin-Dahlem, Acc. 211-48-74-80, 19 Aug 2010, *M. Cubr 47381* (B), KF485548, KF485675, KF485804, KF486059, KF485931, KF486187.

Lactucinae core group: Cicerbita alpina (L.) Wallr.: LAC-012: *AJ633340; LAC-133 [DB7934]: Norway, Hordaland, Hardangarvidda, 15 Aug 2008, T. Dürbye 4350 (B), —, KF485674, KF485803, KF486058, KF485930, KF486186. C. hispida (DC.) Beauverd [= Cephalorrhynchus hispidus (DC.) Boiss.]: LAC-011 [DB0050]: Armenia, Vayotsdzor, Vajk, 24 Jun 2002, C. Oberprieler 10145 (B), KF485547, KF485673, KF485802, —, —, —. —. Cicerbita II: C. auriculiformis (C. Shih) N. Kilian [= Stenoseris auriculiformis C. Shih]: LAC-

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Dong & al. 634 (KUN), KF485625, KF485752, KF485880, KF486136, KF486008, KF486264; LAC-086: China, Yunnan, Lijiang, 17 Sep 2011, H. J. Dong & al. 494 (KUN), KF485620, KF485747, KF485875, KF486131, KF486003, KF486259; LAC-092: China, Yunnan, Lijiang, 1 Sep 2012, H. Peng & al. 2012-368 (KUN), KF485626, KF485753, KF485881, KF486137, KF486009, KF486265; LAC-081: China, Yunnan, Lanping, 3 Oct 2009, H. Peng & al. 715 (KUN), KF485615, KF485742, KF485870, KF486126, KF485998, KF486254; LAC-082: China, Xizang, Bomi, 21 Sep 2012, G. X. Hu & al. 1209094 (KUN), KF485616, KF485743, KF485871, KF486127, KF485999, KF486255. M. cyanea hybrid [= Chaetoseris cyanea hybrid]: LAC-093: China, Yunnan, Eryuan, 22 Sep 2011, H. J. Dong & al. 643 (KUN), KF485627, KF485754, KF485882, KF486138, KF486010, KF486266; LAC-094; China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 479 (KUN), KF485628, KF485755, KF485883, KF486139, KF486011, KF486267; LAC-095: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 481 (KUN), KF485629, KF485756, KF485884, KF486140, KF486012, KF486268; LAC-096: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 480 1 (KUN), KF485630, KF485757, KF485885, KF486141, KF486013, KF486269; LAC-097: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 480 2 (KUN), KF485631, KF485758, KF485886, KF486142, KF486014, KF486270. M. graciliflora (DC.) N. Kilian [= Stenoseris graciliflora (DC.) C. Shih, S. taliensis (Franch.) C. Shih]: LAC-110: China, Sichuan, Tianquan, 8 Sep 2010, Z. H. Wang & L. Chen 159 (KUN), KF485644, KF485771, KF485899, KF486155, KF486027, KF486283; LAC-112: China, Sichuan, Luding, 8 Sep 2010, Z. H. Wang & L. Chen 188 (KUN), KF485646, KF485773, KF485901, KF486157, KF486029, KF486285; LAC-111: China, Sichuan, Luding, 8 Sep 2010, Z. H. Wang & L. Chen 186 (KUN), KF485645, KF485772, KF485900, KF486156, KF486028, KF486284. M. lessertiana (DC.) Decne. [= Mulgedium lessertianum DC.]: LAC-071: Pakistan, Nanga Parbat, 28 Jul 1993, M. Nüsser 157 (B), KF485606, KF485733, KF485861, KF486117, KF485989, KF486245. M. likiangensis (Franch.) N. Kilian & Z. H. Wang [= Chaetoseris likiangensis (Franch.) C. Shih]: LAC-098: China, Yunnan, Lijiang, 1 Sep. 2012, H. Peng & al. 2012-396 (KUN), KF485632, KF485759, KF485887, KF486143, KF486015, KF486271. M. macrantha (C. B. Clarke) N. Kilian & J. W. Zhang [= Chaetoseris macrantha (C. B. Clarke) C. Shih]: LAC-075: Bhutan, 4200 m, 2000, S. Miehe & G. Miehe 00-223-12 (herb. Miehe), KF485610, KF485737, KF485865, KF486121, KF485993, KF486249. M. macrorhiza (Royle) N. Kilian [= Cephalorrhynchus macrorhizus (Royle) Tuisl]: LAC-073: Nepal, Mustang, 2001, S. Miehe, G. Miehe & K. Koch 01-120-03 (herb. Miehe), KF485608, KF485735, KF485863, KF486119, KF485991, KF486247; LAC-074: China, Xizang, Milin, 22 Sep 2012, G. X. Hu & al. 1209101 (KUN), KF485609, KF485736, KF485864, KF486120, KF485992, KF486248. *M. qinghaica* (S. W. Liu & T. N. Ho) N. Kilian & Z. H. Wang [= Mulgedium qinghaicum S. W. Liu & T. N. Ho]: LAC-078: Bhutan, 4200-4300 m, 2001, S. Miehe & G. Miehe 00-346-01 (herb. Miehe), KF485613, KF485740, KF485868, KF486124, KF485996, KF486252. M. souliei (Franch.) N. Kilian [= Parasyncalathium souliei (Franch.) J. W. Zhang & al.]: LAC-069: China, Xizang, Bomi, 7 Aug 2009, Y. S. Chen & Z. H. Wang 9315 (KUN), KF485604, KF485731, KF485859, KF486115, KF485987, KF486243; LAC-070: China, Sichuan, Dege, 3 Aug 2009, Y. S. Chen & Z. H. Wang 9292 (KUN), KF485605, KF485732, KF485860, KF486116, KF485988, KF486244. M. sp. [= Chaetoseris sp.]: LAC-103: China, Yunnan, Zhanyi, 7 Nov 2012, Z. H. Wang & L. Chen 486 (KUN), KF485637, KF485764, KF485892, KF486148, KF486020, KF486276. *M. tenuis* (C. Shih) N. Kilian [= Stenoseris tenuis C. Shih]: LAC-104: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 482_2 (KUN), KF485638, KF485765, KF485893, KF486149, KF486021, KF486277; LAC-105: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 482_1 (KUN), KF485639, KF485766,

KF485894, KF486150, KF486022, KF486278; LAC-106: China, Yunnan, Dali, 17 Oct 2010, Z. J. Yin & al. 3275 (KUN), KF485640, KF485767, KF485895, KF486151, KF486023, KF486279; LAC-107: China, Yunnan, Dali, 10 Sep 2009, Z. J. Yin & al. 1969 (KUN), KF485641, KF485768. KF485896. KF486152. KF486024. KF486280. *M. tenuis* hybrid [= Stenoseris tenuis hybrid]: LAC-102: China, Yunnan, Dali, 17 Oct 2010, Z. J. Yin & al. 3277 (KUN), KF485636, KF485763, KF485891, KF486147, KF486019, KF486275; LAC-108: China, Yunnan, Dali, 7 Nov 2012, Z. H. Wang & L. Chen 476 1 (KUN), KF485642, KF485769, KF485897, KF486153, KF486025, KF486281; LAC-109: China, Yunnan, Dali,7 Nov 2012, Z. H. Wang & L. Chen 476 3 (KUN), KF485643, KF485770, KF485898, KF486154, KF486026, KF486282. M. violifolia (Decne.) N. Kilian [= Cicerbita sikkimensis (Hook. f.) C. Shih]: LAC-076: China, Xizang, Cuona, Mama, 19 Sep 2009, Ze-Long Nie 806 (KUN), KF485611, KF485738, KF485866, KF486122, KF485994, KF486250; LAC-077: China, Xizang, Cuona, 7 Sep 2012, G. X. Hu & al. 1209027 (KUN), KF485612, KF485739, KF485867, KF486123, KF485995, KF486251. — Notoseris henryi (Dunn) C. Shih [= N. porphyrolepis C. Shih]: LAC-056: China, Chongqing, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 255 (KUN), KF485591, KF485718, KF485846, KF486102, KF485974, KF486230; LAC-057; China, Chongging, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 249 (KUN), KF485592, KF485719, KF485847, KF486103, KF485975, KF486231; LAC-062; China, Chongging, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 259 (KUN), KF485597, KF485724, KF485852, KF486108, KF485980, KF486236; LAC-063: China, Chongqing, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 260 (KUN), KF485598, KF485725, KF485853, KF486109, KF485981, KF486237. *N. khasiana* (C. B. Clarke) N. Kilian [= N. rhombiformis C. Shih]: LAC-066: China, Yunnan, Dali, 17 Oct 2010, Z. J. Yin & al. 3263 (KUN), KF485601, KF485728, KF485856, KF486112, KF485984, KF486240; LAC-067; China, Yunnan, Midu, 17 Apr 2010, Z. H. Wang 10 (KUN), KF485602, KF485729, KF485857, KF486113, KF485985, KF486241; LAC-068: China, Yunnan, Yunlong, 23 Sep 2011, Z. H. Wang 473 (KUN), KF485603, KF485730, KF485858, KF486114, KF485986, KF486242. *N. macilenta* (Vaniot & H. Lév.) N. Kilian [= N. psilolepis C. Shih]: LAC-064: China, Chongging, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 235 (KUN), KF485599, KF485726, KF485854, KF486110, KF485982, KF486238; LAC-065: China, Chongging, Nanchuan, 17 Sep 2010, Z. H. Wang & L. Chen 243 (KUN), KF485600, KF485727, KF485855, KF486111, KF485983, KF486239. N. scandens (Hook, f.) N. Kilian [= Prenanthes scandens Hook, f.]: LAC-052: China, Yunnan, Gongshan, 11 Nov 2011, Z. H. Wang, L. Chen & Y. Tang 458 (KUN), KF485587, KF485714, KF485842, KF486098, KF485970, KF486226. N. scandens × yakoensis [= Prenanthes scandens × yakoensis]: LAC-053: China, Yunnan, Gongshan, 11 Nov 2011, Z. H. Wang, L. Chen & Y. Tang 459 (KUN), KF485588, KF485715, KF485843, KF486099, KF485971, KF486227. N. triflora (Hemsl.) C. Shih: LAC-058: China, Chongging, Nanchuan, 18 Sep 2010, Z. H. Wang & L. Chen 262 (KUN), KF485593, KF485720, KF485848, KF486104, KF485976, KF486232; LAC-059: China, Chongqing, Nanchuan, 26 Aug 2009, H. Peng & al. 574 (KUN), KF485594, KF485721, KF485849, KF486105, KF485977, KF486233; LAC-060: China, Yunnan, Qiaojia,

21 Jul 2009, Y. S. Chen & Z. H. Wang 9038 (KUN), KF485595, KF485722, KF485850, KF486106, KF485978, KF486234; LAC-061: China, Sichuan, Emei, 4 Aug 2011, E. D. Liu & al. 3028 (KUN), KF485596, KF485723, KF485851, KF486107, KF485979, KF486235. N. yakoensis (Jeffrey) N. Kilian [= Prenanthes yakoensis Jeffrey]: LAC-054: China, Yunnan, Gongshan, 10 Nov 2011, Z. H. Wang, L. Chen & Y. Tang 463 (KUN), KF485589, KF485716, KF485844, KF486100, KF485972, KF486228; LAC-055: China, Yunnan, Gongshan, 11 Nov 2011, Z. H. Wang, L. Chen & Y. Tang 457 (KUN), KF485590, KF485717, KF485845, KF486101, KF485973, KF486229. — Paraprenanthes diversifolia (Vaniot) N. Kilian [= P. gracilipes C. Shih, P. heptantha C. Shih & D. J. Liou]: LAC-041: China, Sichuan, Dayi, 24 Jun 2011, Z. H. Wang & L. Chen 403 (KUN), KF485576, KF485703, KF485831, KF486087, KF485959, KF486215; LAC-040: China, Sichuan, Dayi, 24 Jun 2011, Z. H. Wang & L. Chen 419 (KUN), KF485575, KF485702, KF485830, KF486086, KF485958, KF486214; LAC-039: China, Sichuan, Dayi, 25 Jul 2009, Y. S. Chen & Z. H. Wang 9106 (KUN), KF485574, KF485701, KF485829, KF486085, KF485957, KF486213. *P. melanantha* (Franch.) Z. H. Wang [= Notoseris melanantha (Franch.) C. Shih]: LAC-046: China, Sichuan, Mianning, 2 Aug 2011, Z. H. Wang & L. Chen 489A (KUN), KF485581, KF485708, KF485836, KF486092, KF485964, KF486220; LAC-047: China, Sichuan, Mianning, 2 Aug 2011, Z. H. Wang & L. Chen 489B (KUN), KF485582, KF485709, KF485837, KF486093, KF485965, KF486221; LAC-048: China, Sichuan, Mianning, 2 Aug 2011, Z. H. Wang & L. Chen 489C (KUN), KF485583, KF485710, KF485838, KF486094, KF485966, KF486222. P. meridionalis (C. Shih) Sennikov [= P. hastata C. Shih]: LAC-036: China, Sichuan, Emei, 8 Jul 2010, Z. H. Wang & L. Chen 57 (KUN), KF485571, KF485698, KF485826, KF486082, KF485954, KF486210. P. oligolepis (C. C. Chang ex C. Shih) Z. H. Wang [= Cicerbita oligolepis C. C. Chang ex C. Shih]: LAC-020: China, Yunnan, Dali, 17 Oct 2010, Z. J. Yin & al. 3268 (KUN), KF485555, KF485682, KF485810, KF486066, KF485938, KF486194; LAC-021: China, Yunnan, Dali, 17 Oct 2010, Z. J. Yin & al. 3264 (KUN), KF485556, KF485683, KF485811, KF486067, KF485939, KF486195; LAC-022: China, Yunnan, Dali, 14 Sep 2011, H. J. Dong & al. 416 (KUN), KF485557, KF485684, KF485812, KF486068, KF485940, KF486196; LAC-023: China, Yunnan, Yunlong, 23 Sep 2011, Z. H. Wang & L. Chen 472 (KUN), KF485558, KF485685, KF485813, KF486069, KF485941, KF486197. P. prenanthoides (Hemsl.) C. Shih [= P. glandulosissima (C. C. Chang) C. Shih, P. luchunensis C. Shih, P. polypodifolia (Franch.) C. Shih]: LAC-034: China, Sichuan, Emei, 8 Jul 2010, Z. H. Wang & L. Chen 67 (KUN), KF485569, KF485696, KF485824, KF486080, KF485952, KF486208; LAC-031: China, Sichuan, Emei, 7 Jul 2010, Z. H. Wang & L. Chen 43 (KUN), KF485566, KF485693, KF485821, KF486077, KF485949, KF486205; LAC-033: China, Sichuan, Emei, 7 Jul 2010, Z. H. Wang & L. Chen 44 (KUN), KF485568, KF485695, KF485823, KF486079, KF485951, KF486207; LAC-035; China, Sichuan, Emei, 7 Jul 2010, Z. H. Wang & L. Chen 52 (KUN), KF485570, KF485697, KF485825, KF486081, KF485953, KF486209. P. sororia (Miq.) C. Shih [= P. multiformis C. Shih, P. pilipes (Migo) C. Shih]: LAC-032: China, Sichuan, Emei, 7 Jul 2010, Z. H. Wang & L. Chen 39 (KUN), KF485567, KF485694, KF485822, KF486078,

KF485950, KF486206; LAC-037: China, Jiangxi, Jinggangshan, 22 May 2011, H. J. Dong & Z. J. Yin 126 (KUN), KF485572, KF485699, KF485827, KF486083, KF485955, KF486211; (b) LAC-038: China, Jiangxi, Dayu, 24 May 2011, H. J. Dong & Z. J. Yin 163 (KUN), KF485573, KF485700, KF485828, KF486084, KF485956, KF486212. P. triflora (Chang & C. Shih) Z. H. Wang & N. Kilian [= Stenoseris triflora Chang & C. Shih, S. leptantha C. Shih]: LAC-024: China, Yunnan, Jingdong, 27 Nov 2010, Z. H. Wang & L. Chen 276 (KUN), KF485559, KF485686, KF485814, KF486070, KF485942, KF486198; LAC-025: China, Yunnan, Jingdong, 27 Nov 2010, Z. H. Wang & L. Chen 272 (KUN), KF485560, KF485687, KF485815, KF486071, KF485943, KF486199; LAC-026: China, Yunnan, Gongshan, 10 Nov 2011, Z. H. Wang, L. Chen & Y. Tang 462 (KUN), KF485561, KF485688, KF485816, KF486072, KF485944, KF486200; LAC-027: China, Yunnan, Eryuan, 22 Sep 2011, H. J. Dong & al. 648 (KUN), KF485562, KF485689, KF485817, KF486073, KF485945, KF486201. P. umbrosa (Dunn) Sennikov [= Mulgedium umbrosum (Dunn) C. Shih, Lactuca parishii Craib]: LAC-028: Thailand, Chiang Mai, Doi Sutep, 27 Jan 2004, N. Kilian 10242 (B), KF485563, KF485690, KF485818, KF486074, KF485946, KF486202; LAC-029: China, Yunnan, Longling, 1 May 2011, Z. H. Wang wzh10 (KUN), KF485564, KF485691, KF485819, KF486075, KF485947, KF486203; LAC-030: China, Yunnan, Jingdong, 1 May 2011, Z. H. Wang wzh09 (KUN), KF485565, KF485692, KF485820, KF486076, KF485948, KF486204. *P. wilsonii* (C. C. Chang) Z. H. Wang [= Notoseris wilsonii (C. C. Chang) C. Shih]: LAC-049: China, Sichuan, Dayi, 26 Jun 2011, Z. H. Wang & L. Chen 344A (KUN), KF485584, KF485711, KF485839, KF486095, KF485967, KF486223; LAC-050: China, Sichuan, Dayi, 25 Jun 2011, Z. H. Wang & L. Chen 344B (KUN), KF485585, KF485712, KF485840, KF486096, KF485968, KF486224; LAC-051: China, Sichuan, Dayi, 25 Jun 2011, Z. H. Wang & L. Chen 344C (KUN), KF485586, KF485713, KF485841, KF486097, KF485969, KF486225. P. yunnanensis (Franch.) C. Shih [= P. longiloba Y. Ling & C. Shih, P. sagittiformis C. Shih]: LAC-042: China, Yunnan, Kunming, 17 Aug 2010, Z. H. Wang wzh01 (KUN), KF485577, KF485704, KF485832, KF486088, KF485960, KF486216; LAC-043: China, Yunnan, Kunming, 1 Jul 2010, C. L. Xiang & al. 429 (KUN), KF485578, KF485705, KF485833, KF486089, KF485961, KF486217; LAC-044; China, Yunnan, Kunming, 1 Jul 2010, C. L. Xiang & al. 432 (KUN), KF485579, KF485706, KF485834, KF486090, KF485962, KF486218; LAC-045: China, Yunnan, Lanping, 26 Jul 2010, H. Peng & al. 1006 (KUN), KF485580, KF485707, KF485835, KF486091, KF485963, KF486219

Appendix S2. Positions of mutational hotspots (= HS) and exons in the individual chloroplast marker sequences excluded from phylogenetic analysis. The position within each marker sequence is calculated without gap; a dash denotes the absence of this sequence portion in the corresponding samples.

Sample name in the tree	petD region: exon	petD region: HS1	psbA- trnH: HS1	psbA- trnH: HS2	psbA- trnH: HS3	psbA- trnH: HS4	psbA- trnH: HS5	<i>trnL-F</i> : exon	trnL-F: HS1	rpl32-trnL: HS1
Launaea sarmentosa_LAC-001	193-200		94-101		357-365		400-417	442-490		128-128
Leontodon tuberosua_LAC-002	193-200		93-99		329-342		377-394	431-479		132-132
Crepis multicaulis_LAC-003	193-200		93-99		338-345		380-387	440-488		132-132
Soroseris erysimoides_LAC-004	188-195		94-101		331-337		372-389	396-444		
Faberiopsis nanchuanensis_LAC-005	200-207		94-101		331-337		372-389	442-490		132-132
Faberia sinensis_LAC-006	200-207		94-100		330-336		371-388	442-490		132-132
Faberia sinensis_LAC-007	200-207		94-101		331-337		372-389	442-490		132-132
Prenanthes faberi_LAC-008	200-207		94-101		331-337		372-389	442-490		132-132
Prenanthes faberi_LAC-009	200-207		94-101		331-336		371-388	442-490		132-132
Prenanthes faberi_LAC-010	200-207		94-101		331-336		371-388	442-490		132-132
Cephalorrhynchus hispidus_LAC-011	193-200		94-101		327-337		372-373	missing	missing	missing
Cicerbita alpina_LAC133	193-200	546-562	94-100		326-334		369-381	442-490		132-132
Prenanthes purpurea_LAC013	193-200	552-568	94-101		331-339		374-388	442-490		132-134
Cicerbita azurea_LAC014	193-200		94-101		327-339		374-385	442-490		132-132
Cicerbita azurea_LAC015	193-200		94-101		327-337		372-389	442-490		132-132
Stenoseris auriculiformis_LAC-016	193-200		missing	missing	missing	missing	missing	442-490		132-132
Stenoseris auriculiformis_LAC-017	193-200		94-101		327-337		372-389	442-490		132-132
Chaetoseris roborowskii_LAC-018	193-200		94-101		327-339		374-391	442-490		132-132
Chaetoseris roborowskii_LAC-019	193-200		94-101		327-337		372-389	442-490		132-132
Cicerbita oligolepis_LAC-020	193-200		94-101		332-344		379-396	442-490		
Cicerbita oligolepis_LAC-021	193-200		94-101		332-344		379-396	442-490		
Cicerbita oligolepis_LAC-022	193-200	544-560	94-101		332-344		379-396	442-490		
Cicerbita oligolepis_LAC-023	193-200		94-101		332-344		379-396	442-490		
Stenoseris leptantha_LAC-024	193-200		95-103		3-5-316		351-358	442-490		
Stenoseris leptantha_LAC-025	193-200		95-103		305-316		351-358	442-490		
Stenoseris triflora_LAC-026	193-200		95-103		305-317		352-359	442-490		
Stenoseris triflora_LAC-027	193-200		95-103		305-317		352-359	442-490		
Lactuca parishii_LAC028	193-200		94-101		327-334		369-386	442-490		132-132

Sample name in the tree	petD region: exon	petD region: HS1	psbA- trnH: HS1	psbA- trnH: HS2	psbA- trnH: HS3	psbA- trnH: HS4	psbA- trnH: HS5	<i>trnL-F</i> : exon	trnL-F: HS1	rpl32-trnL: HS1
Mulgedium umbrosum_LAC-029	193-200		94-101		327-334		369-396	442-490		132-132
Mulgedium umbrosum_LAC-030	193-200		94-101		327-334		369-396	442-490		132-132
Paraprenanthes luchunensis_LAC-031	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes multiformis_LAC-032	193-200	544-560	93-98		324-332		367-384	442-490		132-132
Paraprenanthes polypodifolia_LAC-033	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes glandulosissima_LAC-034	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes prenanthoides_LAC-035	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes hastata_LAC-036	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes pilipes_LAC-037	193-200	544-560	94-101		327-336		371-401	442-490		132-132
Paraprenanthes pilipes_LAC-038	193-200	544-560	94-101		327-335		370-421	442-490		132-132
Paraprenanthes diversifolia_LAC-039	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes heptantha_LAC-040	193-200	544-560	93-98		324-334		369-386	442-490		132-132
Paraprenanthes gracilipes_LAC-041	193-200	544-560	missing	missing	missing	missing	missing	442-490		132-132
Paraprenanthes longiloba_LAC-042	193-200	544-560	94-101		327-336		371-388	442-490		99-99
Paraprenanthes sagittiformis_LAC-043	193-200	544-560	94-101		327-336		371-388	442-490		99-99
Paraprenanthes sagittiformis_LAC-044	193-200	544-560	94-101		327-336		371-388	442-490		99-99
Paraprenanthes yunnanensis_LAC-045	193-200	544-560	94-101		327-337		372-389	442-490		99-99
Notoseris melanantha_LAC-046	193-200	544-560	94-100		353-360		395-412	442-490		132-132
Notoseris melanantha_LAC-047	193-200	544-560	94-100		353-360		395-412	442-490		132-132
Notoseris melanantha_LAC-048	193-200	544-560	94-100		353-360		395-412	442-490		132-132
Notoseris gracilipes_LAC-049	193-200	544-560	93-100		336-345		380-397	442-490		132-132
Notoseris gracilipes_LAC-050	193-200	544-560	93-100		336-345		380-397	442-490		132-132
Notoseris gracilipes_LAC-051	193-200	544-560	93-100		336-345		380-397	442-490		132-132
Prenanthes scandens_LAC-052	193-200	536-552	94-101		327-339		374-391	442-490		153-153
Prenanthes scandens * yakoensis_LAC-053	193-200	536-552	94-101		327-340		375-392	442-490		153-153
Prenanthes yakoensis_LAC-054	193-200	536-552	94-101		327-340		375-392	442-490		153-153
Prenanthes yakoensis_LAC-055	193-200	536-552	94-101		327-342		377-394	442-490		153-153
Notoseris porphyrolepis_LAC-056	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris porphyrolepis_LAC-057	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris triflora_LAC-058	193-200	546-562	94-100		326-338		373-390	442-490		132-132

Sample name in the tree	petD region: exon	petD region: HS1	psbA- trnH: HS1	psbA- trnH: HS2	psbA- trnH: HS3	psbA- trnH: HS4	psbA- trnH: HS5	<i>trnL-F</i> : exon	trnL-F: HS1	rpl32-trnL: HS1
Notoseris triflora_LAC-059	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris triflora_LAC-060	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris triflora_LAC-061	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris henryi_LAC-062	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris henryi_LAC-063	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris psilolepis_LAC-064	193-200	546-562	94-100		326-336		371-388	442-490		132-132
Notoseris psilolepis_LAC-065	193-200	546-562	94-100		326-337		372-389	442-490		132-132
Notoseris rhombiformis_LAC-066	193-200	544-560	94-100		326-337		372-389	442-490		132-132
Notoseris rhombiformis_LAC-067	193-200	544-560	94-100		326-338		373-390	442-490		132-132
Notoseris rhombiformis_LAC-068	193-200	544-560	94-100		326-335		370-387	442-490		132-132
Parasyncalathium souliei_LAC-069	193-200		94-101		318-332		367-384	442-490	38-40	133-133
Parasyncalathium souliei_LAC-070	193-200		94-101		327-341		376-393	442-490	38-40	133-133
Melanoseris lessertiana_LAC071	193-200	544-560	94-100		333-343		378-408	442-490	38-40	132-132
Mulgedium bracteatum_LAC-072	193-200	549-565	94-101		327-337		372-389	442-490	42-42	132-132
Cephalorrhynchus macrorhizus_LAC-073	193-200	544-560	94-101		327-335		370-400	442-490	38-39	132-132
Cephalorrhynchus macrorhizus_LAC-074	193-200	544-560	94-101		327-335		370-390	443-491	38-40	121-121
Chaetoseris macrantha_LAC075	193-200	544-560	94-101		106-106		141-171	442-490	38-40	132-132
Cicerbita sikkimensis_LAC-076	193-200	544-560	94-100		326-335		370-390	442-490	38-39	132-132
Cicerbita sikkimensis_LAC-077	193-200	544-560	94-100		326-335		370-390	442-490	38-39	132-132
Mulgedium qinghaicum_LAC-078	193-200	544-560	94-100		333-341		376-406	442-490	38-40	132-132
Mulgedium qinghaicum_LAC-079	193-200	544-560	94-100		333-341		376-406	442-490	38-40	132-132
Chaetoseris hastata_LAC-080	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetosris yunnanensis_LAC-081	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetosris yunnanensis_LAC-082	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris cyanea_LAC-083	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetosris lutea_LAC-084	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetosris lutea_LAC-085	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sichuanensis_LAC-086	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris lyriformis_LAC-087	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris lyriformis_LAC-088	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132

Sample name in the tree	<i>petD</i> region: exon	petD region: HS1	psbA- trnH: HS1	psbA- trnH: HS2	psbA- trnH: HS3	psbA- trnH: HS4	psbA- trnH: HS5	trnL-F: exon	trnL-F: HS1	rpl32-trnL: HS1
Chaetoseris cyanea_LAC-089	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris hispida_LAC-090	193-200	544-560	94-100		326-337		372-392	442-490	38-40	132-132
Chaetoseris lyriformis_LAC-091	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sichuanensis_LAC-092	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sp2_LAC093	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sp4_LAC094	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sp5_LAC095	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sp6_LAC096	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris sp7_LAC097	193-200	544-560	94-100		326-336		371-391	442-490	38-40	132-132
Chaetoseris likiangensis_LAC-098	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Chaetoseris grandiflora_LAC-099	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Chaetoseris taliensis_LAC-100	193-200	544-560	94-101		327-336		371-401	442-490	38-40	132-132
Chaetoseris taliensis_LAC-101	193-200	544-560	94-101		327-336		371-401	442-490	38-40	132-132
Chaetoseris sp1_LAC102	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Chaetoseris sp3_LAC103	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris tenuis_LAC-104	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris tenuis_LAC-105	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris tenuis_LAC-106	193-200	544-560	94-101		327-335		370-400	442-490	38-40	
Stenoseris tenuis_LAC-107	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris sp1_LAC-108	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris sp2_LAC-109	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris graciliflora_LAC-110	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris taliensis_LAC-111	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Stenoseris graciliflora_LAC-112	193-200	544-560	94-101		327-335		370-400	442-490	38-40	132-132
Lactuca perennis_LAC334	192-199		94-101		311-320		349-356	442-490	38-39	
Lactuca undulata_LAC-114	193-200		94-102		310-321		incomplete	442-490	38-39	94-96
Lactuca undulata_LAC-115	193-200		94-101		311-322		357-374	442-490	38-39	94-96
Lactuca dissecta_LAC-116	193-200		94-101	301-308	335-347		382-389	442-490	38-40	132-132
Lactuca dolichophylla_LAC-117	193-200		94-101	301-308	335-345		380-392	442-490	38-40	132-132
Steptorhamphus tuberosus_LAC-118	193-200		94-102		328-338	355-355	374-381	442-490	38-42	132-132

Sample name in the tree	<i>petD</i> region: exon	petD region: HS1	psbA- trnH: HS1	psbA- trnH: HS2	psbA- trnH: HS3	psbA- trnH: HS4	psbA- trnH: HS5	<i>trnL-F</i> : exon	trnL-F: HS1	rpl32-trnL: HS1
Lactuca inermis_LAC-119	193-200	544-560	95-102		339-349	366-370	389-396	442-490	38-39	123-123
Pterocypsela indica_LAC-120	193-200		94-101	301-315	342-353	370-374	392-409	438-486	38-42	132-132
Pterocypsela laciniata_LAC-121	193-200		94-101	301-315	342-354	371-375	394-411	438-486	38-42	132-132
Pterocypsela formosana_LAC-122	193-200		94-101		327-342	359-362	381-398	442-490	38-42	132-132
Pterocypsela sonchus_LAC-123	193-200		94-101	301-315	342-357	374-378	397-414	442-490	38-42	132-132
Pterocypsela raddeana_LAC-124	193-200		94-101	301-315	342-355	372-375	394-411	442-490	38-42	132-132
Pterocypsela elata_LAC-125	193-200		94-101	301-315	342-355	372-376	395-412	442-490	38-42	153-153
Scariola orientalis_LAC126	193-200	544-560	94-101	300-303	338-346		380-387	442-490	38-43	132-132
Scariola viminea_LAC-135	193-200	544-560	94-101	296-299	334-344		incomplete	442-490	38-42	132-132
Mulgedium sibiricum_LAC-128	193-200		94-101		327-335		370-387	442-490	26-27	132-132
Mulgedium tataricum_LAC-129	188-195		94-101		327-340		375-392	442-490	26-30	132-132
Lactuca sativa_LAC-132	193-200	545-561	94-102	302-305	340-348		382-389	442-490	38-41	
Lactuca serriola_LAC-131	193-200	544-560	94-101	301-304	339-346		380-387	442-490	38-41	

Sample name in the tree	rpl32- trnL: HS2	rpl32- trnL: HS3	rpl32- trnL: HS4	rpl32- trnL: HS5	rpl32- trnL: HS6	rpl32- trnL: HS7	rpl32- trnL: HS8	trnQ- rps16: HS1	trnQ-rps16: HS2	trnQ- rps16: HS3
Launaea sarmentosa_LAC-001	160-167		377-387		697-697		777-777	102-103		
Leontodon tuberosua_LAC-002	164-173	190-190	406-416	738-744			839-839	102-102		
Crepis multicaulis_LAC-003		148-148	379-388		698-698			96-96		908-917
Soroseris erysimoides_LAC-004	160-167	177-177	393-401				809-809	102-102		
Faberiopsis nanchuanensis_LAC-005	161-167	177-177	393-402				830-831	102-102		
Faberia sinensis_LAC-006	161-167	177-178	394-403				831-832	102-102		
Faberia sinensis_LAC-007	161-167	177-178	394-402				830-831	102-102		
Prenanthes faberi_LAC-008	161-167	177-178	394-404				832-833	102-102		
Prenanthes faberi_LAC-009	161-167	177-178	394-403				831-832	102-102		
Prenanthes faberi_LAC-010	161-167	177-178	394-403				831-832	102-102		
Cephalorrhynchus hispidus_LAC-011	missing	missing	missing							
Cicerbita alpina_LAC133	164-174	184	400-420				831-832	102-102		

Sample name in the tree	rpl32- trnL: HS2	rpl32- trnL: HS3	rpl32- trnL: HS4	rpl32- trnL: HS5	rpl32- trnL: HS6	rpl32- trnL: HS7	rpl32- trnL: HS8	trnQ- rps16: HS1	trnQ-rps16: HS2	trnQ- rps16: HS3
Prenanthes purpurea_LAC013	166-175	185	401-411				803-804	102-102		885-891
Cicerbita azurea_LAC014	164-171	181	397-417			802-802	829-829	102-102		
Cicerbita azurea_LAC015	164-171	181	397-417			802-802	829-829	102-102		
Stenoseris auriculiformis_LAC-016	164-174	184	383-403		736-736	789-789	816-817	102-102		
Stenoseris auriculiformis_LAC-017	164-174	184	383-403		736-736	789-789	816-817	102-102		
Chaetoseris roborowskii_LAC-018	164-174	184	383-403			788-788	815-816	102-102		
Chaetoseris roborowskii_LAC-019	164-174	184	383-403			788-788	815-816	102-102		
Cicerbita oligolepis_LAC-020	163-169	179	395-415			800-801	828-829	102-102		
Cicerbita oligolepis_LAC-021	163-169	179	395-415			800-801	828-829	102-102		
Cicerbita oligolepis_LAC-022	163-169	179	395-415			800-801	828-829	102-102		
Cicerbita oligolepis_LAC-023	163-169	179	395-415			800-801	828-829	102-102		
Stenoseris leptantha_LAC-024	157-163	173	397-417			802-802	829-830	102-102		
Stenoseris leptantha_LAC-025	157-163	173	397-417			802-802	829-830	102-102		
Stenoseris triflora_LAC-026	157-163	173	397-417			802-802	829-830	102-102		
Stenoseris triflora_LAC-027	157-163	173	397-417			802-802	829-830	102-102		
Lactuca parishii_LAC028	164-175		400-420	763-770		836-837	864-864	102-102		
Mulgedium umbrosum_LAC-029	164-175		400-420	740-747		813-813	840-840	102-102		
Mulgedium umbrosum_LAC-030	164-175		400-420	740-747		813-814	841-841	102-102		
Paraprenanthes luchunensis_LAC-031	164-165	175	391-411	731-738	752-752	793-793	820-820	102-102		
Paraprenanthes multiformis_LAC-032	164-165	175	391-411	731-738	752-752	793-793	820-820	102-102		
Paraprenanthes polypodifolia_LAC-033	164-165	175	391-411	731-738	752-752	796-796	823-823	102-102		
Paraprenanthes glandulosissima_LAC-034	164-165	175	391-411	731-738	752-752	793-793	820-820	102-102		
Paraprenanthes prenanthoides_LAC-035	164-165	175	391-411	731-738	752-752	793-793	820-820	102-102		
Paraprenanthes hastata_LAC-036	164-171	181	397-417	737-744	758-758	799-799	826-826	102-102		
Paraprenanthes pilipes_LAC-037	164-175	185	401-421	741-748		814-814	841-841	102-102		
Paraprenanthes pilipes_LAC-038	164-175	185	401-421	741-748		814-814	841-841	102-102		
Paraprenanthes diversifolia_LAC-039	164-173	183	399-419	739-746	760-760	801-801	828-828	102-102		
Paraprenanthes heptantha_LAC-040	164-173	183	399-419	739-746	760-760	801-801	828-828	102-102		
Paraprenanthes gracilipes_LAC-041	164-172	182	398-418	738-745	759-759	800-800	827-827	102-102		
Paraprenanthes longiloba_LAC-042	131-138	148	364-384	704-711		761-761	788-788	102-102		

Sample name in the tree	rpl32- trnL: HS2	rpl32- trnL: HS3	rpl32- trnL: HS4	rpl32- trnL: HS5	rpl32- trnL: HS6	rpl32- trnL: HS7	rpl32- trnL: HS8	trnQ- rps16: HS1	trnQ-rps16: HS2	trnQ- rps16: HS3
Paraprenanthes sagittiformis_LAC-043	131-139	149	365-385	705-712		762-762	789-789	102-102		
Paraprenanthes sagittiformis_LAC-044	131-139	149	365-385	705-712		762-762	789-789	102-102		
Paraprenanthes yunnanensis_LAC-045	131-139	149	365-385	705-712		762-762	789-789	102-102		
Notoseris melanantha_LAC-046	164-172	182	398-418	738-745		811-811	838-838	102-102		
Notoseris melanantha_LAC-047	164-172	182	398-418	738-745		811-811	838-838	102-102		
Notoseris melanantha_LAC-048	164-172	182	398-418	738-745		811-811	838-838	102-102		
Notoseris gracilipes_LAC-049	164-172	182	398-418	738-745	759-759	800-800	827-827	102-102		
Notoseris gracilipes_LAC-050	164-172	182	398-418	738-745	759-759	800-800	827-827	102-102		
Notoseris gracilipes_LAC-051	164-172	182	398-418	738-745	759-759	800-800	827-827	102-102		
Prenanthes scandens_LAC-052	185-195	206	422-432			817-817	844-845	102-102		
Prenanthes scandens * yakoensis_LAC-053	185-194	205	421-431			816-816	843-844	102-102		
Prenanthes yakoensis_LAC-054	185-194	205	421-431			816-816	843-844	102-102		
Prenanthes yakoensis_LAC-055	185-193	204	420-430			815-815	842-843	102-102		
Notoseris porphyrolepis_LAC-056	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris porphyrolepis_LAC-057	164-174	184	400-420	745-752		818-818	845-845	102-102		
Notoseris triflora_LAC-058	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris triflora_LAC-059	164-174	184	400-420	745-752		818-818	845-845	102-102		
Notoseris triflora_LAC-060	164-174	184	400-420	745-752		818-819	846-846	102-102		
Notoseris triflora_LAC-061	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris henryi_LAC-062	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris henryi_LAC-063	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris psilolepis_LAC-064	164-175	185	401-421	746-753		819-819	846-846	102-102		
Notoseris psilolepis_LAC-065	164-175	185	401-421			811-811	838-838	102-102		
Notoseris rhombiformis_LAC-066	164-175	185	401-421	741-748		814-814	841-841	102-102		
Notoseris rhombiformis_LAC-067	164-175	185	401-421	741-748		814-814	841-841	102-102		
Notoseris rhombiformis_LAC-068	164-177	187	403-423	748-755		821-821	848-848	102-102		
Parasyncalathium souliei_LAC-069	165-178	192	408-418		748-748	801-801	828-828	102-102		
Parasyncalathium souliei_LAC-070	165-177	191	407-417		747-747	800-800	827-827	102-102		
Melanoseris lessertiana_LAC071	164-174	184	400-420		753-753	806-806	833-833	102-102		
Mulgedium bracteatum_LAC-072		148	364-384			776-776	803-803	102-102		

Sample name in the tree	rpl32- trnL: HS2	rpl32- trnL: HS3	rpl32- trnL: HS4	rpl32- trnL: HS5	rpl32- trnL: HS6	rpl32- trnL: HS7	rpl32- trnL: HS8	trnQ- rps16: HS1	trnQ-rps16: HS2	trnQ- rps16: HS3
Cephalorrhynchus macrorhizus_LAC-073	164-172	182	398-408			800-800	827-827	102-102	746-751	
Cephalorrhynchus macrorhizus_LAC-074	153-160	170	386-396			781-781	808-808	102-102		890-900
Chaetoseris macrantha_LAC075	164-174	184	400-420		753-753		832-832	102-102		890-900
Cicerbita sikkimensis_LAC-076	164-174	184	400-420		753-753			102-102		890-900
Cicerbita sikkimensis_LAC-077	164-174	184	400-420		753-753		826-826	102-102		890-900
Mulgedium qinghaicum_LAC-078	164-174	184	400-420		753-753	806-806	833-833	102-102		890-900
Mulgedium qinghaicum_LAC-079	164-173	183	399-419		752-752	805-805	832-832	102-102		890-900
Chaetoseris hastata_LAC-080	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetosris yunnanensis_LAC-081	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetosris yunnanensis_LAC-082	164-174	184	400-420		753-753		826-826			888-898
Chaetoseris cyanea_LAC-083	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetosris lutea_LAC-084	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetosris lutea_LAC-085	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sichuanensis_LAC-086	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris lyriformis_LAC-087	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris lyriformis_LAC-088	164-171	181	397-417		750-750		823-823	102-102		890-900
Chaetoseris cyanea_LAC-089	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris hispida_LAC-090	164-172	182	398-418		751-751		824-824	102-102	746-751	896-906
Chaetoseris lyriformis_LAC-091	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sichuanensis_LAC-092	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sp2_LAC093	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sp4_LAC094	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sp5_LAC095	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sp6_LAC096	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris sp7_LAC097	164-172	182	398-418		751-751		824-824	102-102		890-900
Chaetoseris likiangensis_LAC-098	164-173	183	398-419		758-758	811-811		102-102		890-900
Chaetoseris grandiflora_LAC-099	164-173	183	398-419		758-758	811-811		102-102		890-900
Chaetoseris taliensis_LAC-100	164-173	183	398-419		758-758	811-811		102-102		895-905
Chaetoseris taliensis_LAC-101	164-173	183	398-419		758-758	811-811		102-102		890-900
Chaetoseris sp1_LAC102	164-173	183	398-419		758-758	811-811		102-102		890-900

Sample name in the tree	rpl32- trnL: HS2	rpl32- trnL: HS3	rpl32- trnL: HS4	rpl32- trnL: HS5	rpl32- trnL: HS6	rpl32- trnL: HS7	rpl32- trnL: HS8	trnQ- rps16: HS1	trnQ-rps16: HS2	trnQ- rps16: HS3
Chaetoseris sp3_LAC103	164-173	183	398-419		758-758	811-811		102-102		
Stenoseris tenuis_LAC-104	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris tenuis_LAC-105	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris tenuis_LAC-106	163-171	181	397-417		756-756	809-809		102-102		890-900
Stenoseris tenuis_LAC-107	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris sp1_LAC-108	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris sp2_LAC-109	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris graciliflora_LAC-110	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris taliensis_LAC-111	164-173	183	398-419		758-758	811-811		102-102		890-900
Stenoseris graciliflora_LAC-112	164-173	183	398-419		758-758	811-811		102-102		890-900
Lactuca perennis_LAC334	94-104	114	323-333			746-747	774-774	102-102		
Lactuca undulata_LAC-114	128-136	146	355-375			760-760	787-787	102-102		
Lactuca undulata_LAC-115	128-136	146	355-375			760-760	787-787	102-102		
Lactuca dissecta_LAC-116	164-178	188	403-423			822-822	849-849	102-102		
Lactuca dolichophylla_LAC-117	164-174	184	398-418			817-817	844-844	102-102		
Steptorhamphus tuberosus_LAC-118	164-174	184	400-420			819-819	846-846	102-102		
Lactuca inermis_LAC-119	155-165	175	402-412			766-766	800-800	102-102		
Pterocypsela indica_LAC-120	164-173	183	399-419			805-805	832-832	102-102	746-751	
Pterocypsela laciniata_LAC-121	164-173	183	399-419			805-805	832-832	102-102	746-751	
Pterocypsela formosana_LAC-122	164-173	183	399-419			805-805	832-832	102-102	746-751	
Pterocypsela sonchus_LAC-123	164-173	183	399-419			805-805	832-832	102-102	746-751	
Pterocypsela raddeana_LAC-124	164-173	183	399-419			805-805	832-832	102-102		
Pterocypsela elata_LAC-125	185-193	203	419-439			825-825	852-852	102-102	746-751	
Scariola orientalis_LAC126	164-175	185	401-411			799-799	826-826	104-105		913-923
Scariola viminea_LAC-135	164-174	184	400-410			799-799	826-826	104-106		898-908
Mulgedium sibiricum_LAC-128	164-173	183	400-420			805-805	832-832	102-102		
Mulgedium tataricum_LAC-129	160-172	182	400-421			806-806	833-833	102-102		
Lactuca sativa_LAC-132			319-339			728-728	755-755	102-104		
Lactuca serriola_LAC-131			319-339			745-745	772-772	102-104		

Table S2. Indels coded in the phylogenetic analysis. For each marker, position, length [nt] and description of the coded indels are given according to the sequences alignment matrix.

ITS region

No.	Position	Length (nt)	Description
1	4-4	1	Gap in Faberia sinensis, Prenanthes yakoensis
2	16-16	1	Insertion of "A" in Scariola orientalis, gap in all other taxa
3	40-40	1	Insertion of "C" in <i>Prenanthes yakoensis</i> , gap in all other taxa
4	40-41	2	Gap in Launaea sarmentosa
5	43-43	1	Gap in Prenanthes yakoensis
6	49-49	1	Gap in Crepis multicaulis
7	59-59	1	Insertion of "T" in Cicerbita oligolepis and Stenoseris triflora, Stenoseris leptantha, gap in all other taxa
8	59-60	2	Gap in Lactuca perennis
9	83-83	1	Gap in Soroseris erysimoides and Crepis multicaulis
10	85-85	1	Gap in Soroseris erysimoides and Prenanthes purpurea
11	87-87	1	Gap in Prenanthes purpurea
12	89-89	1	Gap in Prenanthes purpurea, Cicerbita oligolepis
13	91-91	1	Gap in Lactuca inermis, all Pterocypsela except P. raddeana, Scariola orientalis, S. viminea, Lagedium sibiricum, Mulgedium tataricum, L. sativa, L. serriola
14	94-94	1	Insertion of "T(C)" in all Pterocypsela, Scariola orientalis, S. viminea, Lagedium sibiricum, Mulgedium tataricum, L. sativa, L. serriola, gap in all other taxa
15	102-103	2	Gap in Scariola orientalis and S. viminea
16	103-103	1	Insertion of "T" in Parasyncalathium souliei, gap in all other taxa
17	112-112	1	Gap in Melanoseris lessertiana
18	118-119	2	Gap in Launaea sarmentosa, Crepis multicaulis, Soroseris erysimoides
19	119-119	1	Insertion of "T" in Faberia sinensis, gap in all other taxa
20	119-120	2	Gap in Lactuca sativa and L. serriola
21	122-122	1	Gap in Prenanthes purpurea
22	124-124	1	Gap in Leontodon tuberosus
23	128-129	2	Gap in Parasyncalathium souliei, Lactuc perennis, L. undulata, L. dissecta, L. dolichophylla, L. inermis, Steptorhamphus tuberosus
24	129-129	1	Insertion of "G" in Cicerbita azurea and Pterocypsela indica, gap in all other taxa
25	129-130	2	Gap in Launaea sarmentosa and Leontodon tuberosus
26	133-134	2	Insertion of "AG" in Crepis multicaulis, gap in all other taxa
27	141-142	2	Gap in all Lactuca clade
28	151-151	1	Insertion of "G" in Crepis multicaulis, gap in all other taxa
29	154-156	3	Gap in Faberiopsis nanchuanensis
30	165-165	1	Insertion of "A" in Leontodon tuberosus, Scariola orientalis, S. viminea, Lagedium sibiricum, Mulgeium tataricum, L. sativa, L. serriola, gap in all other taxa

ITS region

No.	Position	Length (nt)	Description
31	198-198	1	Gap in Prenanthes faberi, Cephalorrhynchus hispidus, Melanoseris lessertiana, all Lactuca clade except L. perennis and L. undulata
32	201-201	1	Insertion of "A" in <i>Prenanthes purpurea</i> , gap in all other taxa
33	227-236	9	Gap in one of Faberia sinensis
34	232-232	1	Insertion of "T" in Launaea sarmentosa
35	246-246	1	Gap in Lactuca parshii and Mulgedium umbrosum
36	249-249	1	Gap in Lactuca perennis
37	257-257	1	Insertion of "A" in Cephalorrhynchus hispidus, gap in all other taxa
38	443-443	1	Insertion of "C" in Lagedium sibiricum and Mulgedium tataricum, gap in all other taxa
39	449-449	1	Gap in Lagedium sibiricum and Mulgedium tataricum
40	449-450	2	Gap in Parasyncalathium souliei
41	459-460	2	Insertion of "AT" in Leontodon tuberosus and "CT" in Crepis multicaulis, gap in all other taxa
42	460-460	1	Gap in Launaea sarmentosa
43	463-463	1	Insertion of "T" in Leontodon tuberosus and "C" in Lactuca inermis, gap in all other taxa
44	463-465	3	Gap in Crepis multicaulis
45	465-465	1	Gap in Soroseris erysimoides
46	468-468	1	Gap in Notoseris triflora
47	472-476	5	Gap in Lactuca inermis
48	477-477	1	Gap in Launaea sarmentosa and Prenanthes purpurea
49	478-478	1	Insertion of "T(C, A)" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Prenanthes purpurea, Soroseris erysimoides, all Faberia, gap in all other taxa
50	483-485	3	Gap in all Faberia
51	484-484	1	Insertion of "T" in Soroseris erysimoides and "C" in Crepis multicaulis, gap in all other taxa
52	485-485	1	Insertion of "A(T)" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides and Prenanthes purpurea, gap in all other taxa
53	487-487	1	Gap in Cicerbita alpina
54	515-516	2	Gap in Notoseris rhombiformis
55	516-517	2	Gap in Leontodon tuberosus, Cephalorrhynchus hispidus and Cicerbita alpina
56	540-540	1	Gap in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides, all Faberia, Prenanthes purpurea, Cephalorrhynchus hispidus, Cicerbita alpina and one of Stenoseris auriculiformis
57	544-544	1	Insertion of "C" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides, all Faberia, Prenanthes purpurea, Cephalorrhynchus hispidus and Cicerbita clade, gap in all other taxa
58	590-590	1	Gap in Leontodon tuberosus
59	590-591	2	Gap in Cicerbita oligolepis

ITS region

No.	Position	Length (nt)	Description
60	598-604	7	Gap in Lactuca inermis
61	619-619	1	Insertion of "A" in Faberiopsis nanchuanensis, gap in all other taxa
62	619-620	2	Gap in Launaea sarmentosa and Cephalorrhynchus hispidus
63	633-633	1	Gap in Launaea sarmentosa
64	637-637	1	Insertion of "C(T)" in Notoseris wilsonii and N melanantha, gap in all other taxa
65	640-640	1	Insertion of "A" in one of Notoseris melanantha, gap in all other taxa
66	650-651	2	Gap in Cicerbita sikkimensis

petD region

No.	Position	Length (nt)	Description
1	6-11	6	Inserton of "TATAGA" in Soroseris erysimoides and all Faberia (replicate), gap in all other taxa
2	27-31	5	Gap in Mulgedium tataricum
3	152-162	11	Gap in Soroseris erysimoides
4	162-162	1	Gap in all Faberia, insertion of "A" in all other taxa
5	166-167	2	Insertion of "TA" (replicate) in all Faberia, gap in all other taxa
6	182-182	1	Gap in Lactuca perennis, insertion of "T" in all other taxa
7	256-256	1	Insertion of "G" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides, all Faberia and Prenanthes purpurea, gap in all other taxa
8	349-353	5	Gap in Cephalorrhynchus hispidus and Cicerbita alpina, insertion of "AGATA" in all other taxa
9	418-418	1	Insertion of "T" in Lactuca sativa, gap in all other taxa
10	512-512	1	Insertion of "C" in Parasyncalathium souliei, gap in all other taxa
11	550-554	5	Insertion of "TTTAT" (replicate) in Mulgedium bracteatum, gap in all other taxa
12	561-575	15	Gap in Prenanthes yakoensis and P. scandens
13	565-571	7	Insertion of "TAAAAAT" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides, all Faberia, Prenanthes purpurea, Cephalorrynchus tuberosus, Cicerbita alpina, gap in all other taxa
14	624-625	2	Insertion of "TC" (replicate) in Notoseris henryi, N. triflora, N. psilolepis, gap in all other taxa
15	765-770	6	Gap in all Faberia
16	776-779	4	Insetion of "TATA" (replicate) in one small clade of <i>Melanoseris</i> , i.e. <i>Chaetoseris likiangensis</i> , <i>Ch. grandiflora</i> , <i>Stenoseris tenuis</i> , <i>S. graciliflora</i> , including hybrids between them
17	801-801	1	Insertion of "A" in Mulgedium bracteatum, gap in all other taxa
18	819-819	1	Insertion of "T" in Launaea sarmentosa, gap in all other taxa
19	819-832	14	Insertion of "TTTATATGGATTCA" (replicate) in Launaea sarmentosa
20	915-915	1	Gap in Lactuca perennis and Scariola viminea, insertion of "T" in all other taxa

psbA-trnH

No.	Position	Length (nt)	Description
1	4-4	1	Insertion of "C" in Lactuca inermis, gap in all other taxa
2	25-25	1	Insertion of "A" in Stenoseris triflora and S. leptantha, gap in all other taxa
3	48-48	1	Gap in Leontodon tuberosus, Crepis multicaulis, insertion of "A" in all other taxa
4	83-83	1	Gap in Paraprenanthes diversifolia, P. prenanthoies, P. hastata
5	96-105	10	Insertion of "ATTTTTTT" in Notoseris wilsonii, gap in all other taxa
6	96-117	22	Gap in all taxa except Notoseris wilsonii, Lactuca inermis
7	106-117	12	Insertion of "ATTTTTTTT" in Lactuca inermis, gap in all other taxa
8	122-127	6	Gap in Stenseris triflora and S. leptantha
9	122-430	309	Gap in Chaetoseris macrantha
10	155-155	1	Gap in Launaea sarmentosa
11	161-175	15	Gap in one of Parasyncalathium souliei
12	169-174	6	Gap in Cephalorrhynchus hispidus, Cicerbita alpina
13	169-175	7	Gap in Lactuca inermis
14	170-171	2	Insertion of "TT" in Leontodon tuberosus
15	195-210	16	Insertion of "TTTCTCTTTGTATAAA" in Launaea sarmentosa, gap in all other taxa
16	213-213	1	Gap in Scariola orientalis and S. viminea
17	252-252	1	Gap in Leontodon tuberosus
18	263-269	7	Insertion of "GTTTTAT" in Melanoseris lessertiana and Mulgedium qinghaicum, gap in all other taxa
19	263-273	11	Gap in Scariola viminea
20	314-325	12	Gap in Lactuca perennis and L. undulata
21	332-341	10	Insertion of "ATTTTATTAT" in Launaea sarmentosa, gap in all other taxa
22	332-350	19	Gap in all taxa except Launaea sarmentosa, Crepis multicaulis
23	342-350	9	Insertion of "TTTACATTT" in Crepis multicaulis, gap in all other taxa
24	356-386	31	Gap in Stenseris triflora and S. leptantha
25	365-369	5	Insertion of "ATTTT" in Cicerbita oligolepis, gap in all other taxa
26	374-381	8	Insertion of "GAAAG(T)AAA" in Scariola orientalis, S. viminea, Lactuca sativa and L. serriola
27	374-385	12	Gap in Lactuca perennis and L. undulata
28	390-416	27	Insertion of "TATTACTTTGATTTCATAAATAAGAAA" in Notoseris melanantha, gap in all other taxa
29	425-426	2	Gap in one of Lactuca undulata
30	431-431	1	Gap in Scariola orientalis and S. viminea
31	431-436	6	Gap in Lactuca perennis
32	432-432	1	Gap in Lactuca sativa and L. serriola
33	454-454	1	Gap in Pterocypsela indica

trnL-F

No.	Position	Length (nt)	Description
1	110-119	10	Gap in Soroseris erysimoides
2	182-218	37	Gap in Soroseris erysimoides
3	191-191	1	Insertion of "T" in one of Cephalorrhynchus macrorhizus, gap in all other taxa
4	191-196	6	Gap in Leontodon tuberosus
5	217-222	6	Gap in Leontodon tuberosus
6	248-251	4	Gap in Pterocypsela indica and P. laciniata
7	263-266	4	Insertion of "CATA" (replicate) in Crepis multicaulis, gap in all other taxa
8	290-290	1	Gap in Crepis multicaulis
9	292-295	4	Gap in Crepis multicaulis
10	308-308	1	Gap in Crepis multicaulis
11	472-487	16	Gap in Lagedium sibiricum and Mulgedium tataricum
12	484-487	4	Insertion of "TTTA" in Mulgedium bracteatum, gap in all other taxa
13	484-507	24	Gap in Launaea sarmentosa
14	484-508	25	Gap in Crepis multicaulis
15	494-506	13	Gap in Scariola orientalis
16	500-506	7	Insertion of "CTTTATC" in Prenanthes scandens and P. yakoensis
17	508-508	1	Gap in Soroseris erysimoides, insertion of "T" in all other taxa
18	570-571	2	Gap in all Pterocypsela
19	575-577	3	Insertion of "TAA" in Crepis multicaulis
20	575-588	14	Gap in Cicerbita azurea
21	575-589	15	Gap in all but Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Soroseris erysimoides, all Faberia, Prenanthes purpurea, Cicerbita alpina, C. azurea
22	596-597	2	Insertion of "TT" in Mulgedium bracteatum, gap in all other taxa
23	609-612	4	Gap in Lactuca perennis
24	610-611	2	Insertion of "AT" (SSR) in Lactuca inermis, gap in all other taxa
25	610-619	10	Gap in Crepis multicaulis, Soroseris erysimoides and all Faberia
26	643-647	5	Insertion of "GGAAT" (replicate) in Lagedium sibiricum and Mulgedium tataricum, gap in all other taxa
27	667-674	8	Gap in Crepis multicaulis
28	743-747	5	Gap in Scariola orientalis
29	789-794	6	Gap in one of Notoseris rhombiformis, insertion of "ATGAAA" in all other taxa
30	793-798	6	Gap in Crepis multicaulis
31	808-811	4	Gap in all Pterocypsela, insertion of "ATGA" in all other taxa
32	813-817	5	Insertion of "GAATA" (replicate) in Lactuca perennis and L. undulata, gap in all other taxa

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No.	Position	Length (nt)	Description
1	79-84	6	Gap in Crepis multicaulis
2	94-95	2	Insertion of "AT" (replicate) in Scariola orientalis and S. viminea, gap in all other taxa
3	115-130	16	Insertion of "TTTTTCTTAGCTACAT" (replicate) in <i>Scariola orientalis</i> , gap in all other taxa
4	139-152	14	Insertion of "AAACATAGATAATC" in Soroseris erysimoides, gap in all other taxa
5	161-161	1	Insertion of "T" in <i>Cicerbita</i> clade, <i>Melanoseris</i> clade and <i>Lactuca</i> clade, gap in other clade
6	161-162	2	Gap in Paraprenanthes longiloba
7	196-196	1	Gap in <i>Crepis multicaulis</i> , <i>Lactuca sativa</i> and <i>L. serriola</i> , insertion of "T" in all other taxa
8	209-215	7	Gap in Lagedium sibiricum
9	240-240	, 1	Gap in Crepis multicaulis
10	265-269	5	Insertion of "AAAAT" in all <i>Faberia</i> , gap in all other taxa
11	287-304	18	Gap in Lactuca perennis
12	317-317	10	Insertion of "A" in <i>Cicerbita alpina</i> , gap in all other taxa
13	332-339	8	Insertion of "TTGACTCC" in <i>Crepis multicaulis</i>
14	354-362	9	Gap in Prenanthes purpurea
15	359-362	4	Insertion of "TCAA" in Scariola orientalis, S. viminea, Lactuca sativa and L. serriola, gap in all other taxa
16	456-465	10	Insertion of "ATCATTATTT" (replicate) in <i>Soroseris erysimoides</i> , gap in all other taxa
17	472-476	5	Insertion of "ATTTG" (replicate) in one of <i>Chaetoseris taliensis</i> , gap in all other taxa
18	495-502	8	Gap in Launaea sarmentosa
19	495-503	9	Insertion of "TTTTACAGT" (replicate) in Soroseris erysimoides, gap in all other taxa
20	522-559	38	Gap in all but Mulgedium umbrosum and Lactuca dissecta
21	547-559	13	Gap in Lactuca dissecta
22	571-578	8	Insertion of "ATAAGATC" in Lactuca perennis, gap in all other taxa
23	571-586	16	Gap in all but Lactuca perennis, Notoseris henryi, N. triflora, N. psilolepis
24	579-586	8	Insertion of "ATAAGAAA" in <i>Notoseris henryi</i> , <i>N. triflora</i> , <i>N. psilolepis</i> , gap in all other taxa
25	614-617	4	Insertion of "TCGA" in all <i>Paraprenanthes</i> but <i>P. pilipes</i> , <i>Notoseris melanantha</i> , gap in all other taxa
26	639-643	5	Insertion of "TTAAA" (replicate) in <i>Lactuca perennis</i> and <i>L. undulata</i> , gap in all other taxa
27	668-673	6	Insertion of "ATAGAT" (replicate) in Lactuca dissecta and L. dolichophylla
28	698-710	13	Insertion of "GTTAAATATTTAA" (replicate) in one of <i>Notoseris rhombiformis</i> , gap in all other taxa
29	786-786	1	Insertion of "T" in Launaea sarmentosa
30	805-836	32	Gap in all taxa but Crepis multicaulis and Soroseris erysimoides and all Faberia
31	814-832	19	Gap in Soroseris erysimoides
32	814-836	23	Gap in all taxa but Crepis multicaulis and Soroseris erysimoides

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No.	Position	Length (nt)	Description
33	839-839	1	Insertion of "T" in Prenanthes purpurea
34	845-851	7	Insertion of "CCTATAG" (replicate) in one of Notoseris wilsonii, gap in all other taxa
35	857-857	1	Gap in Soroseris erysimoides
36	867-867	1	Gap in Crepis multicaulis
37	887-891	5	Insertion of "TTTTG" (replicate) in Lactuca parshii and Mulgedium umbrosum, gap in all other taxa
38	901-904	4	Gap in Steptorhamphus tuberosus, insertion of "ACCT" in all other taxa
39	947-951	5	Gap in Lactuca inermis, insertion of "TTATC" in all other taxa
40	963-965	3	Gap in Lactuca perennis and L. undulata, insertion of "CAC" in all other taxa
41	1014- 1030	17	Gap in Crepis multicaulis
42	1024- 1028	5	Insertion of "AACTG" (replicate) in Lactuca undulata, gap in all other taxa
43	1066- 1079	14	Insertion of "AAAAAAGAAAGAAG" (replicate) in Pterocypsela indica, gap in all other taxa
44	1086- 1109	24	Gap in Leontodon tuberosus
45	1087- 1090	4	Gap in Mulgedium bracteatum
46	1097- 1099	3	Gap in Cicerbita clade II
47	1110- 1117	8	Gap in Prenanths yakoensis and P. scandens

rpl32-trnL

No.	Position	Length	Tpi32-trnL Description
		(nt)	
1	9-62	54	Gap in Paraprenanthes yunnanensis, P. saggitiformis, P. longiloba
2	42-62	21	Insertion of "ATCAATACTTTTTAATATAA" in <i>Prenanthes yakoensis</i> , <i>P. scandens</i> , gap in all other taxa
3	73-81	9	Gap in Launaea sarmentosa
4	75-80	6	Gap in Stenoseris triflora, S. leptantha
5	75-85	11	Gap in one of Cephalorrhynchus macrorhizus
6	75-134	60	Gap in Lactuca perennis, L. undulata
7	78-86	9	Gap in Lactuca inermis
8	103-123	21	Insertion of "CTATTCTAATCAATATGAACC" (replicate) in Pterocypsela elata, gap in all other taxa
9	128-128	1	Insertion of "A" in Parasyncalathium souliei, gap in all other taxa
10	153-237	85	Gap in Lactuca sativa and L. serriola
11	157-162	6	Insertion of "AGGTTT" (replicate) in Launaea sarmentosa, gap in all other taxa
12	167-167	1	Gap in Launaea sarmentosa
13	181-222	42	Gap in Lactuca perennis
14	192-227	36	Gap in Mulgedium bracteatum
15	208-222	15	Gap in Mulgedium tataricum
16	209-222	14	Gap in Soroseris erysimoides and all Faberia
17	212-213	2	Insertion of "GC" in Leontodeon tuberosus, gap in all other taxa
18	212-222	11	Gap in all taxa but Leontodon tuberosus, Parasyncalathium souliei
19	212-227	16	Gap in Launaea sarmentosa
20	214-217	4	Insertion of "ACAA' in Parasyncalathium souliei, gap in all other taxa
21	218-222	5	Insertion of "AGAAT" (replicate) in Leontodon tuberosus, gap in all other taxa
22	259-266	8	Insertion of "TTTTTTT" in Crepis multicaulis, gap in all other taxa
23	259-278	20	Gap in all taxa except Crepis multicaulis, Lactuca inermis
24	267-278	12	Insertion of "GTAGTATATTT" (replicate) in Lactuca inermis, gap in all other taxa
25	300-307	8	Insertion of "TTTTGTGG" (replicate) in Stenoseris triflora and S. leptantha, gap in all other taxa
26	300-308	9	Gap in Lactuca inermis
27	314-317	4	Insertion of "GGTG" (replicate) in Lagedium sibiricum and Mulgedium tataricaum, gap in all other taxa
28	331-331	1	Gap in Lactuca dolichophylla
29	350-351	2	Gap in Lagedium sibiricum
30	354-360	7	Gap in <i>Lactuca perennis</i> and <i>L. undulata</i>
31	383-383	1	Gap in Lactuca dissecta and L. dolichophylla

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		1	rpl32-trnL
No.	Position	Length (nt)	Description
32	393-408	16	Gap in Launaea sarmentosa
33	400-400	1	Gap in Lagedium sibiricum and Mulgedium tataricaum
34	402-408	7	Insertion of "AGTTTTT" in Crepis multicaulis, gap in all other taxa
35	420-436	17	Gap in Stenoseris auriculiformis and Chaetoseris roborowskii
36	456-456	1	Gap in Launaea sarmentosa
37	502-524	23	Insertion of "TTTTTAGTAATTACTATATGAAA" in Lactuca parshii, gap in all other taxa
38	545-550	6	Insertion of "AATTTT" (replicate) in in one small clade of <i>Melanoseris</i> , i.e. <i>Chaetoseris likiangensis</i> , <i>Ch. grandiflora</i> , <i>Stenoseris tenuis</i> , <i>S. graciliflora</i> , including hybrids between them, gap in all other taxa
39	581-594	14	Insertion of "ATTGTTGC(A)GATATT" in Lactuca dissecta, L. dolichophylla, Steptorhamphus tuberosus, gap in all other taxa
40	598-643	46	Gap in Crepis multicaulis
41	603-630	28	Gap in Launaea sarmentosa
42	608-630	23	Insertion of "AAAGAACTTCATTGTTGAGATAT" (replicate) in Lactuca perennis, gap in all other taxa
43	674-710	37	Gap in Soroseris erysimoides and Prenanthes purpurea
44	690-695	6	Gap in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis
45	693-693	1	Insertion of "A" in all Faberia, gap in all other taxa
46	696-701	6	Insertion of "T(G)GGTTA" in Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Cicerbita alpina, gap in all other taxa
47	696-706	11	Gap in all taxa but Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis, Cicerbita alpina, all Notoseris taxa but two of N. rhombiformis
48	702-706	5	Insertion of all Notoseris taxa but two of N. rhombiformis, gap in all other taxa
49	731-731	1	Insertion of "C" in Crepis multicaulis, gap in all other taxa
50	737-737	1	Gap in Scariola orientalis, insertion of "T" in all other taxa
51	743-743	1	Insertion of "A" in Soroseris erysimoides, gap in all other taxa
52	743-759	17	Gap in all taxa but Soroseris erysimoides and all Faberia
53	785-789	5	Insertion of "AAATA" (replicate) in Soroseris erysimoides, gap in all other taxa
54	802-806	5	Insertion of "ATCAAGT" in Scariola orientalis and S. viminea
55	802-813	12	Gap in all taxa but Cephalorrhynchus macrorhizus, Scariola orientalis, S. viminea
56	802-833	32	Gap in Launaea sarmentosa
57	807-813	7	Insertion of "ATCAAGT" in Cephalorrhynchus macrorhizus, gap in all other taxa
58	820-826	7	Gap in Cicerbita alpina
59	821-856	36	Gap in Lactuca inermis
60	832-832	1	Insertion of "A" in Leontondon tuberosus, gap in all other taxa
61	832-833	2	Gap in Crepis multicaulis
62	842-842	1	Insertion of "T" in all Pterocypsela and Cicerbita alpina, gap in all other taxa

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No.	Position	Length (nt)	Description
63	850-852	3	Gap in Parasyncalathium souliei
64	861-862	2	Gap in Lactuca inermis
65	891-897	7	Gap in all taxa but Prenanthes purpurea and Mulgedium bracteatum
66	897-897	1	Gap in Prenanths purpurea
67	903-907	5	Insertion of "ATAAT" (replicate) in Lactuca perennis, gap in all other taxa
68	930-951	22	Gap in all taxa but Lactuca sativa and L. serriola
69	935-951	17	Gap in Lactuca sativa
70	964-967	4	Gap in Paraprenanthes yunnanensis, P. saggitiformis, P. longiloba
71	965-965	1	Gap in Crepis multicaulis
72	966-966	1	Gap in Scariola orientalis, S. viminea, Lactuca sativa and L. serriola
73	971-973	3	Insertion of "GAC" in Paraprenanthes polypodifolia, gap in all other taxa
74	971-978	8	Gap in all taxa except Lactuca inermis and Paraprenanthes polypodifolia
75	974-978	5	Insertion of "AGGAC" in Lactuca inermis, gap in all other taxa
76	983-983	1	Insertion of "C" in Crepis multicaulis, gap in all other taxa
77	987-989	3	Gap in all taxa except Launaea sarmentosa, Leontodon tuberosus, Crepis multicaulis
78	988-989	2	Gap in Launaea sarmentosa
79	989-989	1	Insertion of "A" in Leontondon tuberosus, gap in all other taxa
80	1009-1020	12	Gap in all Paraprenanthes but P. pilipes and Notoseris melanantha
81	1013-1018	6	Gap in Cicerbita sikkimensis and Chaetoseris cyanea complex
82	1048-1054	7	Insertion of "AATTTGA" (replicate) in Lactuca inermis, gap in all other taxa
83	1086-1090	5	Gap in Lactuca undulata
84	1101-1101	1	Insertion of "C" in Stenoseris auriculiformis, gap in all other taxa