Himalayan Origin and Evolution of *Myricaria* (Tamaricaeae) in the Neogene

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Abstract

**Background:** *Myricaria* consists of about twelve-thirteen species and occurs in Eurasian North Temperate zone, most species in the Qinghai-Tibet Plateau (QTP) and adjacent areas.

**Methodology/Principal Findings:** Twelve species of *Myricaria* plus two other genera *Tamarix* and *Reaumuria* in Tamaricaeae, were sampled, and four markers, ITS, rps16, psbB-psbH, and trnl-trnF were sequenced. The relaxed Bayesian molecular clock BEAST method was used to perform phylogenetic analysis and molecular dating, and Diva, S-Diva, and maximum likelihood Lagrange were used to estimate the ancestral area. The results indicated that *Myricaria* could be divided into four phylogenetic clades, which correspond to four sections within the genus, of them two are newly described in this paper. The crown age of *Myricaria* was dated to early Miocene ca. 20 Ma, at the probable early uplifting time of the Himalayas. The Himalayas were also shown as the center of origin for *Myricaria* from the optimization of ancestral distribution. Migration and dispersal of *Myricaria* were indicated to have taken place along the Asian Mountains, including the Himalayas, Kunlun, Altun, Hendukosh, Tianshan, Altai, and Caucasus etc., westward to Europe, eastward to Central China, and northward to the Mongolian Plateau.

**Conclusions/Significance:** *Myricaria* spatiotemporal evolution presented here, especially the Himalayan origin at early Miocene ca. 20 Ma, and then migrated westward and eastward along the Asian mountains, offers a significant evolutionary case for QTP and Central Asian biogeography.

Introduction

The Tamaricaeae contains about eighty species [1] and four genera: *Tamarix*, *Myricaria*, *Reaumuria*, and *Hololachna* [2]. This family, and Frankeniaceae, are defined as the salt-gland anatomical lineage [3]. *Myricaria* consists of about twelve - thirteen species [4–7] and occurs in Northern Temperate zone of Eurasia, mainly along the Asian mountains. There are eight species in Himalayas, many are endemic, thus forming a center of diversity for *Myricaria* (see Figure 1).

Desvaux (1825) established the genus *Myricaria* and Niedenzu [8] presented the first classification. Zhang & Zhang [4] studied *Myricaria* in China and recognized ten species; they presumed the Himalayas to be the center of origin based mainly on the distribution of species. Gorschkova [7] described six species belonging to two sections in the flora of the former USSR.

Another issue relevant to *Myricaria* systematics is the species *Myricaria elegans*. Ovezinrlikov & Kinzikaeva [9] erected the genus *Myrtama* based on this species but it caused some controversy. Zhang et al. [10] used ITS sequence data to study the relationships within Tamaricaeae and regarded *Myrtama* as an intermediate genus between *Myricaria* and *Tamarix* [11]. After sampling four species from *Myricaria* and sequencing ITS, rbsL, and tRNAs Ser (GCU) and Gly (UCC), Gaskin et al. [1] found *Myrtama* and *Hololachna* to be distinct within Tamaricaeae, as did Zhang et al. [10]. However, based on additional sequence data, Hua et al. [12] and Wang et al. [13] confirmed that *Myrtama* should be included in *Myricaria*. Sampling ten species of *Myricaria* and sequencing cpDNA *psbA-trnH* and the *rpl16* intron, Liu et al. [14–15] investigated the species-level phylogeographical patterns of *Myricaria* in western China as well as the origin of *M. laxiflora*, a unique subtropical species of conservation concern from the Three Gorges of the Yangtze River in Sichuan and Hubei provinces. The Himalayas were proposed as the center of origin of *Myricaria* by Liu et al. [14] with the estimated age of origin 1.46–2.30 Ma. Closely associated with the distribution pattern of *Myricaria* and related taxa, the QTP and Himalayan uplift during the Neogene are hypothesized to be a major influence on organism evolution in Asia [e.g. 16–17]. Following collision of the Indian and Eurasian continents at ca. 50 Ma, the altitude and range of the QTP near the Oligocene-Miocene boundary became sufficient to trigger a reorganization of the Asian climate, as evidenced by the beginning
of loess deposition in the Chinese Loess Plateau and the Junggar Basin [18–20]. Some evidence confirms that the central areas of the QTP were raised to present altitudes by that time [16,21–22] and uplift of the Himalayas may have also begun at that time [22–23]. Uplift of peripheral portions of the plateau has continued at various intervals [24–28]. A major uplift of QTP is often suggested to have occurred at 8 Ma, which also coupled with global cooling, even though Molnar [29] considered this uplift evidence to be inconclusive. Uplift of the QTP and global climate cooling and aridification [27] have been suggested causes for the evolution of many organisms [30–36]. As these studies have shown, rapid diversification of lineages in the QTP resulted in the migration of some species into other temperate regions, such as Central Asia, the Arctic, the Mediterranean (Caucasus-Alps) and southern Asia. Of these, connections between the QTP and adjacent arid, more northern areas can often be discerned, for example in recent studies on Hippophae rhamnoides (Elaeagnaceae) [37], Caragana (Fabaceae) [38], and Astragalus (Fabaceae) [39]. Linkage of the QTP and Africa and/or the Mediterranean is illustrated by Begonia [40] and Uraria (Annonaceae) [41]. An example linking the QTP and Southeast Asia is Paini (Anura: Dicroglossidae) [42].

The origin of Myricaria has been associated with the QTP and Himalayas but justification has been weak. Zhang & Zhang [4] presumed that Myricaria originated from the Himalayas, only based on species distribution of the genus, whereas same opinion conducted by Liu et al. [14] from a phylogeography. Here we attempt to examine the origin and evolution of this genus and link it to the Himalayan uplift to explain the causes of its evolutionary patterns. In addition, the classification and distribution of Myricaria are examined using molecular phylogeny and biogeography.

**Materials and Methods**

**Taxa sampled**

Twelve species (seventeen samples) of *Myricaria* plus seven species from the outgroups *Tamarix* and *Reummuria* served as sources of DNA material (Table 1). The herbaria utilized in this study were as follows: HNWP (Northwest Institute of Plateau Biology, Chinese Academy of Sciences (CAS), Xining, Qinghai); SHI (Shihezhi University, Shihezhi, Xinjiang); and XJBI (Xinjiang Institute of Ecology and Geography, CAS, Urumqi, Xinjiang), as well as the LE (Komarov Botanical Institute, Russian Academy of Sciences, St. Petersburg, Russia).

**DNA sequencing**

Total genomic DNA was extracted using the CTAB method [43]. The polymerase chain reaction (PCR) was used for amplification of double stranded DNA. The 25 μl reaction system contained 0.25 μl of Ex Taq, 2.5 μl of 10× Ex Taq buffer (Mg²⁺ concentration of 25 mM), 2.0 μl of dNTP mix (2.5 mM concentration for each dNTP), 1 μl of the forward and reverse primers at 5 umol/μl, and 5.3 μl of template DNA. The following primers were used: for ITS: ITS1-f (5’-AGA AGT GTT TGG TTT GCT GTT TCC GTA GC-3’) and ITS4-r (5’-TCC GCT TAT TAT GAT TAT GC-3’), for trnL-trnF; trnL-f (5’-CGA AAT CGG TAG ACG GTA GTG CTA CG-3’) and trnF-r (5’-ATT GTG ACT GGT GAC AGC GAG AG -3’), for the intron of rps16: rps16-f (5’-GTG GTA GAA AGC -3’), and for rps16-r (5’-TCG GGA TCG AAC ATC AAT TGC AAC-3’) [44]; and the intergenic spacer pohB-pohH: pohB-pohH-f (5’-TTCAACAGTTTTGTTAGCCA-3’) and pohB-pohH-f (5’-AGAGTTTTTGTGATGTTGA-3’) [45].

The protocol for amplification consisted of an initial hotstart at 95°C for 2 min, followed by 30 cycles of denaturation at 94°C for 30 s, annealing at 52°C for 30 s, extension at 72°C for 90 s, and a final extension at 72°C for 10 min. PCR products were purified using the PEG precipitation procedure [46]. These were sequenced using an ABI Prism 3730 Genetic Analyzer (Shanghai Sanggon Biological Engineering Technology & Service, Shanghai, China). Sequences were aligned using CLUSTAL X software [47], and then adjusted by hand. All gaps were treated as missing characters. Finally, a combined dataset consisting of four sequences of ITS and three cpDNA trnL-trnF, rps16, and pohB-pohH, was prepared for phylogenetic analysis.

**Phylogenetic analysis and divergence time estimate**

The sequence dataset from twelve species (seventeen samples) of *Myricaria* plus seven species of *Tamarix* and *Reummuria* yielded 3202 aligned nucleotide characters from four genes: ITS, trnL-trnF, rps16, and pohB-pohH. The incongruence length difference (ILD) test of the four gene datasets was carried out using PAUP* [48], to test potential conflicts between the different DNA fragments. This test was implemented with 100 partition-homogeneity test replicates, using a heuristic search option with simple addition of taxa, TBR branch swapping and MaxTrees set to 1000. 0.222 of incongruence length difference (ILD) tests [48] showed that the four gene datasets were not incongruent.

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**Figure 1. Distribution of *Myricaria* species (a,b), the information obtained from floras and herbaria, mainly in China. Geographical division of QTP eastern and western portions is shown in a red broken line (c). The Himalayan origin and dispersal routes along the Asian mountains are illustrated in arrows (a).**

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Estimation of phylogenetic relationships and divergence time was conducted using a Bayesian method implemented in BEAST 1.5.4, employing a relaxed clock model [49–50]. We used the uncorrelated lognormal molecular clock model with a Yule process for the speciation model, GTR+I+G for the substitution model (estimated for the dataset), and a normal distribution with SD of 1 as priors on the calibration nodes to accommodate for calibration uncertainty. Minimum ages for the two normal priors were constrained to the root of all taxa and a node respectively, the family Tamaricaceae 70 Ma, and genus *Tamarix* 25 Ma, with a detailed description as follows. A Markov chain Monte Carlo analysis was run for 50 million generations and sampled every 1,000 generations, and two independent runs were performed to confirm the convergence of the analysis. The stationarity of each run was examined using the effective sampling size of each parameter (>$200$). The last 40 million generations were used to construct the maximum clade credibility tree and the associated 95% highest posterior density distributions around the estimated node ages.

### Table 1. List of Sampled Taxa, Vouchers and Genebank Accession Numbers.

| Species | Voucher | Source | ITS | trnL-trnF | rpo16 | psbA-psbB |
|---------|---------|--------|-----|-----------|-------|-----------|
| *Myricaria Desv.* | | | | | | |
| *M. alopecuroides* Schrenk. | P. Yan 3650 (SHI) | Tashikurgan, Xinjiang, China, alt. 3650m | KJ729654 | KJ729806 | KJ729756 | KJ729705 |
| *M. alopecuroides* Schrenk. 1 | Tibet-Xinjiang Exp. 1034 (HNWP) | Supekpiya, Yecheng, Xinjiang, China, alt. 2800m | KJ808603 | KJ808634 | KJ808619 | KJ808649 |
| *M. bracteata* Royle | Y.H. Wu 36461 (HNWP) | Nuomuhong, Dulan, Qinghai, China, alt. 2840m | KJ729655 | KJ729807 | KJ729757 | KJ729706 |
| *M. elegans* Royle 1 | P. Yan 3999 (SHI) | Bandir, Tashkurgan, Xinjiang, China, alt. 3000m | KJ808604 | KJ808635 | KJ808620 | KJ808650 |
| *M. elegans* Royle 2 | P. Yan 7178 (SHI) | Mazhaxi, Yecheng, Xinjiang, China, alt. 3600m | KJ808605 | KJ808636 | KJ808621 | KJ808651 |
| *M. elegans* Royle 3 | P. Yan 7378 (SHI) | Ritu, Tibet, China, alt. 4600m | KJ808606 | KJ808637 | KJ808622 | KJ808652 |
| *M. germanica* (L.) Desv. | I.O. Baltulin, Asalbaiev s.n. (LE) | Zajasanskaya depression, E. Kazakhstan | KJ808607 | KJ808638 | ---- | ---- |
| *M. luxiflora* (Franch.) Y.Y. Zhang et Y.J. Zhang 1 | Wuhan Bot Gard | Wuhan Bot Gard, Hubei, China | KJ808608 | KJ808639 | KJ808623 | KJ808653 |
| *M. luxiflora* (Franch.) Y.Y. Zhang et Y.J. Zhang 2 | Wuhan Bot Gard | Wuhan Bot Gard, Hubei, China | KJ808609 | KJ808640 | KJ808624 | KJ808654 |
| *M. paniculata* Y.Y. Zhang et Y.J. Zhang | B.Z. Guo; W.Y. Wang 21930 (HNWP) | Linzhi, Tibet, China, alt. 2000m | KJ808610 | ---- | ---- | KJ808655 |
| *M. platyphylla* Maxim. | Z.Y. Yang; L.M. Ke 5711 (XJBI) | Houxia, Urumqi, Xinjiang, China | KJ808611 | KJ808641 | KJ808625 | KJ808656 |
| *M. prostrata* Hook.f. et Thomson ex Benth. et Hook.f. | P. Yan 7242 (SHI) | Hechakou, Hetian, Xinjiang, China, alt. 5000m | KJ808612 | KJ808642 | KJ808626 | KJ808657 |
| *M. pulchera* Batalin | L.M. Ke 121 (XJBI) | Ermuchang, Shaya, Xinjiang, China alt. 4350m | KJ808613 | KJ808643 | KJ808627 | KJ808658 |
| *M. rosse W.W. Sm.* | R.F. Huang G89-485 (HNWP) | Milinpaqui, Tibet, China, alt. 4530m | KJ808614 | ---- | KJ808628 | KJ808659 |
| *M. squamosa* Desv. | P. Yan 4002 (SHI) | Bandir, Tashkurgan, Xinjiang, China, alt. 3000m | KJ729658 | KJ729810 | KJ729760 | KJ729709 |
| *M. squamosa* Desv. 1 | Y.H. Wu 3077 (HNWP) | Beishan, Huzhu, Qinghai, China, alt. 2700m | KJ808615 | KJ808644 | KJ808629 | KJ808660 |
| *M. wardii* C.Marquand Sun X.Y | R.H. Ree, S.K. Wu 30159 (LE) | Lanzhi-Bomi, Tibet, China, alt. 3550m | KJ808616 | KJ808645 | KJ808630 | KJ808661 |
| *Reaumuria Linn.* | | | | | | |
| *R. kaschgarica* Rupr. 1 | Tibet-Xinjiang Exp. Team 5166 (HNWP) | Tashkurgan, Xinjiang, China | KJ808617 | KJ808646 | KJ808631 | KJ808662 |
| *R. kaschgarica* Rupr. 2 | Y.M. Duan 84-A-012 (XJBI) | Ruqiang, Xinjiang, China, alt. 3080m | ---- | KJ808647 | KJ808632 | KJ808663 |
| *R. soongarica* (Pall.) Maxim. | Tibet-Xinjiang Exp. Team 5098(SHI) | Tashkurgan, Xinjiang, China, alt. 2300m | KJ808618 | KJ808648 | KJ808633 | KJ808664 |
| *Tamarix L.* | | | | | | |
| *T. karakalensis* Freyn | K.B. Blinkovskiy 12 VIII 1953 (LE) | C. Kopetdag, Ashikhbad, Turcominia | KJ729659 | KJ729811 | KJ729761 | KJ729710 |
| *T. laxa* Willd. | O.N. Demina 18 V 2001 (LE) | Orlovsky, Bostov, Russia | KJ729660 | ---- | KJ729762 | KJ729711 |
| *T. meyeri* Boiss. | M.R. Tanybaeva 12 V 2007 (LE) | Turkestan Ridge, Kirgiztan | KJ729661 | KJ729812 | KJ729763 | KJ729712 |
| *T. ramosissima* Ledeb. | N.A. Brykova s.n. 10 VII 1998 (LE) | Orlovsky, Bostov, Russia | KJ729662 | KJ729813 | KJ729764 | KJ729713 |
Optimization of ancestral distributions

**Tamaricaceae root constrained.** Tamaricaceae is included in the order Caryophyllales [2,51] and has no reliable macrofossil record. According to molecular dating [52–54], the divergence time of the order has been estimated as ca. 100 Ma. Tilloy [55] considered that the extant woody families originated during the Cretaceous to early Eocene, while herbaceous families appeared during the late Oligocene to Miocene. For instance, the woody families Ulmaceae and Fabaceae appeared at about 70 Ma [54,56] and 70–60 (~50) Ma [54,56,57] respectively. Families related to the Tamaricaceae, such as Polygonaceae and Amaryllidaceae/Chenopodiaceae, have an approximate age of ca. 65 Ma [58]. The two subfamilies of the Caryophyllaceae have an approximate age of 40–55 Ma according to the age of the inflorescence fossil Caryophylla flora palenigena, and the family has an possible age of ca. 75 (60–80) Ma [59–60]. Even though the ancestor of the related families Tamaricaceae and Frankeniaceae has been dated to 43–30 Ma [56], Tamaricaceae itself has had variable dating results. Bell et al. dated it to (72–) 60–58 (~44) [54], while Wikström et al. placed it at 52–37 Ma [56], and Schuster et al. at (125–) 118.7–110 (~90.7) [61]. In the light of these estimates, a balanced age for Tamaricaceae could be suggested as about 70 Ma; this estimate was chosen as the family root for molecular dating.

The earliest reliable fossil record of *Tamarix* is Miocene, from the Yunnan province of China [62]. Most of the available fossils are from the Miocene, therefore, the genus might be hypothesized to have had an origin at least in early Miocene. However, considering its wide distribution in Europe, Africa, Asia, and North America, and our limited samples mainly from China, we conservatively assigned an age of late Oligocene-early Miocene, at ca. 25 Ma for *Tamarix*.

**Areas**

In accordance with the distribution of *Myricaria* species along the Asian mountains (Figure 1), we divided the distribution into six areas, namely, A: the eastern Himalayas, including the eastern QTP, the Hengduan mountains, and northern and central China; B: the western Himalayas, including the western QTP and the Pamir-Alai, Kunlun-Altn, and Hendukosh mountains; C: the Tianshan mountains and Junggar-Turan deserts; D: Altai-Siberia; E: the Mongolian Plateau; and F: Asia Minor-Caucasus-Europe. These six areas are distinct in biodiversity, vegetation, and floristics [16–18,30,32,36–38].

**Optimization of ancestral distributions**

To infer biogeographical events, three methods were used: a parsimony-based procedure Diva [63], S-Diva [64] and a maximum likelihood-based DEC model (Lagrange; [65–66]). These three approaches are simultaneously considered so that to assess the relevant biogeographical processes, such as vicariance, dispersal, and extinction.

Diva. Dispersal–vicariance analysis optimizes distributions for each node of the tree by minimizing the number of assumed dispersals and extinctions, and favors vicariance events [63,67]. The Diva program reconstructs widespread ancestral distributions, restricting them to single areas. Because allopatric speciation by vicariance is the null model in Diva, vicariance and range division would always be the preferred explanation if the ancestors were widespread. To avoid inferring a widespread ancestor at the root because of the presence of widespread extant taxa, a limit of two areas was set (maxareas = 2) in Diva [63]. The phylogenetic typology of the BEAST tree (Figure 2) was input for Diva analysis.

S-Diva. (or Bayes-Diva) [64] is a program which complements Diva and implements the methods of Nylander et al. [68] and Harris et al. [21], determining statistical support for ancestral range reconstructions using multiple trees from Bayesian analysis. This has the advantage that uncertainties in phylogenetic inference can be taken into account. One hundred Bayesian MCMC trees with the last stable typologies from BEAST, and a BEAST tree typology (Figure 2) were input into the S-Diva program.

Lagrange. A valuable new biogeographical method is parametric likelihood analysis, with a dispersal–extinction–cladogenesis model [67], as implemented in Lagrange v. 2.0.1 [63]. This method calculates the likelihood of biogeographical routes and areas occupied by the most recent common ancestor (MRCA) for a given phylogenetic tree topology (BEAST tree, Figure 2) and the present distribution of taxa. Therefore, dispersal and vicariance of lineages, represented by connection areas, can be estimated by the probabilities. This is thus a form of MRCA area reconstruction differing from the parsimony approach of Diva.

**Results**

**Phylogenetic analysis and divergence time estimate**

The phylogenetic tree obtained from Bayesian inference in BEAST showed that *Myricaria* is monophyletic and *Myrtama* should be included in *Myricaria* rather than treated as a distinct genus (Figure 2). Within *Myricaria*, four clades were recognized, two corresponding to the existing sections *Parallelantherae* Ndz. and *Renantherae* Ndz., the other two represent new groups to be named as sections *Alpinae* and *Laxiflorae* (see Appendix S1). Flowers and filaments of the plants are illustrated in Figure 2, to show the characteristics of the four sections. In the present phylogenetic tree, the clades of the genus and the sections have strong support, confirming the validity of taxa at the ranks of genus and section. Section *Renantherae* comprises the most species in the genus, and has two subclades. The two widely distributed species of this section, *M. alopecroides* and *M. squamosa*, are located in each subclade. The crown age of *Myricaria* was ca. 20 Ma, and 8.83–6.35 Ma for four sections.

**Optimization of ancestral distributions**

The results of the three approaches Diva, S-Diva, and Lagrange (Figure 3) showed a consistent and strongly supported pattern, particularly at the ancestral nodes for *Myricaria* (AB), and among the four sections, *Parallelantherae* (B), *Alpinae* (A), *Laxiflorae* (A), and *Renantherae* with AB from Diva and S-Diva, whereas only with ABCDEF/B from Lagrange. On the whole, AB, A, and B, namely the Himalayas and the QTP, should be considered as ancestral areas in *Myricaria*. The events occurring in areas C, D, E, and F, were considered to be dispersals, several of which can be distinguished. *M. prostrata* occurs in the Himalayas, and its western Himalayan distribution was indicated to be a dispersal event from the eastern Himalayas. *M. bracteata* in sand lands of the Mongolian Plateau was shown to be a migrant from the eastern Himalayas, Hengduan Mountains, and Northern China. Whereas the distribution of *M. germanica* in Asia Minor-Caucasus-Europe was came from the western Himalayas.

**Discussion**

Phylogenetic division of sections within *Myricaria*

Niedenzu [8] divided *Myricaria* taxa into two sections: *Parallelantherae* Ndz. and *Renantherae* Ndz. Gorschakova [7] accepted this classification system in the Flora of the USSR. Zhang & Zhang [4], however, considered that the establishment of infrageneric ranks was not appropriate due to its complicated and
variable morphological characters. Therefore, in the Flora of China [5–6] there is no division of infrageneric sections.

However, our phylogenetic tree (Figure 2) yielded a clear phylogenetic division of four sections, including two that are new. Of them the Himalayan and QTP section Alpinae, containing M. prostrata, M. rosea, and M. wardii, is characterized by the prostrate and recumbent, and with an adaptation to high altitudes of 3000–5300 m. Section Laxiflorae, comprises of only species M. laxiflora, endemic to the subtropical area of the Sichuan and Hubei provinces in eastern China. Detailed descriptions of two new sections are given in Appendix S1.

Myricaria elegans Royle was originally described from the Kunawar region of the western Himalayas [69]. Based on this species and its characters of 10 stamens, flat leaves, and no obvious style, the genus Myrtama was established [9]. Qaiser & Ali [70] named it Tamaricaria, whereas Baum [71] moved Myricaria elegans to Tamarix as T. ladachensis. As mentioned, Zhang et al. [10] and Gaskin et al. [1] accepted Myrtama at generic rank. Zhang et al. [10] considered that Myricaria elegans was an intermediate and hybrid genus between Myricaria and Tamarix, related more to Tamarix. However, Hua et al. [12] and Wang et al. [13], based on sequence data, found that it would be appropriately placed in Myricaria. Our results (Figure 1) also show that inclusion of Myricaria elegans in Myricaria is suitable, since the whole of Myricaria, including Myricaria elegans, has strong support (100%) (Figure 2). This is in accordance with evidence from morphological classification [4] and molecular phylogeny [12–14]. The former conclusion supporting retention of Myrtama [10] was only based on ITS

Figure 2. Chronogram of Myricaria and outgroups Tamarix and Reaumuria in Tamariaceae, with maximum clade credibility performed by BEAST. Dating values are plotted at the right of the nodes, and posterior probability support of more than 95% is labeled as ‘*’ at the nodes. Two vertical lines are labeled at 20 Ma and 8 Ma, corresponding respectively to two stages and two high-altitude ranges (blue ranges in A, B) of the QTP uplift, Himalayan motion, and rapid and major-range uplift. Four sections within Myricaria are shown, along with flowers and degree of union of the filaments. In section Alpinae, flower and filament status refers to M. rosea, and in section Renantherae, the flower and filaments, above right, refer to M. bracteata. The filaments below left refer to M. germanica and those below right refer to M. squamosa.
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sequence data, which is probably not sufficient evidence [72]. While in the phylogeny of Gaskin et al. [1] (their Figure 2), only four species were sampled, and four species as a clade had a strong bootstrap support (99%) for the inclusion in *Myricaria* [1].

**Himalayan origin, ancestral inheritance, and multidiversification in the Himalayas**

Our estimated crown age of ca. 20 Ma for *Myricaria* (Figure 2) falls into the probable early range of the Himalayan uplift in early Miocene [22], consequently allows us to speculate that uplift of this mountain range caused the origin of *Myricaria*. The biogeographical analytical result of Diva, S-Diva and Lagrange, showing the combined Himalayan area AB as the ancestral area for *Myricaria* (see Figure 3), which also supports a Himalayan origin. The present molecular dating results are in contrast to previous phylogeographical opinion of a main divergence event at the implausible age of 1.46–2.30 Ma in the Plio-Pleistocene [14].

The southern and northern slopes of the Himalayas differ dramatically in temperature and precipitation. *Myricaria* species occurring on the northern slope are generally xeric, same as those of the main plateau [73]. The western portions of the Himalayas and adjacent QTP are more arid than the eastern parts [27,73], see Figure 1c. Probably these differences are the cause of the persistent diversification between the eastern (A) and western Himalayas (B) for the *Myricaria* lineages (Figures 1 and 3). The eastern A contains the alpine species *M. rosea* and *M. wardii*, and the western B includes *M. elegans* and *M. prostrata*. These four species are endemic to the Himalayas, and occupy two of the four *Myricaria* sections. In particular, the western Himalayas (B) is an important geographical node and dispersal center for *Myricaria*, with movement toward the Pamir-Alai, Hendukosh, Tianshan, and Kunlun-Altun mountains, etc. Noticeable, the Himalayas as a union was divided into A and B two times (see Figure 3), once at the time of generic origin and diversification, and the other at the diversification node of the sections *Renantherae* and *Laxiflorae*. Overall, the Himalayan areas AB, A, and B as the ancestors occurred at least seven times at the nodes in Figure 3. In detail, the Himalayan union AB as an ancestral area appeared at two nodes, with estimated ages of 20.25 Ma (genus crown age) and 10.07 Ma, the eastern Himalayas A at three nodes with ages of 19.84 Ma,
Migration along Asian mountains

For plant migration and dispersal, mountains generally act as a route or corridor [84], such as the Himalayan corridor mentioned above. Whereas the Himalayas for Myricaria are regarded as the center of origin, other distributions outside of the Himalayas and QTP can be understood as dispersals or migrations eastward, westward and northward along Asian mountains (see Figure 1). In fact, the results of vicariance and dispersal from biogeographical analysis (Figure 3) show that except for divergence of phylogenetically basal clades located in the eastern and western Himalayas (A and B), most remaining events, occurring in areas such as the Tianshan-Jungger-Turan (C), Altais-Sibiria (D), Mongolian Plateau (E), and Asia Minor-Caucasus-Europe (F) resulted from dispersal events. As evidenced from Myricaria (see Figure 1), dispersal and migration were possible from the Himalayas to Asia Minor-Caucasus-Europe as shown by M. germanica, and to the sand lands of the Asian mountains very similar to those of Hippophae rhamnoides (Elaeagnaceae) [37], a species with another interesting distribution in North Temperate Eurasia. Hippophae rhamnoides includes nine subspecies, and has been shown to have originated from the QTP, or more exactly, the eastern QTP-Hengduan Mountains, and then to have radiated and dispersed in different directions. Here, Myricaria originated in the Himalayas union, not in the eastern Himalayas only as H. rhamnoides. However, the northwestern Himalayas for H. rhamnoides and Myricaria played an important node role in connecting with Central Asia and Europe, and both dispersal route and direction is very similar.

Supporting Information

Appendix S1 Two new sections within Myricaria. (DOC)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: MLZ. Performed the experiments: HHM. Analyzed the data: MLZ HXZ. Contributed reagents/materials/analysis tools: MLZ BV HXZ. Wrote the paper: MLZ SS.

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