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**EVOLUCIÓ, FILOGÈNIA I SISTEMÀTICA DEL COMPLEX
*ARCTIUM-COUSINIA***

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7. 1. Phylogeny and evolution of the *Arctium-Cousinia* complex (Compositae, Cardueae-Carduinae)

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RESUM. A aquest article s'investiguen la filogènia i l'evolució del complex *Arctium-Cousinia* que inclou els gèneres *Arctium*, *Cousinia*, *Hypacanthium* i *Schmalhausenia* i té la seva màxima diversitat a la regió Irano-Turania i a les muntanyes d'Àsia Central. Per aquest treball s'han generat seqüències dels marcadors ITS, i *rpS4-trnT-trnL* per 138 espècies, incloses 129 espècies de les aproximadament 600 que té el gènere *Cousinia*. Tal i com ja s'havia vist a anàlisis prèvies, *Cousinia* no és monofilètic donat que els subgèneres *Cynaroides* i *Hypacanthodes*, que sumen unes 30 espècies, són més propers a *Arctium*, *Hypacanthium* i *Schmalhausenia* (conjunt que s'ha anomenat clade Arctioide) que a *Cousinia* s.str. (o clade Cousinioide). La distinció entre els clades Arctioide i Cousinioide està recolzada per la morfologia del pol·len i el nombre cromosòmic tal i com havien posat de manifest autors anteriors. Pel que fa a la delimitació genèrica del clade Arctioide, la distribució dels caràcters morfològics tradicionalment considerats més determinants com la forma i armadura de la fulla i la morfologia de les bràctees involucrals, és en part incongruent amb les relacions filogenètiques derivades de les dades moleculars. Amb aquest treball no hem pogut aportar cap solució taxonòmica per aquest conflicte i interpretem que els esmentats caràcters són homoplàsics. Malgrat que la resolució filogenètica del clade Cousinioide és reduïda, es posa de manifest que els marcadors ITS i *rpS4-trnT-trnL* contenen una informació filogenètica limitada però valuosa. Un exemple és que quatre de les cinc espècies anuals de *Cousinia* s.str. incloses a les anàlisis cauen al mateix grup tot i pertànyer a dues seccions diferents. Aquesta pobresa de resolució filogenètica del clade Cousinioide probablement sigui conseqüència del reduït nombre de caràcters informatius combinat amb l'elevat nombre d'espècies d'aquest jove llinatge (uns 8.7 milions d'anys). S'hipotetitza que el mecanisme d'especiació d'aquest llinatge és eminentment al·lopàtric.

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Phylogeny and evolution of the *Arctium-Cousinia* complex (Compositae, Cardueae-Carduinae)

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The phylogeny and evolution of the *Arctium-Cousinia* complex, including *Arctium*, *Cousinia* as one of the largest genera of Asteraceae, *Hypacanthium* and *Schmalhausenia*, is investigated. This group of genera has its highest diversity in the Irano-Turanian region and the mountains of Central Asia. We generated ITS and *rpS4-trnT-trnL* sequences for altogether 138 species, including 129 (of ca. 600) species of *Cousinia*. As found in previous analyses, *Cousinia* is not monophyletic. Instead, *Cousinia* subgg. *Cynaroides* and *Hypacanthodes* with together ca. 30 species are more closely related to *Arctium*, *Hypacanthium* and *Schmalhausenia* (Arctioid clade) than to subg. *Cousinia* (Cousinioid clade). The Arctioid and Cousinioid clades are also supported by pollen morphology and chromosome number as reported earlier. In the Arctioid clade, the distribution of morphological characters important for generic delimitation, mainly leaf shape and armature and morphology of involucre bracts, are highly incongruent with phylogenetic relationships as implied by the molecular data. No taxonomic solution for this conflict can be offered, and the characters named are interpreted as homoplasious. Although phylogenetic resolution in the Cousinioid clade is poor, our ITS and *rpS4-trnT-trnL* sequences contain some phylogenetic information. For example, the six annual species of the Cousinioid clade fall into two groups. Poor phylogenetic resolution probably results from lack of characters and the high number of taxa in this species-rich and comparatively young (ca. 8.7 mya) lineage. We hypothesize that speciation in the Cousinioid clade was mainly allopatric.

KEYWORDS: *Arctium*, *Cousinia*, Irano-Turanian region, ITS, molecular clock, Pamir-Alay, *rpS4-trnT-trnL*, speciation, Tien Shan

INTRODUCTION

The *Arctium-Cousinia* complex (Compositae, Cardueae-Carduinae) in its most recent circumscription (Susanna & Garcia-Jacas, 2007) is a natural group composed of four genera. The largest genus of the group, one of the largest of the Compositae and among the 50 largest of flowering plants (Frodin, 2004), is *Cousinia* Cass. with more than 600 species (Mabberley, 2008). The other genera of the complex are *Arctium* L. (incl. part of *Cousinia*, see below) with 27 species, *Hypacanthium* Juz. with two species (Tscherneva, 1982) and the monotypic *Schmalhausenia* C. Winkl. *Lipskyella* Juz. and *Tiarocarpus* Rech.f., recognized as separate genera by Häffner (2000), were included in *Cousinia* by Susanna & Garcia-Jacas (2007).

Three characters never found in combination elsewhere in the tribe characterize this complex according to Susanna & al. (2003a). The receptacle has strongly twisted

scales, the achenes are always streaky (with wavy fringes), very often winged and without a nectary, and the pappus is formed by free deciduous bristles.

According to Rechinger (1986) and Knapp (1987), the vast majority of species of the *Arctium-Cousinia* complex is distributed in the Turkestan mountain region (Tien Shan and Pamir-Alay) and the Irano-Turanian region (Fig. 1; the “Oriental-Turanian Floristic Region” of Meusel & al., 1965). Within this range, following Knapp (1987), most species of *Cousinia* subgg. *Cynaroides* and *Hypacanthodes* grow only in the mountainous terrain of the Pamir-Alay range and in the western Tien Shan in Central Asia. The two species of *Hypacanthium* are endemic to the western Tien Shan, and the monotypic *Schmalhausenia* is endemic to the subalpine and alpine zone in the northern Tien Shan. *Arctium* s.str. is Eurosiberian in distribution. Those species of *Cousinia* included in *Arctium* by Duistermaat (1996) and Susanna & Garcia-Jacas (2007; see below) are, like the remainder of *Cousinia*, distributed in Central Asia.

Both morphological (Boissier, 1875; Kuntze, 1891; Dittrich, 1977; Tscherneva, 1988a–c; Duistermaat, 1996, 1997; Petit & al., 1996; Petit, 1997; Häffner, 2000) and molecular (Häffner & Hellwig, 1999; Garcia-Jacas & al., 2002; Susanna & al., 2003a, 2006) analyses of the *Arctium-Cousinia* complex concluded that the limits between the biennial *Arctium* and the perennial, biennial and only rarely annual *Cousinia* are unclear. This resulted in many reclassifications with many species changing generic affiliation. As one extreme, Kuntze (1891) proposed the classification of all species of *Cousinia* in *Arctium*.

The difficulties in generic delimitation arise from the incongruent distribution of several morphological, pollen, karyological and molecular characters.

Arctium has leaves which are always unarmed, often large (to 80 cm), and usually cordate. Such leaves are also found in *Cousinia* subg. *Cynaroides* (with the exception of *Cousinia korolkovii* Regel & Schmalh., *C. haesitabunda* Juz. and *C. chlorantha* Kult. with lanceolate leaves) and *C. grandifolia* Kult. of *Cousinia* subg. *Hypacanthodes*. This group of *Cousinia* species was referred to as the “Arctioid” group by Duistermaat (1996). In contrast, *Cousinia* subg. *Cousinia*, nine species of subg. *Hypacanthodes*,

all species of *Hypacanthium* and the monotypic *Schmalhausenia* have smaller leaves which usually are lanceolate and often very spiny. In *Arctium* and part of *Cousinia* subg. *Cynaroides* involucre bracts always end in a recurved hook whereas in most other species of *Cousinia* involucre bracts end in an unhooked spine. Whereas in species with hooked involucre bracts usually the entire capitulum is dispersed as a burr adhering to passing animals, most species of *Cousinia* release their achenes and/or disperse as tumbleweeds. As regards floral morphology, Duistermaat (1996) observed that the stigma of *Cousinia* is very different from that of *Arctium* and the “Arctioid” species of *Cousinia*. Whereas in the former the apical part of the style and the stylar branches are covered with long hairs, and a ring of sweeping hairs at the articulation of the base of the style apex is absent, the style is glabrous and cylindrical in the latter, and a ring of sweeping hairs is present at the thickened articulation. Häffner (2000) noted that a ring of sweeping hairs is also absent in *Schmalhausenia*. Outer florets in *Arctium* often have long and brightly coloured anther-tubes. Many species of *Cousinia* have brightly coloured appendages in the innermost involucre bracts, recalling those of *Carlina* L.

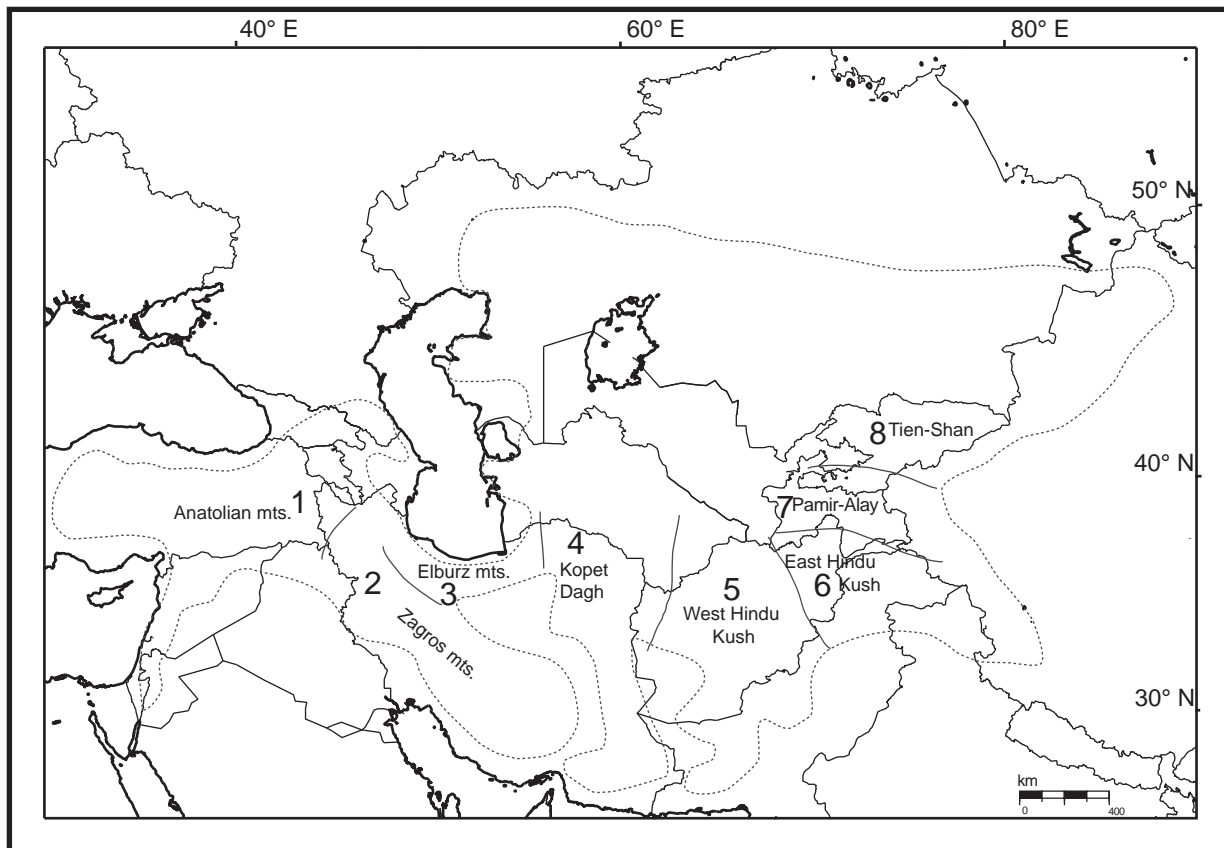


Fig. 1. Geographical distribution of the *Arctium-Cousinia* complex (excl. *Arctium* s.str.). The eight main centres of diversity recognized are based on Rechinger (1986) and Knapp (1987).

This distribution of characters (leaf, involucre bract, style morphology) led Duistermaat (1996) to the conclusion that the obviously “Arctioid” species of *Cousinia* should be transferred to *Arctium* within which their sectional classification by Tscherneva (1988a–c) should be retained. However, later Duistermaat (1997) vaguely suggested that all species of *Cousinia* subg. *Cynaroides* could be placed in *Arctium*. This latter proposal was followed by Susanna & Garcia-Jacas (2007) but does not solve the problematic placement of *Hypacanthium*, *Schmalhausenia* and *Cousinia* subg. *Hypacanthodes*.

Schtepa (1966, 1976) and Kuprianova & Tscherneva (1982) found that *Cousinia* subgg. *Cynaroides* and *Hypacanthodes* have orbicular and spiny pollen grains similar to those of *Arctium*, which they called the *Arctiastrum* pollen type, whereas *Cousinia* subg. *Cousinia* has oblong and smooth pollen grains which they called the *Cousinia* pollen type. *Schmalhausenia* (Susanna & al., 2003a) as well as *Hypacanthium* (S. López-Vinyallonga, unpub. data) have the *Arctiastrum* pollen type.

According to Duistermaat (1996), *Arctium* always has $x = 18$ and $2n = 36$ chromosomes. This number is also found in all species studied of *Cousinia* subgg. *Cynaroides* and *Hypacanthodes* (Tscherneva, 1985) and in *Schmalhausenia* according to Susanna & al. (2003b), and is perfectly correlated with the *Arctiastrum* pollen type. In contrast, *Cousinia* subg. *Cousinia*, with the *Cousinia* pollen type, has $2n = 18, 20, 22, 24$ and 26 chromosomes according to Ghaffari & al. (2006) and earlier authors. Finally, a DNA sequence analysis by Susanna & al. (2003a) grouped all species with $2n = 36$ chromosomes and the *Arctiastrum* pollen type in a monophyletic clade, the Arctioid clade, and species with $2n = 22, 24$ and 26 and the *Cousinia* pollen type in a second monophyletic clade, the Cousinioid clade.

In summary, pollen morphology, chromosome number and DNA data imply a subdivision of the *Arctium-Cousinia* complex into two major lineages which are not congruent with overall morphology particularly because parts of *Cousinia* subgg. *Cynaroides* and *Hypacanthodes* (Arctioid clade) are more similar to subg. *Cousinia* (Cousinioid clade), and other parts (the Arctioid species) are more similar to *Arctium*.

Of the genera and lineages introduced above, *Cousinia* subg. *Cousinia* is most remarkable by containing a high number of species in a comparatively small geographical area. After establishment of *Cousinia* by Cassini (1827), Candolle (1838) recognized 34 species, Bunge (1865) 126 species, Boissier (1875, 1888) 141 species (in the *Flora Orientalis* area), Winkler (1892, 1897) 267 species, and an additional 30 species were added in several publications by Bornmüller (1896–1941; listed by Rechinger, 1953). In her treatment of *Cousinia* for the *Flora of the USSR*, Tscherneva (1962) recognized about 262 species

in about 50 sections, and in his treatment of the genus for *Flora Iranica*, Rechinger (1972) placed more than 350 species in 58 sections. A careful comparison of the systems of Rechinger (1953, 1972, 1979) and Tscherneva (1962, 1988a, b) reveals an approximate number of ca. 630 species in 70 sections and three subgenera. Of these 630 species, ca. 600 belong into subg. *Cousinia*.

The molecular analysis of the *Arctium-Cousinia* complex by Susanna & al. (2003a) included 21 species of *Cousinia* subg. *Cousinia*, 1 species of subg. *Hypacanthodes* and 5 species of subg. *Cynaroides*. By increasing the sample of *Cousinia* subg. *Cousinia* to 106 species (of ca. 600), of subg. *Hypacanthodes* to 6 species (of 10) and of subg. *Cynaroides* to 14 species (of 20), where the sample included represents the large majority of subgeneric units recognized by Tscherneva (1962, 1988c) and Rechinger (1953, 1972, 1979), we pursue several aims. First, we want to investigate whether the subdivision of the *Arctium-Cousinia* complex into two lineages, the Arctioid clade with *Arctium* s.str., *Cousinia* subgg. *Hypacanthodes* and *Cynaroides*, *Hypacanthium* and *Schmalhausenia*, and the Cousinioid clade with *Cousinia* subg. *Cousinia* as found by Susanna & al. (2003a) and as supported by pollen morphology, chromosome number and DNA sequence data can be verified with a strongly enlarged species sample. Second, we aim at obtaining a better understanding of phylogenetic relationships within the Arctioid clade in order to resolve the character incongruences discussed above and to arrive at a new generic classification of this clade. Third, we want to investigate phylogenetic relationships within the large subg. *Cousinia*. The last aims both at examining the validity of the sectional (and subsectional) classification of this group as proposed by Tscherneva (1962, 1988a, b) and Rechinger (1953, 1972, 1979), and at providing a basis for discussing possible evolutionary mechanisms underlying the species-richness of subg. *Cousinia*.

MATERIAL AND METHODS

Plant material. — Samples for analysis were obtained either from silica-gel dried leaves of specimens collected in the wild, from fresh plants cultivated at the Botanic Institute of Barcelona, or from herbarium specimens deposited mainly in B, BC, JE, KAS, LE, M, and MJG (Appendix 1). The data described and discussed here are based on a sample of 138 species for which both ITS and *rpS4-trnT-trnL* could be obtained. These 138 species represent 60 sections of *Cousinia* (out of 70 described by Rechinger, 1972, 1979, and Tscherneva, 1988a–c), three species of *Arctium* and one each of *Hypacanthium* and *Schmalhausenia*. Four outgroup taxa were selected on the basis of previous analyses of the group (Garcia-Jacas

& al., 2002; Susanna & al., 2006). These were *Saussurea elegans* Ledeb., *S. maximowiczii* Herder, *Jurinea albi-caulis* Bunge and *J. humilis* (Desf.) DC. Eighty additional species were sequenced for ITS and 19 for *rpS4-trnT-trnL*. Results obtained for this additional sample are not presented and discussed only briefly. Voucher data and sources of material of the above 138 species are given in Appendix 1. Some of the ITS sequences were taken from previous studies but all *rpS4-trnT-trnL* sequences were newly generated.

DNA extraction, amplification and sequencing.

— Total genomic DNA was extracted either following the protocol of the CTAB method of Doyle & Doyle (1987) and Culling (1992), or following the manufacturer's protocols of the NucleoSpin[®] Plant Kit (Macherey-Nagel GmbH & Co. KG, Düren, Germany) or QIAprep[®] Miniprep (Qiagen Inc., Valencia, California, U.S.A.).

nrDNA ITS. — Double-stranded DNA of the internal transcribed spacer region (ITS1, 5.8S, ITS2) was amplified using 18S as forward primer and 28S as reverse primer (Muir & Schlötterer, 1999). Reactions were performed in 25.0 µl volumes with 10% 10× Biotherm buffer, 5% 50 mM MgCl₂ (GeneCraft, Münster, Germany), 2% of 20 mM dNTPs mix, 2% of each primer at 25 pmol/µl conc., 1.0 unit (0.2 µl) BioTherm[™] polymerase (GeneCraft) and 1.0 µl of template DNA (30–60 ng/µl). This was filled up to 25.0 µl with distilled sterilized water. In some cases 1.0 µl DMSO (dimethyl sulfoxide) per reaction tube was added to improve PCR amplifications (Frackman & al., 1998). The profile used for amplifications consisted of a preheat for 60 s at 94°C, followed by 35 cycles of 94°C for 18 s, 52°C for 30 s, and 72°C for 60 s and a post-treatment of 78 s at 52°C plus 8 min at 72°C. PCR products were purified using the NucleoSpin[®] Extract Kit (Macherey-Nagel, Düren, Germany) following the manufacturer's protocol.

Cycle sequencing reactions were conducted using 0.5–2.5 µl of the purified PCR product and ABI-PRISM[®] BigDye[™] Terminators v3.0 Cycle Sequencing Reagents (BD 3.0 in 10 ml reactions) for 30 cycles for 10 s at 96°C and 4 min at 55°C. Sequences were detected on automated sequencers (ABI 373 or 377).

For sequence analysis, ITS fragments were edited and assembled with the aid of Sequencher 4.1.2 (Gene Codes Corporation, Ann Arbor, Michigan, U.S.A.). Edited sequences were aligned with MacClade 4.1 (Maddison & Maddison, 2000).

cpDNA rpS4-trnT-trnL. — The double-stranded DNA intergenic spacer *rpS4-trnT-trnL* was amplified by PCR with rpS4R2 (Shaw & al., 2005), trnL-b (Taberlet & al., 1991), trnA2 (Cronn & al., 2002) and trnA2R (5'-AGG TTA GAG CAT CGC ATT TG-3' designed for this study) as PCR primers. When the rpS4R2 (forward)/trnL-b (reverse) primer combination failed to generate

double-stranded products, two separate PCR reactions were performed with the following primer combinations: rpS4R2/trnA2R (reverse) and trnA2(forward)/trnL-b. Reactions were performed in 25.0 µl volumes with 10% 10× AmpliTaq buffer, 10% 50 mM MgCl₂, 10% of 20 mM dNTPs mix, 2% of each primer at 25 pmol/µl conc., 1.0 unit (0.2 µl) AmpliTaq DNA polymerase (Applied Biosystems, Foster City, California, U.S.A.), and 5.0 µl of template DNA (30–60 ng/µl). The volume was filled up to 25.0 µl with distilled sterilized water. The profile used for amplification consisted of a preheat for 1 min 35 s at 95°C, followed by 35 cycles of 95°C for 1 min 30 s, 52°C for 1 min 30 s and 72°C for 2 min and a post-treatment of 10 min at 72°C.

Double-stranded PCR products were purified with either the QIAquick[®] Purification Kit (Qiagen Inc., Valencia, California, U.S.A.) or DNA Clean & Concentrator-5 (Zymo Research, Orange, California, U.S.A.) following the manufacturer's protocols. Direct sequencing of the amplified DNA segments was performed using 0.5–1.0 µl of the purified PCR product with a "Big Dye[®] Terminator v3.1 kit" (Applied Biosystems, Foster City, California, U.S.A.). The profile used consisted of a hot start at 96°C for 1 min followed by 25 cycles of 10 s at 96°C, 5 s at 50°C and 4 min at 60°C. Nucleotide sequencing was performed at the Serveis Científic-Tècnics of the University of Barcelona on an ABI PRISM 3700 DNA analyzer (Applied Biosystems, Foster City, California, U.S.A.). Nucleotide sequences were edited with Chromas 2.0 (Technelysium Pty. Ltd., Tewantin, Australia) and aligned visually by sequential pairwise comparison (Swofford & Olsen, 1990).

Data matrices are available on request from the corresponding author.

Analyses. — *Bayesian Inference*. — Bayesian analyses of the ITS1 + ITS2, *rpS4-trnT-trnL* and combined ITS1 + ITS2 + *rpS4-trnT-trnL* datasets were performed with MrBayes software package 3.0b4 (Huelsenbeck & Ronquist, 2001). Bayesian analyses consist of maximum likelihood (ML) comparisons of trees where tree topology and ML parameters are permuted using a Markov Chain Monte Carlo method and sampled periodically. The sample trees are drawn from a posterior probability distribution (Huelsenbeck & Ronquist, 2001). In order to select the best-fit model of substitution, the program Modeltest 3.5 was used (Posada & Crandall, 1998, 2001).

Following the results for DNA evolution of our data as indicated by ModelTest, the ML parameters in MrBayes were set as follows: Nst = 6, Rmat = for ITS dataset: A-C = 0.7720, A-G = 2.7242, A-T = 1.2057, C-G = 0.3372, C-T = 9.8048, G-T = 1.0000 (Proportion of invariable sites [I] = 0.1915; Gamma distribution shape parameter = 0.6024; Model selected: SYM + I + G); for *rpS4-trnT-trnL* dataset: A-C = 0.4777, A-G = 0.6338, A-T = 0.0968,

C-G = 0.6238, C-T = 1.1010, G-T = 1.0000 (Proportion of invariable sites [I] = 0.5280; Gamma distribution shape parameter = 0.9553; Model selected: GTR + I + G); for ITS + *rpS4-trnT-trnL* combined dataset: A-C = 0.5621, A-G = 1.2123, A-T = 0.3214, C-G = 0.3810, C-T = 5.0583, G-T = 1.0000 (Proportion of invariable sites [I] = 0.5978; Gamma distribution shape parameter = 0.5788; Model selected: GTR + I + G) and Rates = gamma.

The Markov Chain Monte Carlo process was set so that four chains ran simultaneously for 2,000,000 generations, with trees being sampled every 100 generations for a total of 20,000 trees in the initial sample. As “stationarity” was achieved by the 3,000th tree, the first 3,000 trees were discarded and the posterior probability of the phylogeny and its branches was determined from the remaining 17,000 trees. Internodes with posterior probabilities $\geq 95\%$ were considered to be well-supported.

Parsimony and Maximum Likelihood. – For the combined dataset (138 taxa), a parsimony analysis and a maximum likelihood analysis (ML) were performed with PAUP Version 4.0b10 (Swofford, 2002). For the parsimony analysis, due to the impracticability of performing standard analyses on our large dataset, we used the PAUPRat approach (Nixon, 1999; Sikes & Lewis, 2001), a tool for implementing Parsimony Ratchet searches using PAUP. Rather than performing many thorough and independent heuristic searches, the Ratchet performs what amounts to a single long search comprising a series of short heuristic searches. We have performed 200 iterations with TBR branch swapping. All most parsimonious trees (MPT) were saved and PAUP was used for computing a strict consensus tree. For the strict consensus tree, consistency index (CI), retention index (RI) and homoplasy index (HI), all excluding uninformative characters, have been calculated.

A bootstrap analysis (BS) (Felsenstein, 1985) was done to obtain support estimates for the nodes in the strict consensus tree. The analysis was performed using the approach by Lidén & al. (1997) using 1,000 replicates, random taxon addition with 10 replicates per replicate and no branch swapping.

For the ML analysis, we started with a tree generated by Neighbor Joining (NJ) and used the SYM + G (Zharkikh, 1994) evolutionary model which best fits the data according to MrModeltest 2.2 (Nylander, 2004). Because of the large size of our dataset we accelerated the analysis using a ratchet approach according to the following parameters suggested by Morrison (2007): SPR branch swapping, re-weight 25% of the characters, 10 iterations of re-weighting, ApproxLim no more than 2% which indicates that approximate scores within 2% of the optimum will be evaluated fully, and MULTREES inactivated.

Partition homogeneity test. – In order to investigate whether the ITS and *rpS4-trnT-trnL* datasets can be combined, a partition homogeneity test (Farris & al., 1994;

implemented in PAUP* 4.0b10; Swofford, 2002) with 10 homogeneity replicates, 10 random addition sequences, tree-bisection-reconnection (TBR) branch swapping on best only and MULTREES on was performed. In order to explore the congruency of the two datasets further, the congruency of individual clades with ≥ 0.95 PP found in the ITS and *rpS4-trnT-trnL* datasets was compared.

Molecular clock analysis. – A likelihood ratio test (Felsenstein, 1988) was performed on the tree resulting from the ML analysis (1,000 random addition sequence, TBR and MULTREES on, after Zhang & al., 2007) of the ITS dataset. Modeltest 3.5 (Posada & Crandall, 1998) was used to perform the likelihood ratio test comparing the log-likelihood scores of the tree with and without enforcing a molecular clock. Degrees of freedom were equal to $s - 2$ where s equals the number of taxa in the analysis. A significant difference at the 0.01 level between two scores was observed. Following this, PAUP* 4.0b10 (Swofford, 2002) was used to calculate genetic distances among taxa. After eliminating all taxa with large genetic distances to *Saussurea elegans*, a clock-like tree with 112 taxa was obtained.

As we could not identify appropriate fossils of the study group to calibrate our molecular clock tree (but see below), published ITS substitution rates were used for calibration. According to Kay & al. (2006), ITS substitution rates in herbaceous angiosperms vary between 1.72×10^{-9} to 8.34×10^{-9} (mean = 4.13×10^{-9}) substitutions/site/year. The approximate age of seven nodes of our molecular clock tree was determined using the above three rates.

RESULTS

Partition homogeneity test. — No significant incongruence ($P = 0.01$) between the two datasets was detected in the partition homogeneity test.

The clades with ≥ 0.95 PP found in the ITS and *rpS4-trnT-trnL* trees, respectively, in the majority of cases are not congruent among the trees (Figs. 2, 3; Appendix 2). Thus, of the seven clades (or nested subclades) found in the *rpS4-trnT-trnL* tree, only one (clade 1) is found in the ITS tree (clade 8, containing one additional species). The remaining six clades are not present in the ITS tree, and the constituent species are separated by at least one branch with ≥ 0.95 PP. Equally, of the 28 clades found in the ITS tree, only three clades (clades 2, 9, 17) plus two species of one clade (clade 8) are recovered as clades with ≥ 0.95 PP by the *rpS4-trnT-trnL* data. The third species of clade 8 of the ITS tree, *C. gmelinii* C. Winkl., is separated from the other two species by several branches in the *rpS4-trnT-trnL* tree although none of these has ≥ 0.95 PP. The remaining 24 ITS clades are not present in the *rpS4-trnT-trnL* tree, and in 13 of these clades the species constituting

the ITS clades are separated by at least one branch with ≥ 0.95 PP in the *rpS4-trnT-trnL* topology. Considering the results of this clade-by-clade comparison, the finding of congruency between the two datasets by the partition homogeneity test is somewhat surprising.

Phylogenetic analysis. — The numerical results of the analyses of the ITS, *rpS4-trnT-trnL* and combined ITS + *rpS4-trnT-trnL* datasets are given in Table 1. The individual datasets were analyzed with Bayesian inference (BI), and the combined dataset with Maximum Parsimony (MP), Bayesian inference (BI) and Maximum Likelihood

(ML). The results of the analyses of all three datasets with BI are shown in Figs. 2–4, the result of the MP analysis is partially shown in Fig. 6 (only the Arctioid clade) and the results of the ML analysis of the combined dataset are not shown.

All analyses confirmed the monophyly of the *Arctium-Cousinia* complex. Support for this was strong in the MP and BI analyses (BS = 89%, PP = 1.00). Except for the BI analysis of *rpS4-trnT-trnL* alone, all analyses also show a clear subdivision of the complex into three groups: an Arctioid clade moderately supported by BS but well

Table 1. Numerical results of the analyses of the ITS, *rpS4-trnT-trnL* and combined ITS + *rpS4-trnT-trnL* datasets.

	ITS				<i>rpS4-trnT-trnL</i>				Combined ITS + <i>rpS4-trnT-trnL</i>			
	tree	–out.	Cous.	Arct.	tree	–out.	Cous.	Arct.	tree	–out.	Cous.	Arct.
Total characters	489				1,109				1,598			
Inf.	172	143	143	51	28	21	21	7	216	171	140	58
Tree length	799	680	546	85	78	55	38	11	980	836	679	105
CI	0.308	0.290	0.300	0.647	0.603	0.545	0.579	0.636	0.299	0.272	0.274	0.590
RI	0.721	0.728	0.624	0.886	0.922	0.931	0.909	0.810	0.711	0.715	0.586	0.849
HI	0.692	0.710	0.700	0.353	0.397	0.455	0.421	0.364	0.701	0.728	0.726	0.410

Tree length, CI, RI and HI were calculated for entire trees, for trees without outgroups and for the Arctioid and the Cousinioid clades. Abbreviations: Arct., Arctioid clade; CI, consistency index; Cous., Cousinioid clade; HI, homoplasy index; Inf., phylogenetically informative characters; –out., outgroups excluded; RI, retention index; tree, the entire tree including outgroups.

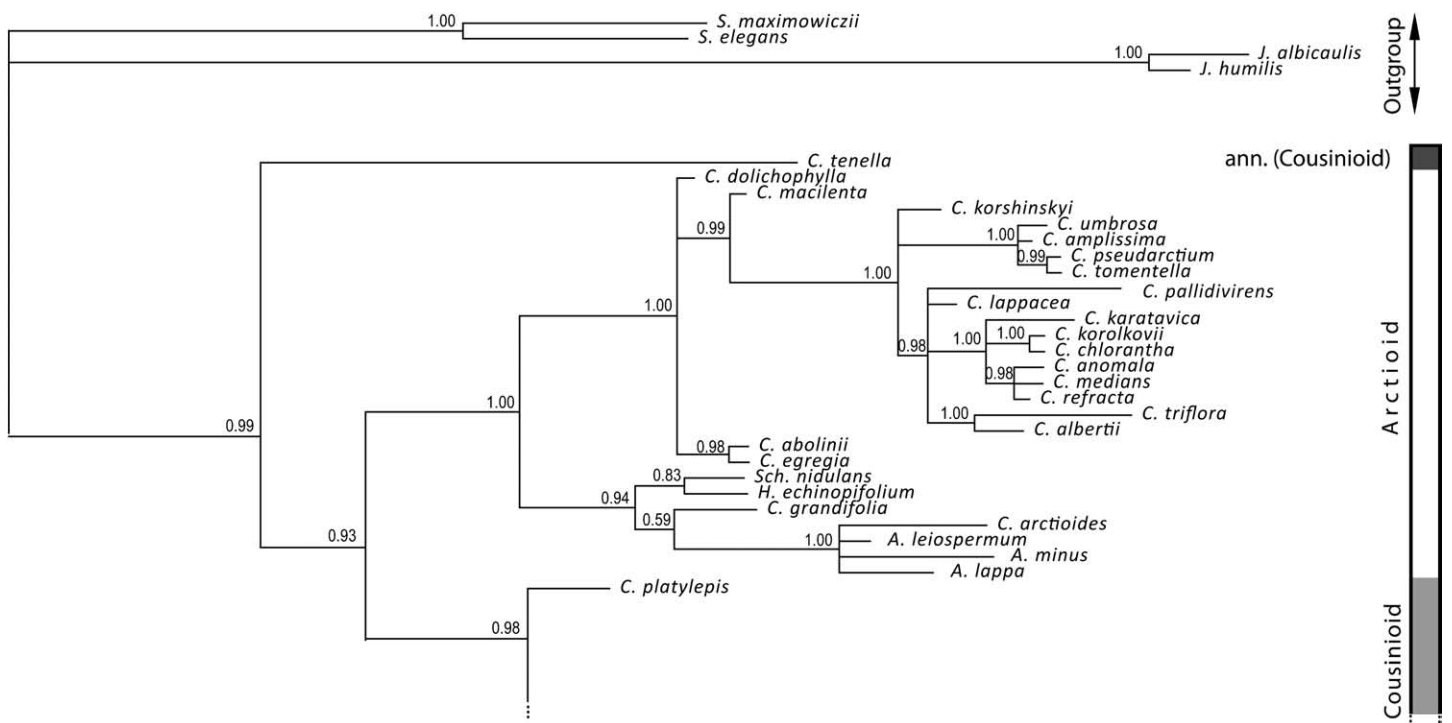
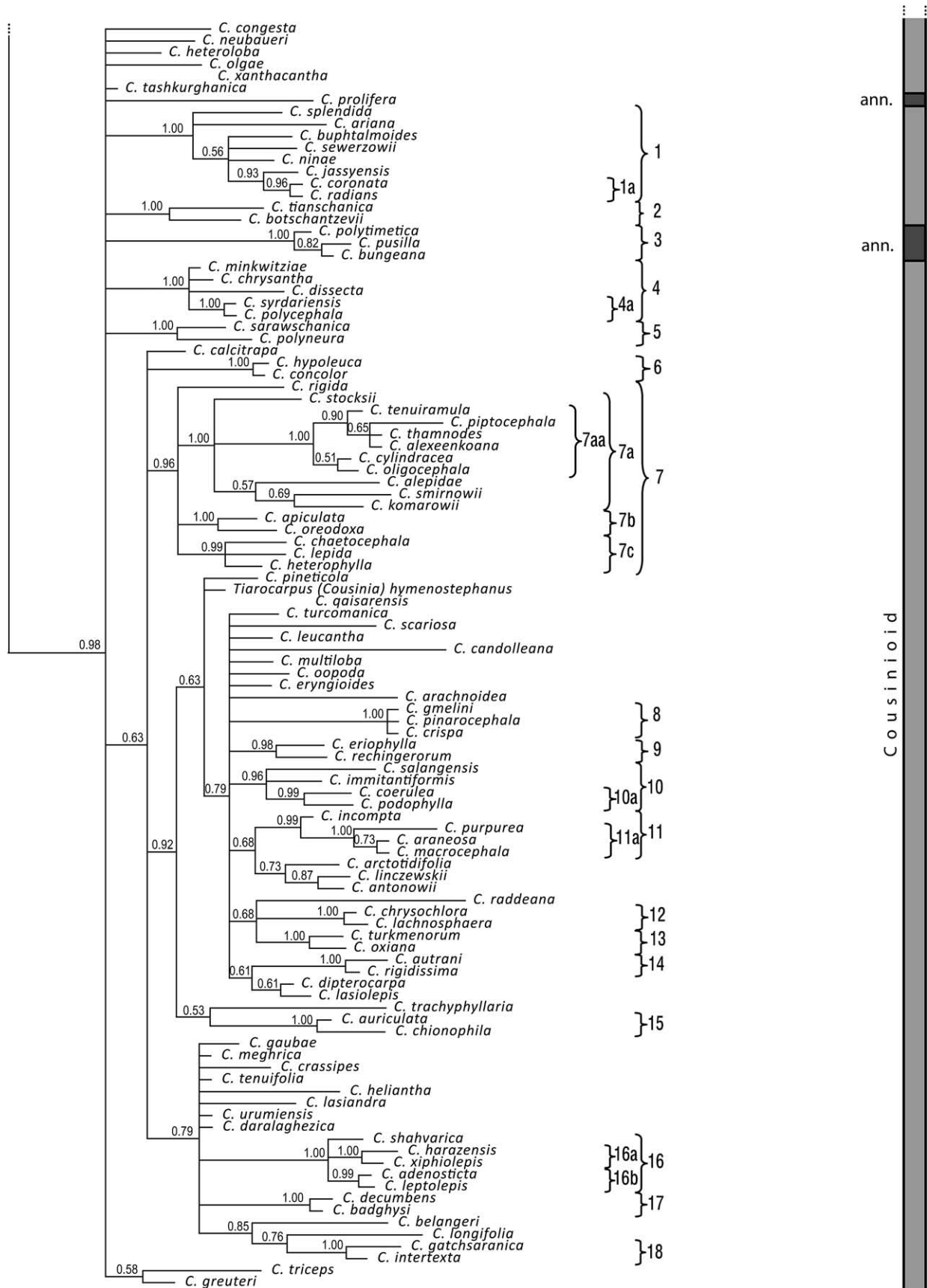


Fig. 2. 50% majority-rule consensus tree obtained from the Bayesian analysis of the ITS dataset. Numbers on branches are posterior probabilities. ann. = annual species. Numbers at curly brackets indicate clades with >0.95 PP discussed in the text.



supported by PP (BS = 72%, PP = 1.00) which contains *Arctium*, *Cousinia* subgg. *Cynaroides* and *Hypacanthodes*, *Hypacanthium* and *Schmalhausenia*, a Cousinioid clade with BS < 50% but with strong Bayesian support (PP = 1.00) which contains most species of *Cousinia* subg. *Cousinia*, and *C. tenella* Fisch. & C.A. Mey. as sister to the above two clades (BS = 54%, PP = 0.89). In the BI analysis of *rpS4-trnT-trnL*, *C. tenella* falls into the Arctioid clade with only low PP.

The Arctioid clade. – The description of the Arctioid clade is based on the tree obtained from the MP analysis of the combined dataset (Fig. 6). This tree is essentially congruent with the BI tree but better suited for character mapping. Within the Arctioid clade relationships among species are reasonably well resolved and this clade is subdivided into two subclades (Fig. 6). The first subclade, with moderate BS (80%) but strong PP support (1.00) is formed by most species of *Cousinia* subgg. *Cynaroides* and *Hypacanthodes*. This subclade consists of an unresolved polytomy containing four species of subg. *Hypacanthodes* (*C. egregia* Juz., *C. dolichophylla* Kult.,

C. abolinii Kult. ex Tscherneva, *C. macilenta* Winkl.) in two lineages plus a well-supported clade (BS = 90%, PP = 1.00) formed by most of species of subg. *Cynaroides* which in turn fall into two unsupported subgroups. The second subclade also has good support (BS = 81%, PP = 1.00) and is formed by *Arctium* together with *C. arctioides* Schrenk of subg. *Cynaroides* (BS = 92%, PP = 1.00), *Cousinia grandifolia* Kult. of subg. *Hypacanthodes*, *Schmalhausenia* and *Hypacanthium*. Relationships in the Arctioid clade as described above are essentially similar in the BI analyses of the ITS and combined dataset and in the ML analysis of the combined dataset.

The Cousinioid clade. – Relationships among species in the Cousinioid clade are essentially unresolved in the MP analysis of the combined dataset. Resolution is better in the BI analyses of the ITS and combined datasets and will be further discussed below (see Discussion).

Molecular clock analysis. — The results of our molecular clock analysis are shown in Table 2 and Fig. 5. Mai (2001) reported fossil *Arctium* achenes from a German Miocene locality. The age of this fossil can be dated

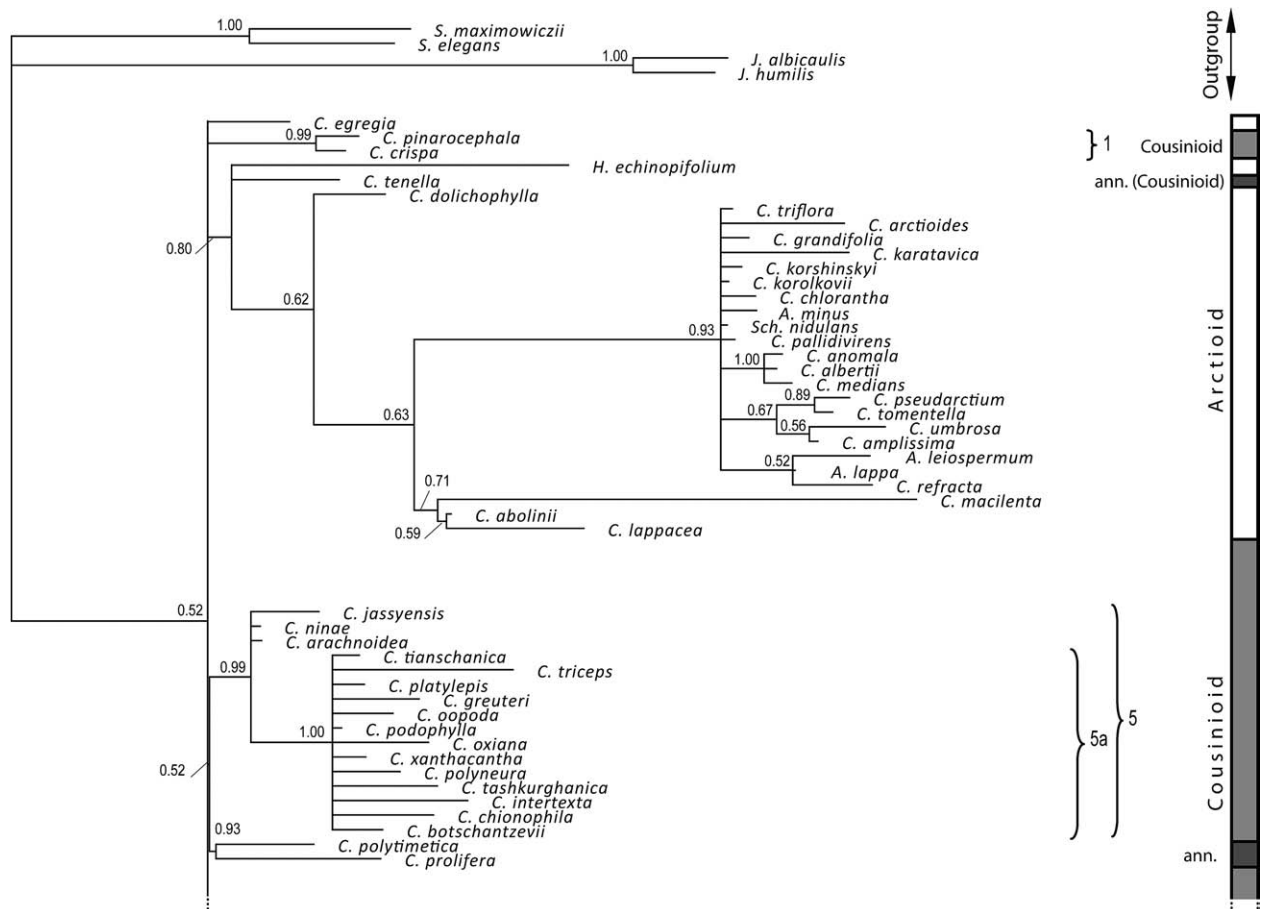


Fig. 3. 50% majority-rule consensus tree obtained from the Bayesian analysis of the *rpS4-trnT-trnL* dataset. Numbers on branches are posterior probabilities. ann. = annual species. Numbers at curly brackets indicate clades with >0.95 PP discussed in the text.

to 7–9 mya (V. Wähner, pers. comm.). Considering that the achenes of *Arctium* can not be reliably distinguished from those of the other genera of the *Arctium-Cousinia* complex, this fossil essentially is compatible with the ages calculated for the complex with all three ITS substitution rates. When considering the occurrence of this achene in Germany, and assuming that the geographical distribution of the complex in the past was similar to its distribution today, with *Arctium* being the only genus occurring in

Europe, the fossil probably is best assigned to *Arctium*. This would imply that of the three ITS substitution rates used for the calibration of the molecular clock, the mean rate fits best. With this rate, the clade containing *Arctium* (node F in Fig. 5) was dated to ca. 6.3 my (crown group age). Based on the above considerations, we will refer mainly to ages obtained when using the mean ITS substitution rate in the following discussion, and these ages are also indicated in Fig. 5.



DISCUSSION

The *Arctium-Cousinia* complex. — As already found by previous authors (Susanna & al., 2003a; Susanna & Garcia-Jacas, 2007), the *Arctium-Cousinia* complex clearly is monophyletic with good support in the MP and BI analyses (BS = 89%, PP = 1.00). In all analyses except the BI analysis of *rpS4-trnT-trnL* a division into three groups, the Arctioid clade, the Cousinioid clade and *Cousinia tenella* was found.

When comparing the Arctioid and Cousinioid clades, two major differences can be observed. First, whereas intraclade relationships are reasonably well-resolved in the Arctioid clade, very little resolution was obtained in the Cousinioid clade. Considering that the two clades are sister to each other, and that crown group diversification of the two clades started at approximately the same time (mean rate: Arctioid clade: ca. 9.7 mya, Cousinioid clade: ca. 8.7 mya; Table 2), differential age of the two groups can not

Table 2. Absolute ages in million years (my) obtained from the molecular clock analysis.

	Rate 1 (my)	Mean rate (my)	Rate 2 (my)
Node A	45.930	19.128	9.472
Node B	35.465	14.770	7.314
Node C	23.256	9.685	4.796
Node D	20.930	8.717	4.317
Node E	16.860	7.022	3.477
Node F	15.116	6.295	3.118
Node G	10.465	4.358	2.158

Rate 1 = 1.72×10^{-9} substitutions/site/year; mean rate = 4.13×10^{-9} substitutions/site/year; rate 2 = 8.34×10^{-9} substitutions/site/year (Kay & al., 2006).

be advocated as explanation for the difference in phylogenetic resolution. Instead, we believe that the striking difference in species number is responsible for the difference

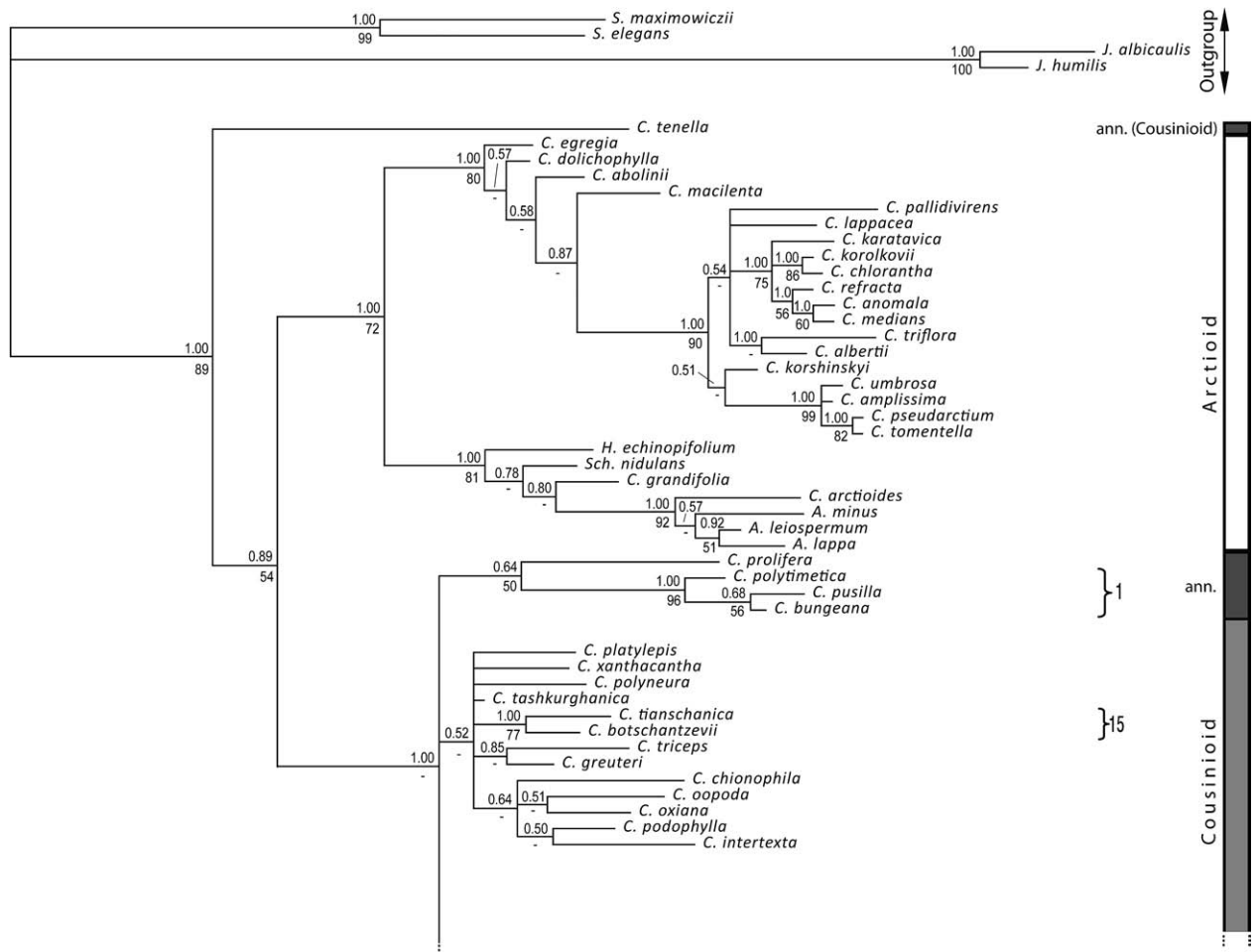
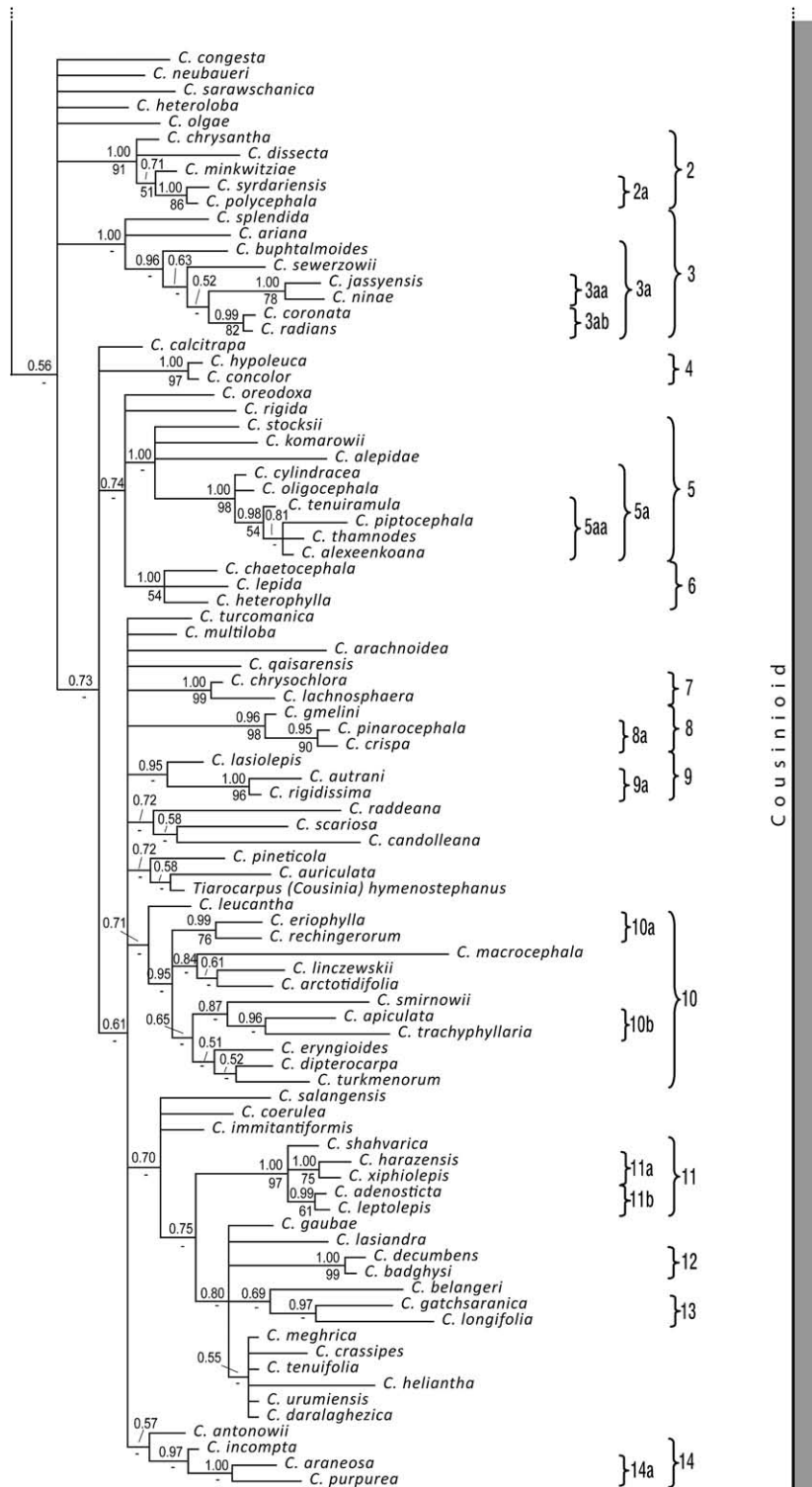


Fig. 4. 50% majority-rule consensus tree obtained from the Bayesian analysis of the combined ITS and *rpS4-trnT-trnL* dataset. Numbers above branches are posterior probabilities and numbers below branches are bootstrap percentages. ann. = annual species. Numbers at curly brackets indicate clades with >0.95 PP discussed in the text.

in phylogenetic resolution. If the Cousinioid clade were to be reduced to a random sample of species equal in number to that of the Arctioid clade, phylogenetic resolution probably would be greatly improved. Indeed, this prediction is to some extent confirmed by the data of Susanna & al.

(2003a), who obtained some phylogenetic structure among the 21 species of the Cousinioid clade included in their analysis.

Second, presumably plesiomorphic and apomorphic characters are distributed unevenly across the two clades.



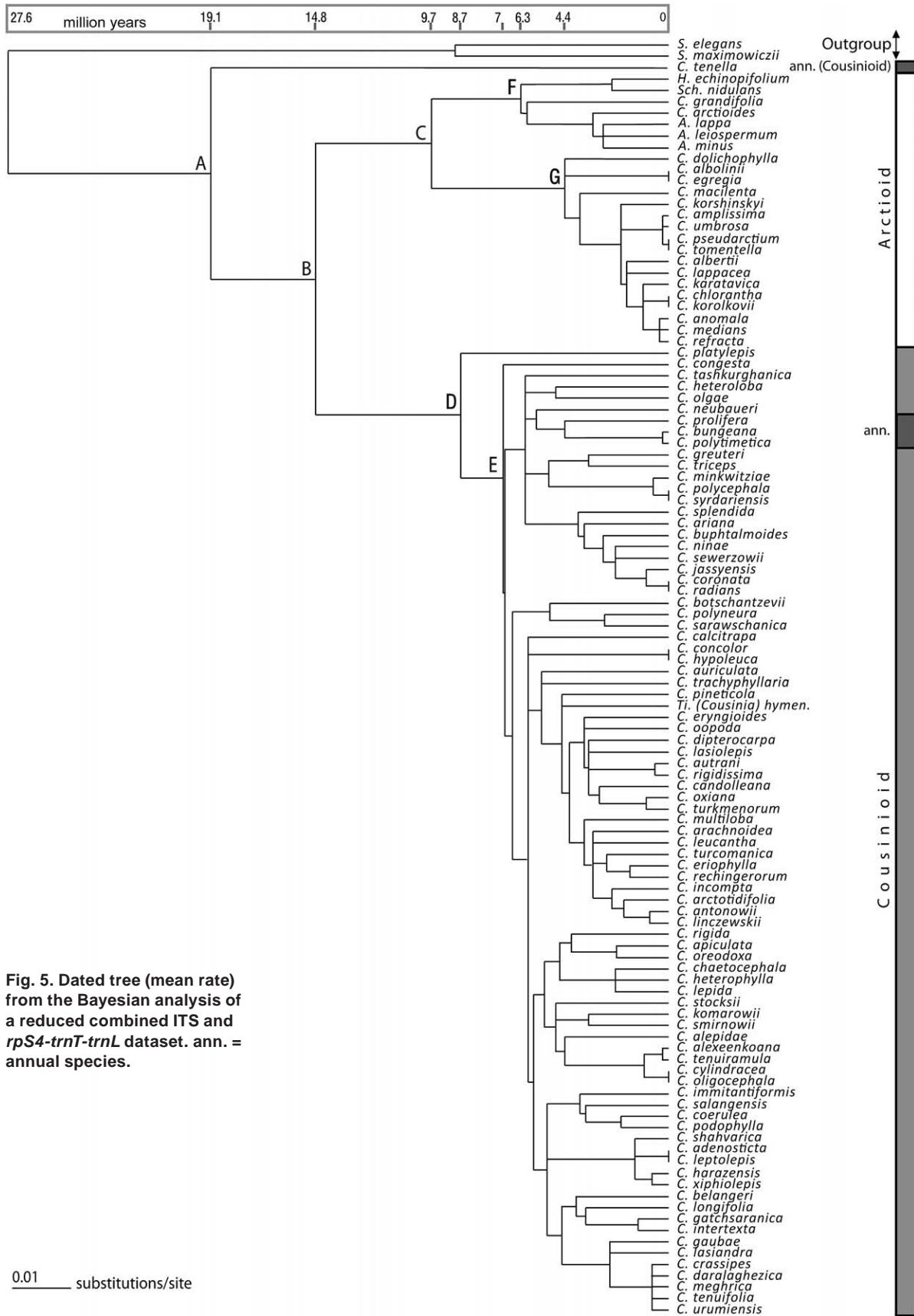


Fig. 5. Dated tree (mean rate) from the Bayesian analysis of a reduced combined ITS and *rpS4-trnT-trnL* dataset. ann. = annual species.

Whereas the spiny pollen (and possibly also long stigmata) of the Arctioid clade can also be found in *Jurinea* and *Saussurea* (Punt & Hoen, 2009) as closest relatives of the *Arctium-Cousinia* complex, the short stigmata and smooth pollen of the Cousinioid clade appear to be apomorphic. In other groups of Cardueae, pollen morphology is associated with the pollen presentation mechanism. In Centaureinae (García-Jacas & al., 2001; Vilatersana & al., 2001), basal groups have spiny pollen and long, usually dry stigmata and lack a touch-sensitive pollen presentation mechanism. More derived groups have very short, sticky stigmata and smooth pollen, and this is associated with touch-sensitive pollen presentation. We hypothesize that a similar transition of characters has taken place in the *Arctium-Cousinia* complex. As regards chromosome numbers, the Arctioid clade is rather stable with a base chromosome number of $x = 18$. Much in contrast to this, considerable descending dysploidy ($x = 9, 10, 11, 12, 13$) is found in the Cousinioid clade. Of the closest relatives of the complex, *Jurinea* has a base chromosome number of $x = 17$ and *Saussurea* of $x = 13$. In general, basal Carduinae have higher chromosome

numbers. The distribution of chromosome numbers in comparison of the Arctioid and Cousinioid clades again is paralleled in subtribe Centaureinae where an evolutionary trend from high to low chromosome numbers has been reported (García-Jacas & al., 2001). Finally, it could be argued that the Arctioid and Cousinioid clades differ in habitat diversity. Whereas species of the Arctioid clade, with the obvious exception of the widespread colonizers mainly of *Arctium*, are more or less confined to high mountain habitats, species of the Cousinioid clade have diversified and grow in widely different habitats including semidesert areas, open hilly areas, alpine ecosystems, high altitude areas transitional between humid and semiarid ecosystems, oak forest floors and rarely in lowlands. Possibly the differential species richness of the Arctioid and Cousinioid clades on the one hand and the unequal distribution of plesiomorphic vs. apomorphic characters among the two clades are not only correlated but causally linked. However, whether transition to apomorphic characters in the Cousinioid clade is cause or effect of its higher species number can not be easily decided with the data available.

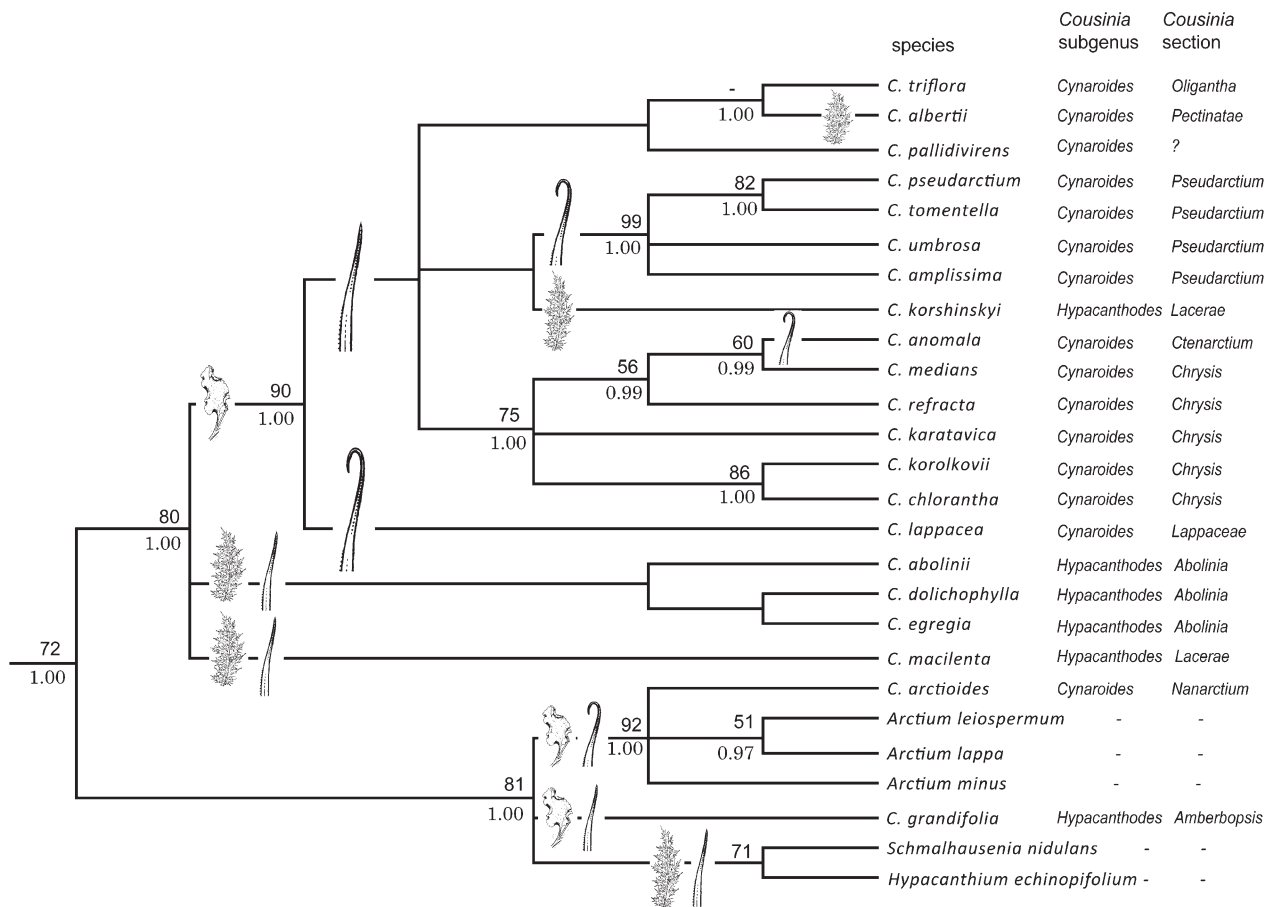


Fig. 6. Distribution of leaf and involucral bract morphology among species of the Arctioid clade mapped on an MP tree of the combined ITS and *rpS4-trnT-trnL* dataset.

The Arctioid clade. — As described in the introduction, the Arctioid clade can not be characterized morphologically beyond pollen and style morphology (neither chromosome number). Part of it (*Arctium* and four species of *Cousinia* subg. *Cynaroides*) have unarmed, large cordate leaves and involucre bracts ending in a recurved hook. Of the remainder of subg. *Cynaroides*, six species have leaves which are not cordate but involucre bracts ending in a hooked spine, and twelve species have involucre bracts ending in a straight spine. The distribution of these two characters precludes the morphological definition of a monophyletic *Arctium*. In subg. *Hypacanthodes* no species has either cordate leaves or hooked involucre bracts. The distribution of spiny leaves also runs across established generic and subgeneric boundaries. All species of *Arctium* and almost all species of *Cousinia* subg. *Cynaroides* have unarmed leaves, and only *C. alberti* Regel & Schmalh. and *C. horrescens* Juz. are spiny. In contrast to this, *Schmalhausenia*, *Hypacanthium* and *Cousinia* subg. *Hypacanthodes* (except *C. grandifolia*) have strongly spiny leaves.

A logical solution to the above character distribution would be to establish a broadly redefined *Schmalhausenia* including *Hypacanthium* and the spiny species of *Cousinia* subg. *Hypacanthodes*, and a broadly redefined *Arctium* including subg. *Cynaroides* and the unarmed *C. grandifolia* from *Cousinia* subg. *Hypacanthodes*. However, this solution clearly is not compatible with molecular evidence presented here. In fact, the tree topology (Fig. 6) suggests the recognition of two to four groups: (1) *Schmalhausenia* plus *Hypacanthium*, (2) *Cousinia grandifolia*, (3) *Arctium* incl. *Cousinia arctioides* and (4) the remaining species of *Cousinia* subg. *Cynaroides* and *Hypacanthodes*. However, this solution is not compatible with morphological evidence. In conclusion, our declared aim to revise generic limits in the Arctioid clade on the basis of a larger taxon sample could not be achieved because of the molecular placement and morphology of the species of *Cousinia* subg. *Cynaroides* and *Hypacanthodes*. When considering the distribution of leaf and involucre bract morphology across the phylogeny of the entire complex (Fig. 6), it is obvious that either cordate and unarmed or spiny leaves and apically hooked or straight involucre bracts—these two characters are not always correlated—evolved several times in parallel. In view of the uncertainties in our phylogeny we did not try to formally optimize the evolution of these characters.

Interestingly, the only colonizing species of the *Arctium-Cousinia* group, i.e., most species of *Arctium*, have capitula with hooked involucre bracts. Such capitula are dispersed epizoochorously and may be direct cause of the colonizing success of the genus. However, some species with hooked involucre bracts are geographically limited to the Tien-Shan. It seems possible that the biennial habit of *Arctium*—most other species of the Arctioid clade are

perennial—contributed to this colonizing success as already suspected by Duistermaat (1996).

The sectional classification of *Cousinia* subg. *Cynaroides* and subg. *Hypacanthodes* is only partly supported by our results, and neither subgenus is monophyletic as presently circumscribed (Fig. 6). One species of subg. *Cynaroides*, *C. arctioides*, is placed within the *Arctium* clade with good parsimony and Bayesian support (BS = 92%, PP = 1.00). This position is fully congruent with morphology as *C. arctioides* has unarmed leaves as well as involucre bracts ending in a recurved hook. We propose to follow the taxonomical criterion of Kuntze (1891) and Duistermaat (1996) which consider this species as *Arctium arctioides* (Schrenk) Kuntze.

Apart from this, our phylogeny is largely congruent with the current sectional classification of subg. *Cynaroides*. The species of sect. *Chrysis* (*C. medians* Juz. through *C. chlorantha*) form a clade with low BS (75%) but high PP (1.0) support. However, this clade also includes *C. anomala* Franch. of sect. *Ctenarctium*. This grouping is not supported by morphological characters as *C. anomala* has involucre bracts ending in a hook and oblong-lanceolate leaves. Section *Pseudarctium* (*C. pseudarctium* Bornm. through *C. amplissima* Boiss.) is a strongly supported clade (BS = 99%, PP = 1.00) containing species with involucre bracts with hooked spines and large, soft and slightly pubescent leaves. Sections *Oligantha* and *Pectinatae* group together with BS < 50% but strong BI support (PP = 1.00). This grouping is not supported by any morphological characters and may result from the poor representation of sect. *Pectinatae* in our sample.

Two species of subg. *Hypacanthodes* were resolved in unexpected position. Thus, *C. korshinskyi* Winkl. of sect. *Lacerae* is part of the subg. *Cynaroides* clade discussed above and separated from the other species of sect. *Lacerae* included in the analysis (*C. macilentata*). This grouping is incongruent with morphology because *C. korshinskyi* is more similar to the rest of subg. *Hypacanthodes* than to subg. *Cynaroides*. *Cousinia grandifolia* is part of the clade containing *Arctium*, *Schmalhausenia* and *Hypacanthium*. In this case, this placement is well correlated with morphology. *Cousinia grandifolia* shares the possession of large, unarmed, oblong-ovate leaves with *Arctium*, and involucre bracts ending in a spine without hook with *Schmalhausenia* and *Hypacanthium*. With these characters *C. grandifolia* is somewhat intermediate between *Arctium* on the one hand and *Schmalhausenia* and *Hypacanthium* on the other hand.

The sectional classification of subg. *Hypacanthodes* is not confirmed by our data. Although the species of sect. *Abolinia* form a clade which is homogeneous morphologically, this clade has no statistical support. *Cousinia macilentata* and *C. korshinskyi* of sect. *Lacerae* are placed in two different and strongly supported clades (see above).

They had been placed in separate series of sect. *Lacerae*, *Macilentae* and *Pterolepideae*, by Tscherneva (1962) and differ in pubescence, height and branching of the stem, leaf size, capitulum pubescence, shape and texture of the involucre bracts and achene ornamentation.

The Cousinioid clade. — As evident from our Bayesian analyses of the ITS, *rpS4-trnT-trnL* and combined datasets (Figs. 2–4), and also from the MP and ML analyses of the combined dataset (not shown), very little phylogenetic resolution can be obtained within *Cousinia* subg. *Cousinia*. Although a varying number of mostly small species groups with high support is recognizable in the different datasets, relationships among species groups are essentially unresolved.

In the following discussion, we first will explore in detail the phylogenetic information contents of our data. This we will do by examining whether clades with ≥ 0.95 PP are plausible in terms of traditional taxonomic assignment (as a proxy for morphological similarity) and/or geographical distribution by growing in the same general area. For this purpose, the following eight major centres of species diversity were defined (Fig. 1): Anatolian mountains, Zagros mountains, Elburz mountains, Kopet Dag and mountains to the south, West Hindu Kush (C & W Afghanistan), East Hindu Kush (NE Afghanistan), Pamir-Alay and Tien-Shan (Rechinger, 1986; Knapp, 1987). Second, we will investigate possible causes for the very limited amount of phylogenetic resolution obtained.

Phylogenetic information in the rpS4-trnT-trnL, ITS and combined datasets. — In the *rpS4-trnT-trnL* dataset, seven clades (or nested subclades) with ≥ 0.95 PP were obtained (Appendix 2). Of these, clades 3 and 4 are not plausible in terms of traditional taxonomic assignment or geographical distribution. Clade 1 is plausible in both terms, and clades 2–2a and 5–5a, with few exceptions (2–2a: *C. araneosa* DC., *C. macrocephala* C.A. Mey., *C. gatchsaranica* Mehregan, Assadi & Attar; 5–5a: *C. oxiana* Tscherneva, *C. intertexta* Freyn & Sint.), contain species from the Kopet Dag area (S Turkmenistan, NE & E Iran and NW Afghanistan) and Central Asia, respectively, but of different sections of *Cousinia* subg. *Cousinia*. In the ITS dataset, 28 clades (or nested subclades) with ≥ 0.95 PP were obtained (Appendix 2). Of these, 16 are plausible both in terms of taxonomic assignment and geographical distribution, 2 are plausible in terms of taxonomic assignment only, 4 are plausible in terms of geographical distribution only, and the remaining 6 clades are not plausible in either terms. In the combined dataset, 28 clades (or nested subclades) with ≥ 0.95 PP were obtained (Appendix 2). Of these, 13 are plausible both in terms of taxonomic assignment and geographical distribution, 4 are plausible in terms of taxonomic assignment only, 4 in terms of geographical distribution only, and the remaining 7 clades are not plausible in either terms.

To summarize the above considerations, our datasets do appear to contain some plausible phylogenetic information mainly in their distal branches. When considering terms of clade plausibility, it also appears that geographical distribution is a more reliable criterion for recognizing phylogenetic relatedness than traditional taxonomic assignment as also found in other studies (e.g., González-Albaladejo & al., 2005).

Most interestingly, and in support of the above conclusion, the annual species (except *C. pygmaea* C. Winkl.—which does not appear in the trees shown—and *C. tenella* which groups outside *Cousinia* subg. *Cousinia* in all our analyses, for discussion see below) group in one clade in the majority of our analyses. In the ITS dataset (Fig. 2), *C. bungeana*, *C. polytimetica* and *C. pusilla* form one clade with 1.0 PP which is part of a polytomy also containing *C. prolifera*. In the *rpS4-trnT-trnL* dataset (Fig. 3), *C. prolifera* Jaub. & Spach and *C. polytimetica* Tscherneva form one clade (0.93 PP) widely separated (although by no branch with ≥ 0.95 PP) from *C. pusilla* C. Winkl. and *C. bungeana* Regel & Schmalh. which are part of a large polytomy. In the Bayesian combined dataset (Fig. 4) these four annual species form one clade with 0.64 PP, of which *C. bungeana*, *C. polytimetica* and *C. pusilla* form a subclade with 1.0 PP. These four annual species are also recovered as a monophyletic group in the MP analysis of the combined dataset (50% BS; *C. bungeana*, *C. polytimetica*, *C. pusilla*: 96% BS; not shown), and *C. bungeana*, *C. polytimetica* and *C. pusilla* are recovered in the ML analysis of the combined dataset (not shown). The monophyly of the annual species (except *C. pygmaea* and *C. tenella*) in the combined dataset even more than the above considerations implies that our ITS and *rpS4-trnT-trnL* sequence data do contain limited but valuable phylogenetic information.

Possible causes for lack of phylogenetic resolution in Cousinia subg. Cousinia. — The first obvious reason for the lack of phylogenetic resolution obtained in the analyses of the ITS, *rpS4-trnT-trnL* and combined datasets is the small number of characters available in relation to the number of taxa analysed. For the 106 species included of *Cousinia* subg. *Cousinia*, the *rpS4-trnT-trnL* dataset yielded 21, the ITS dataset 143 and the combined dataset 140 parsimony informative characters (Table 1). Although in theory the number of parsimony informative characters in the ITS and combined datasets would be sufficient to resolve a phylogeny of 106 species, they are not, because the characters show considerable homoplasy. Although the RI for the *rpS4-trnT-trnL* dataset is high (0.922), it is fairly low for the ITS (0.721) and combined datasets (0.711). The most likely reason for the lack of characters in *Cousinia* subg. *Cousinia* is its comparatively young age (Table 2, Fig. 5) in combination with its high species number. Thus, using the average substitution rate for ITS for herbaceous perennials (Kay & al., 2006) as to some

extend justified by the age of the fossil *Arctium* achene by Mai (2001; see above), a crown group age of ca. 8.7 my (node D) was found for *Cousinia* subg. *Cousinia*, and its major diversification started only ca. 7 mya (node E).

Considering the existence of a large number of closely related and often sympatric species with no obvious mechanism of reproductive isolation in *Cousinia* subg. *Cousinia*, hybridization and hybrid speciation seems possible and clearly would confound the reconstruction of phylogeny (Niето-Feliner & al., 2001; Font & al., 2002; Vriesendorp & Bakker, 2005; Suárez-Santiago & al., 2007). Based on the examination of literature reports of hybrids and intermediate forms and the morphological and molecular examination of some suspected hybrids, Mehregan & Kadereit (in press) concluded that although hybridization does occur in *Cousinia*, it seems to be comparatively rare in the genus. Thus, only 26 hybrids and 12 intermediate forms in *Cousinia* subg. *Cousinia* have been published (Tscherneva 1962; Rechinger, 1972, 1979). These hybrids and intermediate forms involve ca. 11% of the currently recognized species of *Cousinia* subg. *Cousinia*. The two hybrids examined in detail (*C. shahvarica* × *C. decumbens*, *C. eryngioides* × *C. chrysochlora*), confirmed by additivity of their ITS sequences, showed reduced pollen fertility. It is possible that hybridization is much more frequent but undetected because of lack of sufficiently detailed taxonomic knowledge and the existence of several morphologically very similar species. However, the complete absence of polyploid chromosome numbers in *Cousinia* subg. *Cousinia* among the 148 species analyzed (<http://www-asteraceae.cla.kobe-u.ac.jp/search.html>) implies that polyploid hybrid speciation played no role in the evolution of *Cousinia* subg. *Cousinia*. Nothing can be said about the evolutionary significance of homoploid hybrid speciation in this group.

In summary, there is little evidence that hybridization and/or hybrid speciation are important factors hindering the reconstruction of the phylogeny of *Cousinia* subg. *Cousinia*.

The phylogeny and evolution of Cousinia subg. Cousinia. – As evident from the above discussion, very little can be said about the phylogeny of *Cousinia* subg. *Cousinia*. Although a number of species groups can be recognized which may well represent monophyletic lineages as judged from their taxonomic assignment and/or geographical distribution, relationships among species groups are entirely unclear and no major subdivision of *Cousinia* subg. *Cousinia* is evident from our data. This is most likely the result of paucity of phylogenetic informative characters contained in our ITS and *rpS4-trnT-trnL* datasets in relation to species number. This in turn is the result of the comparatively young geological age of the group in combination with its species richness. Apart from the recovery of a number of probably monophyletic species groups, our data contain

valuable information about the evolution of annuality, a very rare condition in *Cousinia* subg. *Cousinia*. It appears that annuality originated twice in *Cousinia* subg. *Cousinia*. The first lineage of annuals, comprising *C. bungeana*, *C. polytimetica*, *C. pusilla* and *C. prolifera* has been discussed above and is well-nested within *Cousinia* subg. *Cousinia*. The second lineage comprises *C. tenella* and *C. pygmaea* and will be discussed further below.

With its comparatively young age and high species number *Cousinia* subg. *Cousinia* represents a massive radiation which is quite unusual because it took place in a comparatively small geographical area. Based on the results of an ongoing taxonomic revision of part of *Cousinia* subg. *Cousinia* (sect. *Cynaroideae*; Mehregan & Kadereit, 2008), mainly two conclusions can be drawn about the evolution of this group. First, it seems very likely that the number of species described (ca. 600) vastly overstates the number of existing species. Thus, our revision reduced the number of species in sect. *Cynaroideae* from 110 to 31. If reduction in species number by two-thirds eventually will also take place in other parts of *Cousinia* subg. *Cousinia*, it would still remain a large genus with about 200 species. Second, of the ca. 30 species recognized by Mehregan & Kadereit (2008), only two are geographically widespread, namely *C. calocephala* Jaub. & Spach and *C. onopordioides* Ledeb. The remaining species have rather narrow and, most importantly, allo- or parapatric geographical ranges where species boundaries are marked by prominent topographical barriers. This may imply that the predominant speciation process in *Cousinia* subg. *Cousinia* may be geographical speciation. Today, the group is found mainly (but not exclusively) in vegetation types classified as ‘Kurdo-Zagrosian and other steppe-forest climaxes’ by Zohary (1973). When this type of vegetation is considered essentially identical with ‘Mediterranean to sub-Mediterranean *Artemisia-Ephedra-Juniperus-Compositae* steppe of mountainous regions’ as defined by Frenzel & al. (1992), the reconstruction of vegetation during the last glacial maximum (20,000 to 18,000 years ago) by Frenzel & al. (1992) suggests that the geographical range of *Cousinia* subg. *Cousinia* in the last glacial maximum was compressed in only the northern parts of its extant range. This in turn implies substantial changes in geographical distribution of species during the Quaternary. Considering the age of the crown group diversification of *Cousinia* subg. *Cousinia* (ca. 8.7 mya; Table 2) and the age of species which in most cases is older than the onset of the Quaternary (Fig. 5), the climatic oscillations of the Quaternary do not appear to have resulted in widespread extinction of the group. Whether these climatic oscillations instead were a stimulus for speciation (through changes in geographical distribution), as found for, e.g., *Primula* L. sect. *Auricula* and *Epimedium* L. (Kadereit & al., 2004; Zhang & al., 2007), we can not say without further analysis.

***Cousinia tenella*.** — In all our analyses except the BI analysis of the *rpS4-trnT-trnL* dataset alone, *C. tenella* is sister to the Arctioid and Cousinioid clades. Support for the sister group relationship between the Arctioid and Cousinioid clades increases (BS = 94%, PP = 1.00; tree not shown) when *C. tenella* is removed from the data matrix.

Although only *C. tenella* was included in the datasets shown here, an enlarged ITS dataset of 216 species (not shown) shows *C. pygmaea* to be sister to *C. tenella* with 0.84 PP. Such relationship is plausible considering the great similarity between the two species which differ only in flower colour, relative length of corolla tube and limb and achene shape (Rechinger, 1972). The placement of *C. tenella* in most of our analyses might suggest segregation of *C. tenella* and *C. pygmaea* from *Cousinia* subg. *Cousinia*. On the basis of karyology, pollen type and stigma morphology, however, *C. tenella* (and probably *C. pygmaea*) clearly falls into the Cousinioid clade. *Cousinia tenella* has $2n = 26$ chromosomes according to Aryavand (1975) and Susanna & al. (2003b), the Cousinioid pollen type and stigma morphology (Susanna, pers. comm.). The fact that this species is resolved as sister to the rest of the *Arctium-Cousinia* complex probably is due to its annuality, autogamy and probably increased mutation rate. There exist other examples for unexpected placements of annuals in tribe Cardueae. In *Echinops* L., annual species together with the annual *Acantholepis* Less. are sister to the rest of the genus (Garnatje, 2005); in *Ptilostemon* Cass., the only annual species, *Ptilostemon stellatus* (L.) Greuter, is sister to the rest of the genus (Vilatersana & al., in prep.); and in subtribe *Centaureinae* the annual *Schischkinia* Iljin and *Stizolophus* Cass. are successive sisters to the rest of the subtribe (García-Jacas & al., 2006). In all these cases, the position of annuals as sister to the remainder of their respective groups is not congruent with morphology. Several authors have pointed out that annual plants often have an elevated rate of molecular evolution compared with perennials, explained by generation time, population size, and speciation rate (Bousquet & al., 1992; Laroche & al., 1997; Andreasen & Baldwin, 2001). If this indeed should be the explanation for the unexpected placement of *C. tenella*, it is interesting to observe that this problem does not appear to exist for the equally annual *C. bungeana*/*C. polytimetica*/*C. pusilla*/*C. prolifera* lineage which clearly falls into the Cousinioid clade as expected from morphology.

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Appendix 1. Species included in the molecular analysis with voucher information and GenBank accession numbers (ITS1; ITS2; *rpS4-trnT-trnL*). For all species of *Cousinia* the subgeneric classification is given. An asterisk (*) indicates sequences previously published.

Hypacanthium echinopifolium (Bornm.) Juz., Kyrgyzstan: Sine loc., *Ijlin s.n.* (LE), AY373746*; AY373713*; EU661125. *Jurinea albicaulis* Bunge, Greece: Makedonia, Halkidiki, between Arnea and Paleokastro, *Susanna 1957 & al.* (BC), AY373747*; AY373714*; EU661092. *Jurinea humilis* DC., Spain: Huesca, Plan, La Barana, *P. Montserrat s.n.* (JACA), L35868*; EU661093. *Saussurea elegans* Ledeb., Kazakhstan: Aksu-Dzabagly reservation, 1,800 m, *Susanna 2179 & al.* (BC), AY373750*; AY373717*; EU661090. *Saussurea maximowiczii* Herder, Japan: Ofuna Botanical Garden, AY373751*; AY373718*; EU661091. *Schmalhausenia nidulans* (Regel) Petrak, Kazakhstan: Almatinskaya oblast, Alatau mt., above Almaty, *Susanna 2088 & al.* (BC), AY373752*; AY373719*; EU661124. *Tiarocarpus hymenostephanus* Rech. f., Afghanistan: W, Herat, Band-e Farsi, 10 SW Farsi, 3,200 m, *Freitag 6841* (KAS), EU923831; EU923945; EU661139. *Tiarocarpus neubaueri* (Rech. f.) Rech. f., Afghanistan: C, Bamian, Band-i Amir, Hochfächen, 3,000 m, *Dieterle 896* (M), EU923775; EU923889; EU661140. *Arctium lappa* L., Iran: NNE, Khorassan: 25 SW Bodjnourd, 1,500 m, *Mehregan 140* (MJG), EU923773; EU923887; EU661123. *Arctium leiospermum* Juz. & C. Serg., Kazakhstan: Zambylskaya oblast, Kurdai pass, 900 m, *Susanna 2154 & al.* (BC), AY373720*; AY373687*; EU661121. *Arctium minus* (Hill.) Bernh., Belgium: Lovaina Botanical Garden, AF19049*; AF19103*. *Cousinia oopoda* Juz. (subg. *Cousinia* sect. *Actinia* Bunge), Tajikistan: South Tajikistan, west mountainside, Khodzha-Kazjyan range, Koy-Pjez-Tau mt., limestone outcrops, 1,240 m, *Kinzikaeva 3585* (LE), EU923839; EU923953; EU661232. *Cousinia tashkurghanica* Rech. f. (subg. *Cousinia* sect. *Actinia* Bunge), Afghanistan: N, Mazar Sharif, Tang-e Sayyad, S of Tashkurghan, 600 m, *Freitag 3905* (KAS), EU923777; EU923891; EU661234. *Cousinia oligocephala* Boiss. (subg. *Cousinia* sect. *Albidae* Rech. f.), Iran: Fars, Inter Shiraz and Sepidan, Shoul pass, 2,000 m, *Mehregan 218* (MJG), EU923796; EU923910; EU661208. *Cousinia chrysantha* Kult. (subg. *Cousinia* sect. *Alpinae* Bunge), Kazakhstan: Shimkientskaya oblast, Aksu Dzabagly reservation, Darbassa canyon, 1,840 m, *Susanna 2198 & al.* (BC), AY373725*; AY373692*; EU661136. *Cousinia dipterocharpa* Bornm. & Rech. f. (subg. *Cousinia* sect. *Alpinae* Bunge), Iran: Khorassan, 27 km from Ghouchan to Bajgiran, Alam Ali pass, 1,820 m, *Mehregan 167* (MJG), EU923862; EU923976; EU661155. *Cousinia lasiolepis* Boiss. (subg. *Cousinia* sect. *Alpinae* Bunge), Iran: Yasoudj, Dena mts., Bizhan Pass, 3,300 m, *Mehregan 202* (MJG), EU923863; EU923977; EU661177. *Cousinia multiloba* DC. (subg. *Cousinia* sect. *Alpinae* Bunge), Iran: Khorassan, SW Bodjnourd, Salouk mt., 2,200 m, *Mehregan 189* (MJG), EU923835; EU923949; EU661202. *Cousinia serawschanica* C. Winkl. (subg. *Cousinia* sect. *Alpinae* Bunge), Tajikistan: Voru, 2,000–2,300 m, *Susanna 2516 & al.* (BC), EU923874; EU923988; EU661192. *Cousinia splendida* C. Winkl. (subg. *Cousinia* sect. *Alpinae* Bunge), Tajikistan: Fan mountains, road above Iskandar-Kul, 2,300 m, *Susanna 2507 & al.* (BC), EU923871; EU923985; EU661197. *Cousinia badghysi* Kult. (subg. *Cousinia* sect. *Badghysia* Tschern.), Turkmenistan: Badghys, Eroylanz lake, SE "sopki" Kazan, *Kamelin 360* (LE), EU923872; EU923986; EU661166. *Cousinia piptocephala* Bunge (subg. *Cousinia* sect. *Badghysia* Tschern.), Iran: Isfahan, 5 km from Mourcheh Khort towards Natanz, 1,500 m, *Mehregan 188* (MJG), EU923801; EU923915; EU661204. *Cousinia ariana* Bornm. (subg. *Cousinia* sect. *Carduncellus* (Juz.) Rech. f.), Afghanistan: E, Kabul, N Salang, 3,150 m, *Freitag 3494* (KAS), EU923788; EU923902; EU661221. *Cousinia buphthalmoides* Regel (subg. *Cousinia* sect. *Carduncellus* (Juz.) Rech. f.), Afghanistan: Unai-Pass, E side, 3,050 m, *Freitag 6268* (KAS), EU923789; EU923903; EU661179. *Cousinia tianschanica* Kult. (subg. *Cousinia* sect. *Carduncellus* (Juz.) Rech. f.), Kazakhstan: Shimkientskaya oblast, Aksu Dzabagly reservation, Aksu canyon, *Susanna 2191 & al.* (BC), AY373743*; AY373710*; EU661127. *Cousinia dissecta* Kar. & Kir. (subg. *Cousinia* sect. *Chrysoptera* Tschern.), Kazakhstan: Almatinskaya oblast, Sugati mts., Kokpekti narrows, *Susanna 2137 & al.* (BC), AY373728*; AY373695*; EU661141. *Cousinia oxiana* Tschern. (subg. *Cousinia* sect. *Chrysoptera* Tschern.), Afghanistan: NW, Herat, 34 km N of Shindand, 1,320 m, *Amin 5252* (KAS), EU923854; EU923968; EU661227. *Cousinia turkmenorum* Bornm. (subg. *Cousinia* sect. *Chrysoptera* Tschern.), Iran: Khorassan, 20 km from Mashhad to Neishabour, 1,150 m, *Mehregan 197* (MJG), EU923855; EU923969; EU661156. *Cousinia greuteri* Rech. f. (subg. *Cousinia* sect. *Congesta* Bunge), Afghanistan: N, Robatak pass between Haibak and Pul-i-Khumri, 1,400 m, *Tonchev 2200* (KAS), EU923783; EU923897; EU661237. *Cousinia coronata* Franch. (subg. *Cousinia* sect. *Coronophora* (Juz.) Rech. f.), Uzbekistan: road Tashkent–Samarkand, 10 km from the cross-roads to Jizzak, 400 m, *Susanna 2039 & al.* (BC), AY373727*; AY373694*; EU661178. *Cousinia radians* Bunge (subg. *Cousinia* sect. *Coronophora* (Juz.) Rech. f.), Iran: Khorassan, 66 km from Mashad to Kalat-Naderi, 1,000 m, *Ghahreman & al. s.n.* (TUH), EU923791; EU923905; EU661217. *Cousinia congesta* Bunge (subg. *Cousinia* sect. *Cousinia*), Uzbekistan: between Samarkand and Kitov, Takhta–Karachi pass, 1,600 m, *Susanna 2059 & al.* (BC), AY373726*; AY373693*; EU661131. *Cousinia daralaghezica* Takht. (subg. *Cousinia* sect. *Cousinia*), Armenia: Vajots Dzor, Eghegnadzor distr., Eghegnadzor town, *Tamanian s.n.* (ERE), EU923813; EU923927; EU661216. *Cousinia intertexta* Freyn & Sint. (subg. *Cousinia* sect. *Cousinia*), Turkey: Sivas, Divrigi-Arapkir, 20 km S Divrigi, 1,410 m, *Nydegger 17154* (M), EU923817; EU923931; EU661235. *Cousinia meghrica* Takht. (subg. *Cousinia* sect. *Cousinia*), Armenia: Sjunik, Meghri distr., Agarak village, *Tamanian s.n.* (ERE), EU923814; EU923928; EU661142. *Cousinia minkwitziae* Bornm. (subg. *Cousinia* sect. *Cousinia*), Kazakhstan: Zambylskaya oblast, Talaski Alatau, 6 km W from Il Tai, 1,000 m, *Susanna 2183 & al.* (BC), AY373735*; AY373702*; EU661135. *Cousinia polycephala* Rupr. (subg. *Cousinia* sect. *Cousinia*), Kazakhstan: Zambylskaya oblast, 30 km from the Kurdai pass, *Susanna 2161 & al.* (BC), AY373738*; AY373705*; EU661138. *Cousinia syrdariensis* Kult. (subg. *Cousinia* sect. *Cousinia*), Kazakhstan: Zambylskaya oblast, 30 km from the Kurdai pass, *Susanna 2159 & al.* (BC), AY373741*; AY373708*; EU661133. *Cousinia tenuifolia* C.A. Mey. (subg. *Cousinia* sect. *Cousinia*), Iran: Azerbaijan, 10 km from Sarab to Ardebil, 1,400 m, *Mehregan 186* (MJG), EU923812; EU923926; EU661171. *Cousinia urumienensis* Bornm. (subg. *Cousinia* sect. *Cousinia*), Iran: Azerbaijan, Urmia, inter Silvana and Razhan, 1,650 m, *Mehregan 201* (MJG), EU923809; EU923923; EU661205. *Cousinia araneosa* DC. (subg. *Cousinia* sect. *Cynaroideae* Bunge), Iran: Hamedan, Alvand mt., 10 km from Hamedan to Toiserkan, Ganjnameh Valley, 2,400 m, *Mehregan 219* (MJG), EU923870; EU923984; EU6611678. *Cousinia macrocephala* C.A. Mey. (subg. *Cousinia* sect. *Cynaroideae* Bunge), Iran: Azerbaijan, Arasbaran, Molouk pass, 1,500 m, *Mehregan 212* (MJG), EU923852; EU923966; EU661169. *Cousinia purpurea* C.A. Mey. in DC. (subg. *Cousinia* sect. *Cynaroideae* Bunge), Armenia: Ararat, near the village of Shahap, roadsides, *Tamanian s.n.* (ERE), AY373739*; AY373706*; EU661137. *Cousinia decumbens* Rech. f. (subg. *Cousinia* sect. *Decumbentes* Rech. f.), Iran: Semnan, Shahroud, mt. Shahvar, SW Slopes, 3,600 m,

Appendix 1. Continued.

Mehregan 110 (MJG), EU923821; EU923935; EU661157. *Cousinia bungeana* Regel & Schmalh. (subg. *Cousinia* sect. *Dichotomae* Bunge), Sine loc.: *Ovczinnikov 16104* (TAD), EU923786; EU923900; EU661130. *Cousinia polytimetica* Tschern. (subg. *Cousinia* sect. *Dichotomae* Bunge), Uzbekistan: Bukharskaya reg., deserted left bank of Zeravshan river, to SE from Uzliskent vil., *Kryakin s.n.* (LE), EU923785; EU923899; EU661244. *Cousinia pusilla* C. Winkl. (subg. *Cousinia* sect. *Dichotomae* Bunge), Tajikistan: S Tajikistan, spackled rock mountains to the S from Besharcha mts. to Babatag range, *Botschantzev 117* (LE), EU923787; EU923901; EU661189. *Cousinia auriculata* Boiss. (subg. *Cousinia* sect. *Eriocousinia* Tschern.), Afghanistan: E, Parvan, Ghowrband, Sorkh-Parsa, 2,050 m, *Freitag 1128* (KAS), EU923830; EU923944; EU661191. *Cousinia chionophila* Rech. & Koeie (subg. *Cousinia* sect. *Eriocousinia* Tschern.), Afghanistan: SE, Ghazni, mts. SE of Dashti Nawor (Sperlu Buli), 3,850 m, *Freitag 1478* (KAS), EU923829; EU923943; EU661247. *Cousinia gmelini* C. Winkl. (subg. *Cousinia* sect. *Eriocousinia* Tschern.), Iran: Kandevar to Yoush, *Djavadi s.n.* (IRAN), EU923842; EU923956; EU661207. *Cousinia polyneura* Rech. f. (subg. *Cousinia* sect. *Eriocousinia* Tschern.), Afghanistan: E, Kabul, mts. above Paghman, 4,000 m, *Freitag 3555* (KAS), EU923781; EU923895; EU661231. *Cousinia xanthacantha* Regel (subg. *Cousinia* sect. *Eriocousinia* Tschern.), Afghanistan: Kabul, in dec. orient. jugi Unai, *Breckle 2286* (M), EU923778; EU923892; EU661230. *Cousinia harazensis* Rech. f. (subg. *Cousinia* sect. *Harazensis* Rech. f.), Iran: Mazandaran, Haraz road, Rineh, 1,900 m, *Mehregan 175* (MJG), EU923825; EU923939; EU661146. *Cousinia gatchsaranica* I. Mehregan, Assadi & Attar (subg. *Cousinia* sect. *Hausknechtii* Rech. f.), Iran: Gachsaran, N slopes of Khami mt., above Tol Chegah, 2,100 m, *Mehregan 83246* (MJG), EU923816; EU923930; EU661159. *Cousinia heliantha* Bunge (subg. *Cousinia* sect. *Helianthae* Bunge), Iran: Khorassan, 30 km from Sabzevar to Esferaien, 1,550 m, *Mehregan 170* (MJG), EU923810; EU923924; EU661172. *Cousinia coerulea* Kult. ex Tschern. (subg. *Cousinia* sect. *Homalochaete* C. Winkl.), Tajikistan: Vorzov canyon, Kudratov, *Susanna 2459 & al.* (BC), EU923873; EU923987; EU661144. *Cousinia crassipes* Kult. (subg. *Cousinia* sect. *Homalochaete* C. Winkl.), Iran: Khorassan, Kalat-e Naderi, Ghareh-Sou, 1,400 m, *Mehregan 144* (MJG), EU923811; EU923925; EU661143. *Cousinia podophylla* Tschern. (subg. *Cousinia* sect. *Homalochaete* C. Winkl.), Sine loc., *Ouchimiku 258* (TAD), EU923961; EU661233. *Cousinia raddeana* C. Winkl. (subg. *Cousinia* sect. *Hoplophylla* Tschern.), Iran: Khorassan, Mashhad, Kalat-e Naderi road, Sanganeh, Chahchaheh, 1,600 m, *Ghahreman & al. 27315* (TUH), EU923853; EU923967; EU661194. *Cousinia immitantiformis* Rech. f. (subg. *Cousinia* sect. *Immitantes* Rech. f.), Afghanistan: NW, Sabzak pass, NE Herat, 2,400 m, *Freitag 6784* (KAS), EU923846; EU923960; EU661149. *Cousinia salangensis* Rech. f. (subg. *Cousinia* sect. *Immitantes* Rech. f.), Afghanistan: E, Kabul, S Salang valley, marble area, 2,500 m, *Freitag 3490* (KAS), EU923845; EU923959; EU661134. *Cousinia jassyensis* C. Winkl. (subg. *Cousinia* sect. *Jurineopsis* (Juz.) Tschern.), Kyrgyzstan: SW, Jalal Abad Oblast, Toktogul Rayoh, 790 m, *Martins 872* (JE), EU923790; EU923904; EU661240. *Cousinia smirnowii* Trautv. (subg. *Cousinia* sect. *Kopetdagia* Tschern.), Iran: Khorassan, SW Bodjnour, Salouk mt., 2,300 m, *Mehregan 152* (MJG), EU923802; EU923916; EU661154. *Cousinia chrysochlora* Rech. f. & Koeie (subg. *Cousinia* sect. *Lachnosphaerae* Rech. f.), Iran: Khorassan, 30 km from Sedeh to Asad Abad, Mahousak, 2,000 m, *Mehregan 198* (MJG), EU923856; EU923970; EU661148. *Cousinia lachnosphaera* Bunge (subg. *Cousinia* sect. *Lachnosphaerae* Rech. f.), Iran: Khorassan, 40 km from Gonabad to Ferdous, 1,900 m, *Mehregan 145* (MJG), EU923857; EU923971; EU661209. *Cousinia alepidea* Boiss. (subg. *Cousinia* sect. *Lasiandrae* Bunge), Afghanistan: SW, Ghazni, Jamroud, 45 km SW of Ghazni, 2,010 m, *Freitag 2008* (KAS), EU923794; EU923908; EU661182. *Cousinia lasiandra* Bunge (subg. *Cousinia* sect. *Lasiandrae* Bunge), Iran: Khorassan, 3 km from Doust Abad to Birjand, 1,500 m, *Mehregan 127* (MJG), EU923822; EU923936; EU661185. *Cousinia triceps* Kult. (subg. *Cousinia* sect. *Leiacanthos* Tschern.), Uzbekistan: Kughitang range, Tanghydevol canyon, rocky area, 870 m, *Kamelin 870* (LE), EU923784; EU923898; EU661129. *Cousinia antonowii* C. Winkl. (subg. *Cousinia* sect. *Leiocaules* Bunge), Iran: Khorassan, 10 km from Tivan to Bajgiran, 1,590 m, *Mehregan 196* (MJG), EU923849; EU923963; EU661200. *Cousinia arctotidifolia* Bunge (subg. *Cousinia* sect. *Leiocaules* Bunge), Iran: Khorassan, Shirvan, 10 km from Ziarat to Lojelli, 1,215 m, *Mehregan 120* (MJG), EU923848; EU923962; EU661165. *Cousinia heterophylla* Boiss. (subg. *Cousinia* sect. *Lepidae* Bunge), Afghanistan: W, N-Kattawz, between Kotanni-Kotal and Patanna, 2,150 m, *Freitag 3345* (KAS), EU923804; EU923918; EU661184. *Cousinia lepida* (Bunge ex) Boiss. (subg. *Cousinia* sect. *Lepidae* Bunge), Iran: Khorassan, Esferaien, 5 km Dahaneh Ojagh to Esferaien, 1,420 m, *Mehregan 114* (MJG), EU923805; EU923919; EU661183. *Cousinia linczewskii* Juz. (subg. *Cousinia* sect. *Leucocaulon* Tschern.), Iran: Khorassan, Inter Mashhad and Sarakhs, Mazd-Avand pass, 1,000 m, *Mehregan 138* (MJG), EU923850; EU923964; EU661153. *Cousinia turcomanica* C. Winkl. (subg. *Cousinia* sect. *Leucocaulon* Tschern.), Iran: Khorassan, 110 km W of Bojnour, between Spakhou and Kastan, 2,000 m, *Mehregan 105* (MJG), EU923834; EU923948; EU661174. *Cousinia rigida* Kult. (subg. *Cousinia* sect. *Lopholepis* Tschern.), Kyrgyzstan: Talasskaya reg., north mountainside of Talassky Alatau, Kur-Bokair canyon, south rocky slope, *Popova s.n.* (LE), EU923869; EU923983; EU661222. *Cousinia arachnoidea* Fisch. & C.A. Mey. (subg. *Cousinia* sect. *Microcarpae* Bunge), Kazakhstan: Almatinskaya oblast, Malai Sary pass, 700 m, *Susanna 2140 & al.* (BC), AY373722*; AY373689*; EU661248. *Cousinia ninae* Juz. (subg. *Cousinia* sect. *Microcarpae* Bunge), Kyrgyzstan: Oshskaya, Torgulsky reg., Oitaya area north from Shoporovo vil., speckled soils, *Sultanova s.n.* (LE), EU923868; EU923982; EU661224. *Cousinia platylepis* Schrenk. ex Fisch. & C.A. Mey. (subg. *Cousinia* sect. *Microcarpae* Bunge), Kazakhstan: Zambylskaya oblast, 30 km from the Kurdai pass, *Susanna 2158 & al.* (BC), AY373737*; AY373704*; EU661236. *Cousinia sewertzowii* Regel (subg. *Cousinia* sect. *Microcarpae* Bunge), Kazakhstan: Aksu-Dzabagly reservation, 1,800 m, *Susanna 2178 & al.* (BC), AY373740*; AY373707*; EU661175. *Cousinia prolifera* Jaub. & Spach (subg. *Cousinia* sect. *Microcousinia* Tschern.), Iran: Khorassan, 110 km from Birjand to Ferdous, 1,500 m, *Mehregan 233* (MJG), EU923779; EU923893; EU661245. *Cousinia heteroloba* Rech. f. (subg. *Cousinia* sect. *Molles* Schrenk.), Afghanistan: NE, Andarab valley below Khinjan, near Ghazan, 1,000 m, *Freitag 3043* (KAS), EU923776; EU923890; EU661198. *Cousinia candolleana* Jaub. & Spach. (subg. *Cousinia* sect. *Myriotemae* Rech. f.), Iran: Ghazvin, 10 km from Abgarm to Avadj, 1,950 m, *Mehregan 239* (MJG), EU923858; EU923972; EU661195. *Cousinia olgae* Regel. & Schmalh. (subg. *Cousinia* sect. *Olgaeanthae* Tschern.), Iran: Khorassan, 40 km from Torbat-e Djam to Saleh Abad, 1,550 m, *Mehregan 194* (MJG), EU923780; EU923894; EU661203. *Cousinia eriophylla* (Kult.) Bornm. (subg. *Cousinia* sect. *Platyacanthae* Rech. f.), Iran: Khorassan, Inter Ghouchan and Bajgiran, 5 km S of Dorbadam, 1,500 m, *Mehregan 190* (MJG), EU923840; EU923954; EU661160. *Cousinia re-chingerorum* Bornm. (subg. *Cousinia* sect. *Platyacanthae* Rech. f.), Iran: Khorassan: 30 km W of Torbat-e Djam, E slopes of Bezd

Appendix 1. Continued.

mt., 2,000 m, *Mehregan 163* (MJG), EU923841; EU923955; EU66116. *Cousinia trachyphyllaria* Bornm. & Rech. f. (subg. *Cousinia* sect. *Platyacanthae* Rech. f.), Iran: Khorassan, Neishabour, Khorw, 2,100 m, *Mehregan 217* (MJG), EU923828; EU923942; EU661163. *Cousinia apiculata* Tschern. (subg. *Cousinia* sect. *Pseudactinia* Tschern.), Iran: Khorassan, Shirvan, Sarani, 2,300 m, *Gahreman & Attar s.n.* (MJG), EU923792; EU923906; EU661162. *Cousinia oreodoxa* Bornm. & Sint. (subg. *Cousinia* sect. *Pseudactinia* Tschern.), Iran: Khorassan, Inter Bodjnour and Raz, 7 km from Tangeh Torkeman to Ashkhaneh, 1,000 m, *Mehregan 150* (MJG), EU923793; EU923907; EU661151. *Cousinia belangeri* DC. (subg. *Cousinia* sect. *Pugioniferae* Bunge), Iran: Tehran, 15 km to Karaj, 1,200 m, *Mehregan s.n.* (MJG), EU923819; EU923933; EU661210. *Cousinia calcitrapa* DC. (subg. *Cousinia* sect. *Pugioniferae* Bunge), Iran: Fars, pass inter Sepidan and Yassoudj, 20 km from Sepidan, 2,400 m, *Mehregan 203* (MJG), EU923820; EU923934; EU661220. *Cousinia incompta* DC. (subg. *Cousinia* sect. *Pugioniferae* Bunge), Iran: 20 km Borujen to Isfahan, 2,400 m, *Mehregan 128* (MJG), EU923851; EU923965; EU661186. *Cousinia pineticola* Rech. f. & Gilli (subg. *Cousinia* sect. *Racemosae* Rech. f.), Afghanistan: E, Kabul, Korogh Koh, SW of Kabul, 2,600 m, *Freitag 1568* (KAS), EU923832; EU923946; EU661107. *Cousinia botschantzevii* Juz. ex Tschern. (subg. *Cousinia* sect. *Regelianae* (Juz.) Tschern.), Uzbekistan: Kuramit mt., Koitash range, S slope above the Koitash village, 1,850 m, *Kamelin 29* (LE), EU923782; EU923896; EU661250. *Cousinia autranii* C. Winkl. (subg. *Cousinia* sect. *Rigidissimae* Rech. f.), Afghanistan: W, Herat, Ghorat, Bande Frasi, 15 km SSE Farsi, 2,400–2,500 m, *Freitag 6831* (KAS), EU923860; EU923974; EU661105. *Cousinia rigidissima* Rech. f. (subg. *Cousinia* sect. *Rigidissimae* Rech. f.), Afghanistan: W, Herat, Ghorat, in summo jugo 5 km S Dahan-e Kaftarkhan ad viam inter Tulak et Farsi, 2,350 m, *Podlech 19193* (M), EU923861; EU923975; EU661187. *Cousinia scariosa* Regel (subg. *Cousinia* sect. *Scariosae* Rech. f.), Afghanistan: N, Sar-e Pol, Sangcharak, 40 km SSW mts. above Damdaran, 1,900–2,500 m, *Freitag 6594* (KAS), EU923838; EU923952; EU661190. *Cousinia eryngioides* Boiss. (subg. *Cousinia* sect. *Sciadocousinia* Tschern.), Iran: Khorassan, 50 km from Mashhad to Neishabour, 1,500 m, *Mehregan 168* (MJG), EU923836; EU923950; EU661164. *Cousinia concolor* Bunge (subg. *Cousinia* sect. *Serratuloides* Bunge), Iran: Semnan, Shahroud, S slopes of Shahvar mt., above Nekarman, 2,500 m, *Mehregan 158* (MJG), EU923808; EU923922; EU661150. *Cousinia crispa* Jaub. & Spach. (subg. *Cousinia* sect. *Serratuloides* Bunge), Iran: Chalous road, 10 km from Kandavan to Chalous, 2,000 m, *Mehregan 185* (MJG), EU923843; EU923957; EU661241. *Cousinia hypoleuca* Boiss. (subg. *Cousinia* sect. *Serratuloides* Bunge), Iran: Tehran, Inter Firouzkouh and Polur, 5 km W of Lasem, 2,500 m, *Mehregan 177* (MJG), EU923807; EU923921; EU661176. *Cousinia pinarocephala* Boiss. (subg. *Cousinia* sect. *Serratuloides* Bunge), Iran: Tehran, Firouzkouh, Gadouk pass, 2,200 m, *Mehregan 192* (MJG), EU923844; EU923958; EU661223. *Cousinia adenosticta* Bornm. (subg. *Cousinia* sect. *Sphaerocephalae* Bunge), Iran: Tehran, between Shemshak and Dizin, 3,000 m, *Mehregan 165* (MJG), EU923827; EU923941; EU661110. *Cousinia leptolepis* (Bornm. & Gauba) Rech. f. (subg. *Cousinia* sect. *Sphaerocephalae* Bunge), Iran: Tehran, Karadj, Kuh Dashteh, S slopes, 2,200 m, *Mehregan 111* (MJG), EU923826; EU923940; EU661206. *Cousinia shahvarica* Rech.f. (subg. *Cousinia* sect. *Sphaerocephalae* Bunge), Iran: Semnan, Shahroud, mt. Shahvar, S slopes, 3,000 m, *Mehregan 117* (MJG), EU923823; EU923937; EU661180. *Cousinia xiphiolepis* Boiss. (subg. *Cousinia* sect. *Sphaerocephalae* Bunge), Iran: Tehran, inter Firouzkouh and Polur, 5 km W of Lasem, 2,600 m, *Mehregan 179* (MJG), EU923824; EU923938; EU661188. *Cousinia longifolia* C. Winkl. & Bornm. (subg. *Cousinia* sect. *Spinuliferae* Rech. f.), Iran: Kerman, Laleh zar to Rabor, *Attar 226* (MJG), EU923815; EU923929; EU661158. *Cousinia qaisarensis* Rech. f. (subg. *Cousinia* sect. *Spinuliferae* Rech.f.), Afghanistan: NW, Fariab, Maimana, Qaisar, 20 km, mts. above Shakh (Gala Shakh), 2,200 m, *Freitag 6745* (KAS), EU923833; EU923947; EU661201. *Cousinia alexeenkoana* Bornm. (subg. *Cousinia* sect. *Stenocephalae* Bunge), Iran: Isfahan, 10 km from Khaansaar to Boein, 2,500 m, *Mehregan 132* (MJG), EU923799; EU923913; EU661181. *Cousinia cylindracea* Boiss. (subg. *Cousinia* sect. *Stenocephalae* Bunge), Iran: Fars, inter Shiraz and Sepidan, Shoul pass, 2,000 m, *Mehregan 213* (MJG), EU923797; EU923911; EU661108. *Cousinia gaubae* Bornm. (subg. *Cousinia* sect. *Stenocephalae* Bunge), Iran: Ghazvin, Karadj, in declivibus lapidosis, 1,400 m, *Rechinger 5504* (M), EU923818; EU923932; EU661109. *Cousinia tenuiramula* Rech. f. (subg. *Cousinia* sect. *Stenocephalae* Bunge), Iran: Bakhtiari, Rokh pass, 2,200 m, *Mehregan 206* (MJG), EU923798; EU923912; EU661214. *Cousinia thamnoides* Boiss. & Hausskn. (subg. *Cousinia* sect. *Stenocephalae* Bunge), Iran: Gachsaran, Khami mt. above Tol Chegah, 3,170 m, *Mehregan 237* (MJG), EU923800; EU923914; EU661212. *Cousinia komarovii* (O. Kuntze) C. Winkl. (subg. *Cousinia* sect. *Stenoloma* Juz.), Iran: Khorassan, 15 km Bajgiran to Ghouchan, 1,700 m, *Mehregan 156* (MJG), EU923803; EU923917; EU661152. *Cousinia leucantha* Bornm. & Sint. (subg. *Cousinia* sect. *Stenoloma* Juz.), Iran: Golestan Nat. Park, 12 km ENE of Tange Gol, 1,900–2,100 m, *Akhani 11357* (M), EU923837; EU923951; EU661170. *Cousinia stocksii* C. Winkl. (subg. *Cousinia* sect. *Stocksianae* Rech. f.), Afghanistan: W, Farah, Schucht, 1 km nördlich Hakumate Purchaman, 1,880 m, *Podlech 21787* (M), EU923795; EU923909; EU661116. *Cousinia tenella* Fisch. & C.A. Mey. (subg. *Cousinia* sect. *Tenellae* Bunge), Iran: Golestan Nat. Park, between Sharlegh and Cheshmeh Khan, *Akhani 243* (MJG), EU923774; EU923888; EU661104. *Cousinia chaetocephala* Kult. (subg. *Cousinia* sect. *unknown*), Iran: Khorassan, Tchenaran, 3km from Meritchehan to Radekan, 1,500 m, *Mehregan 115* (MJG), EU923806; EU923920; EU661145. *Cousinia pallidivirens* Kult. (subg. *Cynaroides* Tschern. unassigned to any section), Uzbekistan: Sine loc., *Botschantzev s.n.* (LE), EU923768; EU923882; EU661126. *Cousinia chlorantha* Kult. (subg. *Cynaroides* Tschern. sect. *Chrysis* Juz.), Uzbekistan: Malguzar mt., S slope, by Tashkesken "say", *Kamelin 199* (LE), EU923864; EU923978; EU661120. *Cousinia karatavica* Regel et Schmalh. (subg. *Cynaroides* Tschern. sect. *Chrysis* Juz.), Kazakhstan: Zambylskaya oblast, Karatau mt., Kuyuk pass, 1,000 m, *Susanna 2162* & al. (BC), AY373732*; AY373699*; EU661101. *Cousinia kolkovii* Regel & Schmalh. (subg. *Cynaroides* Tschern. sect. *Chrysis* Juz.), Uzbekistan: N macro-mountainside, Nuratau range, Sintob kishlak surroundings, rocky slopes, *Botschantzev 427* (LE), EU923865; EU923979; EU661103. *Cousinia medians* Juz. (subg. *Cynaroides* Tschern. sect. *Chrysis* Juz.), Uzbekistan: S, Supkhandarbinskaya reg., between kishlaks Sajrob and Shurob, 121–122 km of the road from Termez, Speckled rock outcrops, *Botschantzev 240* (LE), EU923769; EU923883; EU661100. *Cousinia refracta* (Bornm.) Juz. (subg. *Cynaroides* Tschern. sect. *Chrysis* Juz.), Tajikistan: Kondara river canyon, Varzowski Rayon reservation, *Susanna 2456* & al. (BC), EU923867; EU923981; EU661111. *Cousinia anomala* Franch. (subg. *Cynaroides* Tschern. sect. *Ctenarctium* Juz.), Tajikistan: Slopes above Voru, above 300 m above Kishlak, 2,200–2,300 m, *Susanna 2521* & al. (BC), EU923770; EU923884; EU661115. *Cousinia lappacea* Schrenk (subg. *Cynaroides* Tschern. sect. *Lappaceae* Bunge), Kazakhstan: Zambylskaya oblast, Kurdai pass, 900 m, *Susanna 2150* & al. (BC), AY373733*;

Appendix 1. Continued.

AY373700*; EU661112. *Cousinia arctioides* Schrenk (subg. *Cynaroides* Tschern. sect. *Nanarctium* Juz. ex. Tschern.), Kazakhstan: Dzhezkazganskaya reg., Turgajskaya lowland, 49 km to SW from Dzhezkazgana, right bank of Kumula river, Tamarix bushland, *Kamelin 6434* (LE), EU923772; EU923886; EU661118. *Cousinia triflora* Schrenk (subg. *Cynaroides* Tschern. sect. *Oligantha* Juz.), Iran: Golestan Nat. Park, Yakhbala pass, *Akhani 102* (MJG), EU923771; EU923885; EU661094. *Cousinia albertii* Regel et Schmalh. (subg. *Cynaroides* Tschern. sect. *Pectinatae* C. Winkl.), Kazakhstan: Shimkientskaya oblast, Mashat canyon, 1,840 m, *Susanna 2206* & al. (BC), AY373721*; AY373688*; EU661099. *Cousinia amplissima* (Boiss.) Boiss. (subg. *Cynaroides* Tschern. sect. *Pseudarctium* Juz.), Iran: Dena, 15 km from Meimand to Yassoudj, 2,500 m, *Mehregan 174* (MJG), EU923766; EU923880; EU661098. *Cousinia pseudarctium* Bornm. (subg. *Cynaroides* Tschern. sect. *Pseudarctium* Juz.), Tajikistan: Vorzov valley, 2 km N kishlak Ziddy, *Susanna 2477* & al. (BC), EU923876; EU923990; EU661095. *Cousinia tomentella* C. Winkl. (subg. *Cynaroides* Tschern. sect. *Pseudarctium* Juz.), Tajikistan: S mountainside of Guissar range, left bank of Varzob river, Deamalik kishlak surroundings, 1,800 m, *Tschukavina 10512* (LE), EU923767; EU923881; EU661097. *Cousinia umbrosa* Bunge (subg. *Cynaroides* Tschern. sect. *Pseudarctium* Juz.), Kazakhstan: Almatinskaya oblast, Alatau mt. above Almaty, 1,200 m, *Susanna 2100* & al. (BC), AY373745*; AY373712*; EU661096. *Cousinia abolinii* Kult. ex Tscherneva (subg. *Hypacanthodes* Tschern. sect. *Abolinia* Tschern.), Kyrgyzstan: SW, Jalal Abad Oblast, Kara Saj Tal, Aksy Rayan, 1,030 m, *Lazkov s.n.* (JE), EU923763; EU923877; EU661113. *Cousinia dolichophylla* Kult. (subg. *Hypacanthodes* Tschern. sect. *Abolinia* Tschern.), Uzbekistan: Ugamsky range above Nanaj, right edge of Pskem valley, “shiblyak”, *Kamelin 265* (LE), EU923875; EU923989; EU661117. *Cousinia egregia* Juz. (subg. *Hypacanthodes* Tschern. sect. *Abolinia* Tschern.), Uzbekistan: Angren valley, rise to Kamchik pass, rubby slope, *Kamelin 420* (LE), EU923866; EU923980; EU661196. *Cousinia grandifolia* Kult. (subg. *Hypacanthodes* Tschern. sect. *Amberbopsis* Tschern.), Kazakhstan: Zambylskaya oblast, Talaski Alatau, 6 km W from Il Tai, 1,000 m, *Susanna 2181* & al. (BC), AY373730*; AY373697*; EU661114. *Cousinia korshinskyi* C. Winkl. (subg. *Hypacanthodes* Tschern. sect. *Lacerae* C. Winkl.), Kyrgyzstan: isolated terrain feature Kanka, upper waters of river Kanka, near snow pot, h. 2,300 m, *E.M. Il'ina s.n.* (LE), EU923765; EU923879; EU661102. *Cousinia macilenta* C. Winkl. (subg. *Hypacanthodes* Tschern. sect. *Lacerae* C. Winkl.), Tajikistan: SW, Jugum Hissaricum (Gissar), divorticum aquarum inter flumina Ljuzob et Unou, 3,000 m, *Zaprojagaev s.n.* (M), EU923764; EU923878; EU661119.

Appendix 2. Geographical distribution and taxonomic composition of clades found in the BI analyses of the ITS, *rps4-trnT-trnL* and combined ITS + *rps4-trnT-trnL* datasets. Numbers refer to clades or subclades of the Cousinioid group with PP ≥ 0.95 in Figs. 2–4. Geographical distribution according to the eight main centres of diversity of *Cousinia* as shown in Fig. 1.

***rps4-trnT-trnL*:** CLADE 1: *C. pinarocephala*, sect. *Serratuloides*, Reg. 3; *C. crispa*, sect. *Serratuloides*, Reg. 3. CLADE 2: *C. araneosa*, sect. *Cynaroides*, Reg. 2; *C. leucantha*, sect. *Stenoloma*, Reg. 4; (NESTED SUBCLADE 2a: *C. smirnowii*, sect. *Kopetdagia*, Reg. 4; *C. komarowii*, sect. *Stenoloma*, Reg. 4; *C. linczewskii*, sect. *Leucocaulon*, Reg. 4; *C. dipterocarpa*, sect. *Alpinae*, Reg. 4; *C. turkmenorum*, sect. *Chrysoptera*, Reg. 4; *C. decumbens*, sect. *Decumbentes*, Reg. 4; *C. eriophylla*, sect. *Platyacanthae*, Reg. 4; *C. rechingerorum*, sect. *Platyacanthae*, Reg. 4; *C. apiculata*, sect. *Pseudactinia*, Reg. 4; *C. trachyphyllaria*, sect. *Platyacanthae*, Reg. 4; *C. gatchsaranica*, sect. *Hausknechtii*, Reg. 2; *C. macrocephala*, sect. *Cynaroides*, Reg. 1; *C. eryngioides*, sect. *Sciadocousinia*, Reg. 3, 4; *C. arctotidifolia*, sect. *Leiocaules*, Reg. 4; *C. badghysi*, sect. *Badghysia*, Reg. 4; *C. longifolia*, sect. *Spinuliferae*, Reg. 4). CLADE 3: *C. sewertzowii*, sect. *Microcarpae*, Reg. 7; *C. turcomanica*, sect. *Leucocaulon*, Reg. 4. CLADE 4: *C. serawschanica*, sect. *Alpinae*, Reg. 7; *C. raddeana*, sect. *Hoplophylla*, Reg. 4, 5, 7; *C. splendida*, sect. *Alpinae*, Reg. 7; *C. heteroloba*, sect. *Molles*, Reg. 5, 6. CLADE 5: *C. jassyensis*, sect. *Jurineopsis*, Reg. 8; *C. ninae*, sect. *Microcarpae*, Reg. 7, 8; *C. arachnoidea*, sect. *Microcarpae*, Reg. 8; (NESTED SUBCLADE 5a: *C. tianschanica*, sect. *Carduncellus*, Reg. 8; *C. triceps*, sect. *Leiacanthos*, Reg. 7; *C. platylepis*, sect. *Microcarpae*, Reg. 5, 7, 8; *C. greuteri*, sect. *Congesta*, Reg. 5, 6; *C. oopoda*, sect. *Actinia*, Reg. 7; *C. podophylla*, sect. *Homalochaete*, Reg. 7; *C. oxiana*, sect. *Chrysoptera*, Reg. 4; *C. xanthacantha*, sect. *Eriocousinia*, Reg. 6; *C. polyneura*, sect. *Eriocousinia*, Reg. 6; *C. tashkurghanica*, sect. *Actinia*, Reg. 5; *C. intertexta*, sect. *Cousinia*, Reg. 1; *C. chionophila*, sect. *Eriocousinia*, Reg. 5; *C. botschantzevii*, sect. *Regelianae*, Reg. 7). **ITS:** CLADE 1: *C. splendida*, sect. *Alpinae*, Reg. 7; *C. ariana*, sect. *Carduncellus*, Reg. 5, 6; *C. bupthalmoides*, sect. *Carduncellus*, Reg. 5, 6, 7; *C. sewertzowii*, sect. *Microcarpae*, Reg. 7; *C. ninae*, sect. *Microcarpae*, Reg. 8, 7; *C. jassyensis*, sect. *Jurineopsis*, Reg. 8; (NESTED SUBCLADE 1a: *C. coronata*, sect. *Coronophora*, Reg. 7; *C. radians*, sect. *Coronophora*, Reg. 4). CLADE 2: *C. tianschanica*, sect. *Carduncellus*, Reg. 8; *C. botschantzevii*, sect. *Regelianae*, Reg. 7. CLADE 3: *C. polytimetica*, sect. *Dichotomae*, Reg. 7; *C. pusilla*, sect. *Dichotomae*, Reg. 7; *C. bungeana*, sect. *Dichotomae*, Reg. 8. CLADE 4: *C. minkwitziae*, sect. *Cousinia*, Reg. 8; *C. chrysantha*, sect. *Alpinae*, Reg. 8; *C. dissecta*, sect. *Chrysoptera*, Reg. 8; (NESTED SUBCLADE 4a: *C. syrdariensis*, sect. *Cousinia*, Reg. 8; *C. polycephala*, sect. *Cousinia*, Reg. 7, 8). CLADE 5: *C. serawschanica*, sect. *Alpinae*, Reg. 7; *C. polyneura*, sect. *Eriocousinia*, Reg. 6. CLADE 6: *C. hypoleuca*, sect. *Serratuloides*, Reg. 3; *C. concolor*, sect. *Serratuloides*, Reg. 3, 4. CLADE 7: *C. rigida*, sect. *Lopholepis*, Reg. 8; (NESTED SUBCLADE 7a: *C. stocksii*, sect. *Stocksianae*, Reg. 4, 5; (NESTED SUBCLADE 7aa: *C. tenuiramula*, sect. *Stenocephalae* s.l., Reg. 2; *C. piptocephala*, sect. *Badghysia*, Reg. 2; *C. thamnoides*, sect. *Stenocephalae* s.l., Reg. 2; *C. alexeenkoana*, sect. *Stenocephalae* s.l., Reg. 2; *C. cylindracea*, sect. *Stenocephalae* s.l., Reg. 2; *C. oligocephala*, sect. *Stenocephalae* s.l., Reg. 2)); (NESTED SUBCLADE 7b: *C. apiculata*, sect. *Pseudactinia*, Reg. 4; *C. oreodoxa*, sect. *Pseudactinia*, Reg. 4); (NESTED SUBCLADE 7c: *C. chaetocephala*, sect. unknown, Reg. 4; *C. lepida*, sect. *Lepidae*, Reg. 4; *C. heterophylla*, sect. *Lepidae*, Reg. 4, 5). CLADE 8: *C. gmelinii*, sect. *Serratuloides*, Reg. 3; *C. pinarocephala*, sect. *Serratuloides*, Reg. 3; *C. crispa*, sect. *Serratuloides*, Reg. 3. CLADE 9: *C. eriophylla*, sect. *Platyacanthae*, Reg. 4; *C. rechingerorum*, sect. *Platyacanthae*, Reg. 4. CLADE 10: *C. salangensis*, sect. *Immitantes*, Reg. 6; *C. immitantiformis*, sect. *Immitantes*, Reg. 5; (NESTED SUBCLADE 10a: *C. coerulea*, sect. *Homalochaete*, Reg. 7; *C. podophylla*,

Appendix 2. Continued.

sect. *Homalochaete*, Reg. 7). CLADE 11: *C. incompta*, sect. *Pugioniferae*, Reg. 2; (NESTED SUBCLADE 11a: *C. purpurea*, sect. *Cynaroideae*, Reg. 1; *C. araneosa*, sect. *Cynaroideae*, Reg. 2; *C. macrocephala*, sect. *Cynaroideae*, Reg. 1). CLADE 12: *C. chrysochlora*, sect. *Lachnosphaerae*, Reg. 4; *C. lachnosphaera*, sect. *Lachnosphaerae*, Reg. 4, 5. CLADE 13: *C. turkmenorum*, sect. *Chrysoptera*, Reg. 4; *C. oxiana*, sect. *Chrysoptera*, Reg. 4. CLADE 14: *C. autranii*, sect. *Rigidissimae*, Reg. 5; *C. rigidissimae*, sect. *Rigidissimae*, Reg. 5. CLADE 15: *C. auriculata*, sect. *Eriocousinia*, Reg. 5, 6, 7; *C. chionophila*, sect. *Eriocousinia*, Reg. 5. CLADE 16: *C. shahvarica*, sect. *Sphaerocephalae*, Reg. 3; (NESTED SUBCLADE 16a: *C. harazensis*, sect. *Sphaerocephalae*, Reg. 3; *C. xiphiolepis*, sect. *Sphaerocephalae*, Reg. 3); (NESTED SUBCLADE 16b: *C. adenosticta*, sect. *Sphaerocephalae*, Reg. 3; *C. leptolepis*, sect. *Sphaerocephalae*, Reg. 3). CLADE 17: *C. decumbens*, sect. *Decumbentes*, Reg. 3; *C. badphysi*, sect. *Badghysia*, Reg. 4. CLADE 18: *C. gatchsaranica*, sect. *Hausknechtii*, Reg. 2; *C. intertexta*, sect. *Cousinia*, Reg. 1. **Combined ITS + rps4-trnT-trnL:** CLADE 1: *C. polytimetica*, sect. *Dichotomae*, Reg. 7; *C. pusilla*, sect. *Dichotomae*, Reg. 7; *C. bungeana*, sect. *Dichotomae*, Reg. 8. CLADE 2: *C. chrysantha*, sect. *Alpinae*, Reg. 8; *C. dissecta*, sect. *Chrysoptera*, Reg. 8; (NESTED SUBCLADE 2a: *C. minkwitziae*, sect. *Cousinia*, Reg. 8; *C. syrdariensis*, sect. *Cousinia*, Reg. 8; *C. polycephala*, sect. *Cousinia*, Reg. 7, 8). CLADE 3: *C. splendida*, sect. *Alpinae*, Reg. 7; *C. ariana*, sect. *Carduncellus*, Reg. 6, 5; (NESTED SUBCLADE 3a: *C. bupthalmoides*, sect. *Carduncellus*, Reg. 5, 6, 7; *C. sewertzowii*, sect. *Microcarpae*, Reg. 7; (NESTED SUBCLADE 3aa: *C. jassyensis*, sect. *Jurineopsis*, Reg. 8; *C. ninae*, sect. *Microcarpae*, Reg. 7, 8); (NESTED SUBCLADE 3ab: *C. coronata*, sect. *Coronophora*, Reg. 7; *C. radians*, sect. *Coronophora*, Reg. 4)). CLADE 4: *C. hypoleuca*, sect. *Serratuloides*, Reg. 3; *C. concolor*, sect. *Serratuloides*, Reg. 3, 4. CLADE 5: *C. stocksii*, sect. *Stocksianae*, Reg. 4, 5; *C. komarowii*, sect. *Stenoloma*, Reg. 4; *C. alepideae*, sect. *Lasiandrae*, Reg. 5; (NESTED SUBCLADE 5a: *C. cylindracea*, sect. *Stenocephalae* s.l., Reg. 2; *C. oligocephala*, sect. *Stenocephalae* s.l., Reg. 2; (NESTED SUBCLADE 5aa: *C. tenuiramula*, sect. *Stenocephalae* s.l., Reg. 2; *C. piptocephala*, sect. *Badghysia*, Reg. 2; *C. thamnoides*, sect. *Stenocephalae* s.l., Reg. 2; *C. alexeenkoana*, sect. *Stenocephalae* s.l., Reg. 2)). CLADE 6: *C. chaetocephala*, sect. unknown, Reg. 4; *C. lepida*, sect. *Lepidae*, Reg. 4; *C. heterophylla*, sect. *Lepidae*, Reg. 4, 5. CLADE 7: *C. chrysochlora*, sect. *Lachnosphaerae*, Reg. 4; *C. lachnosphaera*, sect. *Lachnosphaerae*, Reg. 4, 5. CLADE 8: *C. gmelinii*, sect. *Serratuloides*, Reg. 3; (NESTED SUBCLADE 8a: *C. pinarocephala*, sect. *Serratuloides*, Reg. 3; *C. crispa*, sect. *Serratuloides*, Reg. 3). CLADE 9: *C. lasiolepis*, sect. *Alpinae*, Reg. 2, 4, 5; (NESTED SUBCLADE 9a: *C. autranii*, sect. *Rigidissimae*, Reg. 5; *C. rigidissimae*, sect. *Rigidissimae*, Reg. 5). CLADE 10: *C. araneosa*, sect. *Cynaroideae*, Reg. 2; *C. linczewskii*, sect. *Leucocaulon*, Reg. 4; *C. arctotidifolia*, sect. *Leiocaules*, Reg. 4; *C. smirnowii*, sect. *Kopetdagia*, Reg. 4; *C. eryngioides*, sect. *Sciadocousinia*, Reg. 3, 4; *C. dipterocharpa*, sect. *Alpinae*, Reg. 4; *C. turkmenorum*, sect. *Chrysoptera*, Reg. 4; (NESTED SUBCLADE 10a: *C. eriophylla*, sect. *Platyacanthae*, Reg. 4; *C. rechingerorum*, sect. *Platyacanthae*, Reg. 4); (NESTED SUBCLADE 10b: *C. apiculata*, sect. *Pseudactinia*, Reg. 4; *C. trachyphyllaria*, sect. *Platyacanthae*, Reg. 4). CLADE 11: *C. shahvarica*, sect. *Sphaerocephalae*, Reg. 3; (NESTED SUBCLADE 11a: *C. harazensis*, sect. *Sphaerocephalae*, Reg. 3; *C. xiphiolepis*, sect. *Sphaerocephalae*, Reg. 3); (NESTED SUBCLADE 11b: *C. adenosticta*, sect. *Sphaerocephalae*, Reg. 3; *C. leptolepis*, sect. *Sphaerocephalae*, Reg. 3). CLADE 12: *C. decumbens*, sect. *Decumbentes*, Reg. 3; *C. badphysi*, sect. *Badghysia*, Reg. 4. CLADE 13: *C. gatchsaranica*, sect. *Hausknechtii*, Reg. 2; *C. longifolia*, sect. *Spinuliferae*, Reg. 4. CLADE 14: *C. incompta*, sect. *Pugioniferae*, Reg. 2; (NESTED SUBCLADE 14a: *C. araneosa*, sect. *Cynaroideae*, Reg. 2; *C. purpurea*, sect. *Cynaroideae*, Reg. 1). CLADE 15: *C. tianschanica*, sect. *Carduncellus*, Reg. 8; *C. botschantzevii*, sect. *Regelianae*, Reg. 7.