

# Molecular systematics of *Echinops* L. (Asteraceae, Cynareae): A phylogeny based on ITS and *trnL-trnF* sequences with emphasis on sectional delimitation

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**Abstract** The monogeneric subtribe Echinopsinae of Cynareae (Asteraceae) was analysed using nrDNA ITS and plastid *trnL-trnF* for 81 of the ca. 120 species, representing all *Echinops* sections except *Cenchrrolepis* and *Pterolepis*. Maximum parsimony and Bayesian analyses were used for each dataset and for the combined data. The resulting molecular phylogenetic framework resolves *Echinops* as monophyletic and confirms the inclusion of *E. acantholepis* (= *Acantholepis orientalis*) within the genus. *Echinops* is divided into two lineages, one consisting of *E.* sect. *Chamaechinops* and sect. *Acantholepis* (almost exclusively annual or biennial), and the other the remaining sections (almost all perennial). Our results support an infrageneric classification of *Echinops* into nine previously recognized sections: *Acantholepis* (= *Nanechinops*), *Chamaechinops*, *Echinops* (= *Terma*), *Hamolepis*, *Hololeuce*, *Oligolepis*, *Phaeochaete*, *Psectra* and *Ritropsis*, all resolved as strongly monophyletic after some species relocations. The status of *E.* sect. *Phaeochaete* and the sectional assignment of *E. onopordum*, *E. pungens* and *E. transcaucasicus* remain to be clarified. The evolution of involucre bracts is congruent with the phylogenetic framework and supports the sectional classification. Annual pollen type does not constitute an autapomorphy of the clade grouping the annuals but may represent the ancestral character state for the entire genus.

**Keywords** *Acantholepis*; Bayesian inference; Compositae; Echinopsinae; morphological characters

## INTRODUCTION

The genus *Echinops* s.l. (including *Acantholepis* Less.) is the only member of Echinopsinae (Cass.) Dumort., one of the five subtribes of Cynareae Lam. & DC., otherwise known as Cardueae Cass. (Susanna & al., 2006). It is characterized by the presence of unflowered capitula aggregated into second order spherical or oval heads, this syncephalia being a unique feature within the tribe (Petit, 1997). *Echinops* comprises ca. 120 species (Bobrov, 1997; Susanna & Garcia-Jacas, 2007) distributed in tropical Africa, the Mediterranean basin, temperate regions of Eurasia, reaching Central Asia, Mongolia and north-eastern China, with the greater number of species occurring in the Caucasus and the Middle East (for a distribution map of the genus, see Jäger, 1987).

The strong morphological uniformity of *Echinops* makes its taxonomical delimitation to be almost unquestioned, but it also hinders the attempts at establishing natural groups and an infrageneric classification. The most complete analysis of the genus was made by Bunge (1863) who recognized twelve sections, seven of them new. After Bunge, revisions were published by Fries (1923), Jeffrey (1968) and Tadesse (1997) for tropical Africa, Hedge (1975) for the Turkish flora, Kožuharov (1976) for the European taxa, Rechinger (1979) and Mozaffarian (2008) for the Iranian flora, and Bobrov (1997) for the former U.S.S.R.

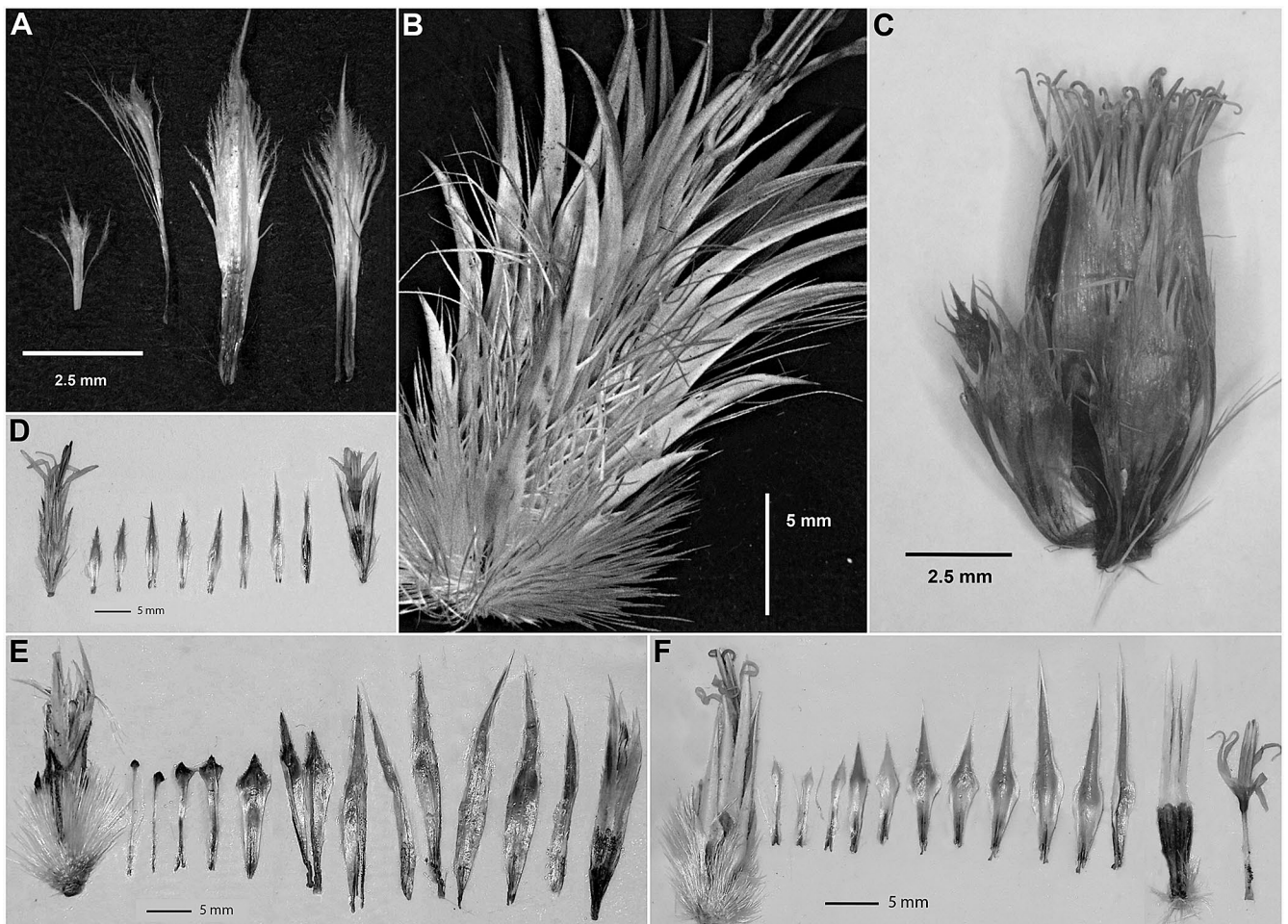
territories. Morphological characters used for infrageneric classifications of *Echinops* are almost limited to the bracts of the unflowered capitula, such as their number, or the degree of connation of the inner bracts (Fig. 1; Table 1; Hedge, 1975; Kožuharov, 1976; Rechinger, 1979; Bobrov, 1997). In this genus, the diversity of the involucre bracts seems to be related to the fact that the one-seeded capitulum is the unit of dispersal, and therefore has an adaptive value (Davis, 1956). On the contrary, the pappus—a main source of key taxonomical characters for other Cynareae—plays no role in dissemination and is very short and quite uniform throughout *Echinops* species. Other features like the type and density of indumentum on stems, leaves and phyllaries display a certain amount of variability that provides taxonomically useful characters (Mozaffarian, 2006).

A previous attempt at molecular phylogenetic reconstruction for *Echinops* (Garnatje & al., 2005) consisted of a parsimony analysis based on the ITS region and representing 30 species and the monotypic genus *Acantholepis*; that is, hardly a quarter of all the members in the group. *Echinops* (*Acantholepis* included) was consistently resolved as a natural group in this study, as well as in molecular phylogenies of the tribe Cynareae (Garcia-Jacas & al., 2002; Susanna & al., 2006). The inclusion of *Acantholepis orientalis* Less. within *Echinops* species (Garnatje & al., 2005) agreed with its treatment as *E. acantholepis* (Jaubert & Spach, 1848). Nevertheless, most authors had considered

this taxon as an independent genus close to *Echinops* (Hedge, 1975; Dittrich, 1977, 1996; Rechinger, 1979; Petit, 1988, 1997; Bremer, 1994; Bobrov, 1997). The molecular evidence of Garnatje & al. (2005) suggested a tight relationship between *Acantholepis* and *E. nanus*, an assumption consistent with life cycles (they are both annual), and pollen type data (Garnatje & Martín, 2007). Nevertheless, using traditional morphological characters, *Acantholepis* is well distinguished from *Echinops* species by having second-order head involucre bracts well developed and patent; the outer ones are similar to the leaves and exceed notably the remaining parts of the syncephalia, whereas these bracts in *Echinops* are comparatively small and hidden. The karyological data also differentiate *Acantholepis* and *Echinops*, although both *Acantholepis* and *E. gmelini* present a metacentric chromosome pair notably larger than the rest, which is not found in other species (Garnatje & al., 2004; Sánchez-Jiménez & al., 2009). However there exists the doubt that, in the previous molecular phylogenetic reconstruction, the association of the two annual taxa at the base of the tree could be produced

by long-branch attraction, the parsimony method being particularly sensitive to this bias (Philippe & al., 2005). Therefore, the relationship between *Acantholepis* and annual *Echinops* needs to be confirmed. As stated above, the taxonomical delimitation of *Echinops* has not been the object of many doubts and restructurations, and in fact, apart from *E. acantholepis*, the only other species whose position as a member of the genus has been questioned is *E. strigosus*, an Ibero-mauritanian biennial species. According to palynological and leaf features, this species is isolated within the genus (Petit, 1988), which led Tomšovic (1997) to suggest its segregation from *Echinops* to form the monotypic genus *Psectra* (Endl.) Tomšovic. The ITS analysis performed by Garnatje & al. (2005) fully resolves the systematic position of this taxon as nested within *Echinops*, and therefore better considered as *E. strigosus*. Due to the limited species sampling, the previous molecular study did not allow to test the suitability of traditional infrageneric classifications.

The ITS region has been chosen for carrying out the present study because it has provided good results for *Echinops*



**Fig. 1.** Bract diversity in the genus *Echinops*. **A**, plumose bracts of *E. gmelini* (sect. *Nanechinops*); **B**, uniflowered capitulum of *E. strigosus* (sect. *Psectra*) with 8–9 rows of free bracts; **C**, hooked bracts of *E. hoehnelii* (sect. *Hamolepis*); **D**, dissection of *E. koeltzii* Rech. f. (sect. *Echinops*) uniflowered capitulum with the inner bracts free or slightly connate at the base; **E**, dissection of *E. spinosissimus* subsp. *spinosissimus* (sect. *Ritropsis*) uniflowered capitulum with the inner bracts united in a membranous tube; **F**, dissection of *E. elymaiticus* (sect. *Oligolepis*) uniflowered capitulum with the inner bracts connate forming a leather-like tube.

(Garnatje & al., 2005) and other genera of Cynareae (e.g., Susanna & al., 1999; Vilatersana & al., 2000; Wang & al., 2005, 2007; Hidalgo & al., 2006). The *trnL-trnF* region has also been used successfully in Cynareae (e.g., Wang & Liu, 2004; Hidalgo & al., 2006; Wang & al., 2007) and in other groups of Asteraceae (e.g., Kiers & al., 1999; Álvarez & al., 2001; Bayer & al., 2002; Liu & al., 2006; Katinas & al., 2008; Mort & al., 2008). The main goal of this study is to establish a comprehensive molecular phylogenetic framework of *Echinops* in order to (1) discuss whether or not the sections of the genus form natural groups, thus evaluating the suitability of the current classifications, and (2) link the findings to the distribution of some key morphological characters traditionally used in the infrageneric treatments of this genus.

## ■ MATERIALS AND METHODS

**Plant material.** — We analyzed the sequences of the internal transcribed spacers region of nrDNA (ITS1, 5.8S, ITS2) and the *trnL-trnF* region of cpDNA in 89 ingroup specimens, with eight additional specimens for the ITS region and four for the *trnL-trnF*. Eighty-eight taxa corresponding to 81 species and 10 subspecies (three for *E. ritro*, two for *E. sphaerocephalus*, five for *E. spinosissimus*) were represented. We have sequenced a well-rounded representative sample of the genus, which includes the type species of all sections except for the two African sections *Cenchrrolepis* (described as monotypic)

and *Pterolepis* (three species assigned; Fries, 1923). The entire geographic distributional range of the genus is covered, although the tropical African area is still somewhat weakly represented in our sampling (4 of the 25 species considered by Tadesse, 1997). Four species, *Brachylaena discolor*, from tribe Tarchonantheae Kostel, *Cardopatum corymbosum* and *Cousiniopsis atractyloides*, from subtribe Cardopatiinae Less., and *Tugarinovia mongolica*, from subtribe Carlininae (Cass.) Dumort. were chosen as outgroup members according to previous works based on morphological (Petit, 1988) and molecular characters (Susanna & al., 2006). The source of the investigated species is shown in the Appendix in the Electronic Supplement to the online version of this article. Both previously published and new sequences were used in the analysis; the present study contributes 71 new sequences for the ITS region and 89 for the *trnL-trnF* one. The species have been named according to Greuter (2006–2009).

### DNA isolation, PCR amplification and sequencing. —

Total genomic DNA was extracted using a Nucleospin Plant II kit (Macherey-Nagel, GmbH & Co., Düren, Germany) from sheets provided by different herbaria, silica gel-dried material collected during our expeditions, and young plants from germinated cypselas cultivated in the greenhouse of the Botanical Institute of Barcelona (see Appendix in the Electronic Supplement).

The plastid *trnL-trnF* region includes the *trnL* intron, the 3' *trnL* (UAA) exon, and the intergenic spacer between *trnL* (UAA) and *trnF* (GAA), which were amplified and sequenced

**Table 1.** Description and type species of *Echinops* sections according to Bunge (1863), Fries (1923), Rechinger (1979) and Bobrov (1997). Classification according to Rechinger (1979), with sections from outside the Iranian region added (Fries, 1923; Bobrov, 1997). *Echinops* sect. *Psectra* follows Bunge (1863) and Tomšovic (1997). Number of species in *E.* sect. *Echinops*, sect. *Oligolepis* and sect. *Ritropsis* are estimates.

Section	Type	Description
<i>Acantholepis</i> (Less.) Jaub. & Spach	<i>E. acantholepis</i> Jaub. & Spach	Bracts of the common involucre large, external ones leaf-shaped, exceeding and surrounding the head. Annual plants. 1 species.
<i>Chamaechinops</i> Bunge	<i>E. humilis</i> M. Bieb.	Inner involucre bracts free; pappus bristles awn-shaped, remotely toothed above, not bearded. Biennial and perennial plants. 5 species.
<i>Echinops</i>	<i>E. sphaerocephalus</i> L.	Involucre bracts 16–25, the inner free to base or slightly connate. Perennial plants. About 50 species.
<i>Hamolepis</i> R.E. Fr.	<i>E. hoehnelii</i> Schweinf.	Involucre bracts hooked above, inner ones connate. Perennial plants. 1 species.
<i>Hololeuce</i> Rech. f.	<i>E. hololeucus</i> Rech. f.	Involucre bracts 15–20, free, the outer largely plumose. Perennial plants. 1 species.
<i>Nanechinops</i> Bunge	<i>E. nanus</i> Bunge	Involucre bracts 16–20, free, largely plumose, perennial; anther appendages awn-shaped below, bearded above. Annual plants. 3 species.
<i>Oligolepis</i> Bunge	<i>E. leucographus</i> Bunge	Involucre bracts 12–15, in 3 rows, inner involucre bracts united to form a leather-like tube. Perennial plants. About 50 species.
<i>Phaeochaete</i> Bunge	<i>E. longifolius</i> A. Rich.	Involucre bracts up to 25, innermost connate; penicillate bristles complanate, connate. Perennial plants. 3 species.
<i>Psectra</i> Endl.	<i>E. strigosus</i> L.	Involucre bracts free, in 8–9 rows. Perennial plants. 1 species.
<i>Ritropsis</i> Greuter & Rech. f.	<i>E. orientalis</i> Trautv.	Involucres of 16–25 bracts; inner united in a membranous cylindrical tube. Perennial plants. About 25 species.
<i>Terma</i> Endl.	<i>E. exaltatus</i> Schrad.	Inner involucre bracts free; pappus cup-shaped, split only above. Perennial plants. 4 species.

together. Universal primers *trnL*-c, forward, and *trnL*-f, reverse, and, in some cases, *trnL*-d, reverse, and *trnL*-e, forward, were used for amplifying and sequencing the *trnL-trnF* region (Taberlet & al., 1991). The polymerase chain reaction (PCR) procedure included a hot start at 95°C for 1 min 35 s, 34 cycles of 1 min denaturation at 93°C, 1 min annealing at 58°C, 1 min extension at 72°C, and a final 10 min extension at 72°C. The ITS1 spacer, 5.8S gene and ITS2 spacer (ITS region) were amplified and sequenced together. The ITS region was amplified by PCR with the forward primer ITS1, and the reverse primer ITS4 (White & al., 1990), as described in Soltis & Kuzoff (1993). In some cases, we used the 1406F (Nickrent & al., 1994) as forward primer. Sequencing primers ITS1, ITS4 and sometimes ITS2 and ITS3 (reverse and forward, respectively; White & al., 1990) were used.

Both ITS and *trnL-trnF* products were purified with a QIAquick® PCR Purification Kit (Qiagen Inc., Valencia, California, U.S.A.) or DNA Clean & Concentrator™-5 D4004 (Zymo Research, Orange, California, U.S.A.). Direct sequencing of the amplified DNA segments was performed using the BigDye® Terminator Cycle Sequencing v.3.1 (PE Biosystems, Foster City, California, U.S.A.), following the protocol recommended by the manufacturer. Nucleotide sequencing was carried out at the *Serveis Científicotècnics* of the University of Barcelona on an ABI PRISM 3700 DNA analyzer (PE Biosystems, Foster City, California, U.S.A.).

**Phylogenetic analyses.** — Nucleotide sequences were edited and aligned manually with SeaView v.4 (Galtier & al., 1996) and BioEdit v.7.0.9 (Hall, 1999). ITS and *trnL-trnF* analyses were performed both separately and combined. Positions 38–39 and 119–121 of ITS matrix were excluded for the analyses because of their ambiguous alignment.

**Bayesian analyses.** — Datasets were analysed using MrModeltest v.2.3 (Nylander, 2004) to determine the sequence evolution models that best described the present data. These models were used to perform Bayesian analysis using MrBayes v.3.1.2 (Ronquist & Huelsenbeck, 2003), which runs two parallel analyses simultaneously. Four Markov chains were run simultaneously for 106 generations, and these were sampled every 100 generations. Data from the first 1000 generations were discarded as the “burn-in” period, after confirming that likelihood values were stabilized prior to the 1000th generation. The 50% majority rule consensus tree and posterior probability (PP) of nodes were calculated from the remaining 9001 trees sampled.

**Parsimony analysis.** — Heuristic analyses of the ITS and the combined data using PAUP\* v.4.0b10 (Swofford, 2003) failed because of tree storage limitations. As a result, heuristic analysis was carried out with the PAUPRat approach (Nixon, 1999; Sikes & Lewis, 2001), a tool for implementing parsimony ratchet searches using PAUP\*. For parsimony ratchet analyses, uninformative characters were deactivated and 15% of the informative characters were perturbed. The analyses consisted of 10 runs of 200 iterations with tree bisection reconnection (TBR) branch swapping, one tree held at each iteration. On the other hand, parsimony analyses of *trnL-trnF* region involved heuristic searches conducted with PAUP\* using TBR branch swapping with character states specified as unordered

and unweighted. To locate islands of most parsimonious trees (Maddison, 1991), 1000 replicates were performed with random taxon addition. All most parsimonious trees (MPTs) were saved and PAUP\* was used to compute a strict consensus. Tree lengths, consistency index (CI) and retention index (RI) were calculated excluding uninformative characters.

Bootstrap (BS; Felsenstein, 1985) was carried out to obtain support estimates of the nodes of the trees selected. A fast stepwise-addition bootstrap analysis was performed in PAUP\* using 2,000,000 replicates with the default options because of the size of the dataset. The fast stepwise-addition bootstrap method usually provides underestimates as compared to those obtained with branch swapping bootstrap analyses (Mort & al., 2000).

## ■ RESULTS

**Bayesian analyses.** — In some cases Akaike information criterion (AIC) and hierarchical Likelihood Ratio Tests (hLRT) implemented in MrModeltest v.2.3 (Nylander, 2004) determined different models as best fitting the datasets (Table 2). When this occurred, we performed analyses with all models. No inconsistencies were detected between the resulting trees, this leading us to show only the results obtained with the AIC model (Fig. 2), because this approach presents several important advantages over the hLRTs for model selection (Posada & Buckley, 2004).

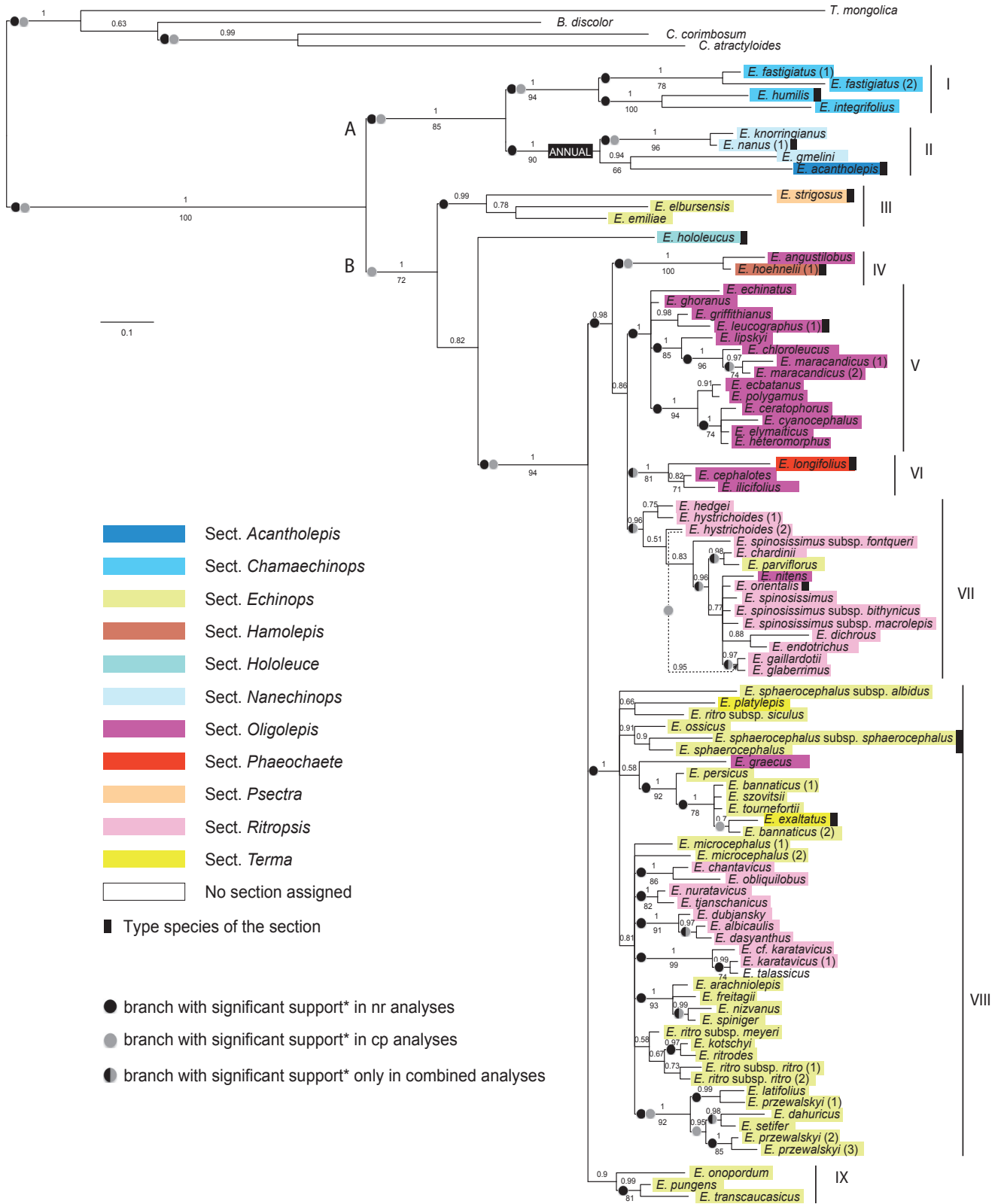
**Parsimony analyses.** — The numerical results of the combined ITS and *trnL-trnF* dataset, as well as for separated regions, are given in Table 2.

**Congruence of the trees.** — Bayesian analyses produced trees with better resolution than parsimony; however they do not show topological discordance for significantly supported

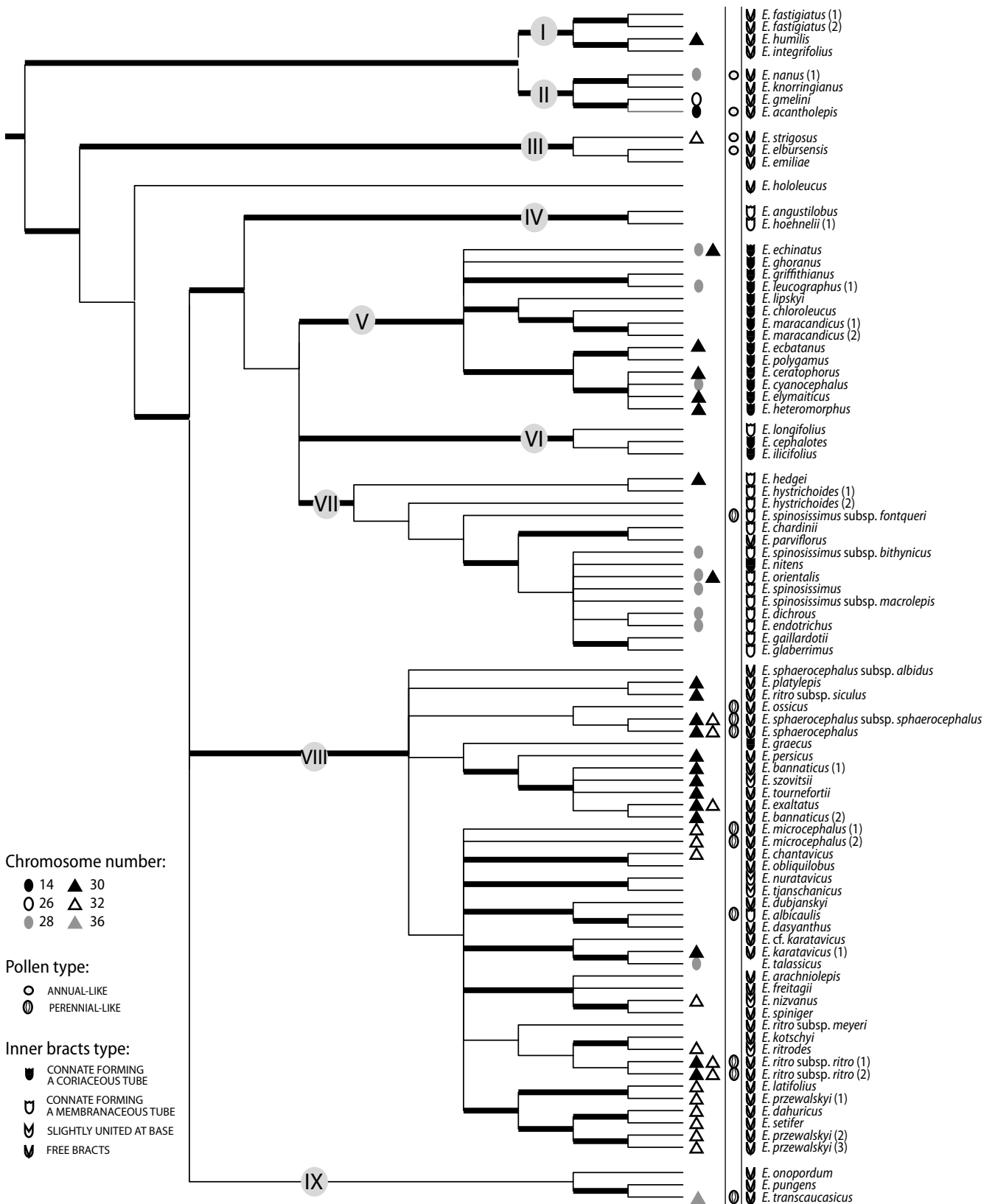
**Table 2.** Statistics of the PAUP\* and Bayesian analyses. Consistency and homoplasy indexes are calculated excluding uninformative characters.

Dataset	Combined <sup>a</sup>	ITS <sup>a</sup>	<i>trnL-trnF</i>	
Ingroup taxa	89	97	93	
Total characters	1519	659	860	
Informative substitutions	240	216	25	
Number of MPTs	1954	1983	32	
Number of steps	681	652	33	
Consistency index (CI)	0.5051	0.4923	0.8182	
Retention index (RI)	0.7643	0.7690	0.9455	
Rescaled consistency index (RC)	0.3861	0.3786	0.7736	
Models selected for Bayesian Inference	AIC	GTR+I+G	GTR+I+G	GTR+G
	hLRT	GTR+I+G	GTR+G	F81+G
	hLRT2	GTR+I+G	SYM+G	F81+I
	hLRT3	GTR+I+G	SYM+I+G	F81+G
	hLRT4	GTR+I+G	SYM+G	F81+I

<sup>a</sup> PAUPRat approach (Nixon, 1999; Sikes & Lewis, 2001) performed.



**Fig. 2.** Majority rule consensus tree resulting from Bayesian analysis of the combined ITS + *trnL-trnF* dataset; numbers above branches are Bayesian posterior probabilities and below branches Bootstrap ( $\geq 70\%$ ). \* indicates  $PP \geq 0.95$  and/or  $BS \geq 80\%$ . Sections are those considered by various authors before this study. Numbers in parentheses after species names differentiate individuals of the same species; A and B denote the two main lineages; roman numerals on the tree indicate the sections as treated in this work. Discontinuous line indicates the incongruence between combined and *trnL-trnF* trees (see text for explanation).



**Fig. 3.** Distribution of selected morphological characters in the phylogeny. See text for bract morphology, Garnatje & Martín (2007) for pollen and Sánchez-Jiménez & al. (2009) and references therein for chromosome numbers. Numbers in parentheses after species names differentiate individuals of the same species; roman numerals on the tree indicate the sections as treated in this work. Thick lines are branches with PP ≥ 0.95.

branches. This led us to present only the tree obtained with the Bayesian analysis of the combined dataset (Fig. 2). Posterior probabilities and bootstrap values are indicated on this tree. Moreover, inspection of topologies and branch support values do not show any conflict between the plastid and nuclear DNA phylogenetic reconstructions. The only inconsistency is between the combined and plastid trees, and concerns the position of *E. hystrioides* (specimen 2). This specimen groups with *E. gaillardotii* and *E. glaberrimus* in *trnL-trnF* analyses (PP = 0.95), while its position in combined analyses is not compatible with such an association. The great compatibility of the nuclear and chloroplastic datasets is also expressed in terms of branch supports, as 20% of supported branches occur only in combined analyses. Notwithstanding, the branch grouping *E. bannaticus* (specimen 2) and *E. exaltatus* in *trnL-trnF* analyses (PP = 0.98) lost significance in combined analyses (PP = 0.70), this indicating a certain level of incongruence, even quite localized.

**Morphological characters.** — The diversity of inner involucre bracts of the unflowered capitula within *Echinops* is distributed into four main categories (Fig. 3): free bracts (FB); bracts slightly united at the base (SUB); connate, forming a membranous tube (CMT); and connate, forming a coriaceous tube (CCT).

## DISCUSSION

### Phylogenetic and taxonomic implications

The new phylogenetic framework established by these analyses shows *Echinops* to be monophyletic (PP = 1.00, BS = 100%; Fig. 2), and confirms previous results pointing towards the inclusion of *E. acantholepis* (= *Acantholepis orientalis*) within the genus (Garnatje & al., 2005). *Echinops* is divided into two main lineages, one comprising the representatives of *E. sect. Chamaechinops*, *sect. Nanechinops* and *E. acantholepis* (lineage A, PP = 1.00, BS = 85%; Fig. 2), the other including the remaining *Echinops* species (lineage B, PP = 1.00, BS = 72%; Fig. 2). The tree provides a valuable resolution at the sectional level, permitting the comparison of the molecular phylogenetic framework with the traditional classifications of the genus. Our results resolved the infrageneric classification of *Echinops* in nine sections, and highlighted the need to do some species relocations to make these sections monophyletic:

***Echinops* sect. *Chamaechinops*.** — (Clade I, PP = 1.00, BS = 94%; Fig. 2) This includes plants restricted to the Tian Shan and Sayan mountains, in northeast Central Asia, which are characterized by pappus bristles that are awn-shaped and remotely serrate above and not bearded (Table 1; Bobrov, 1997).

***Echinops* sect. *Acantholepis*.** — (Clade II, PP = 1.00, BS = 90%; Fig. 2) All the species from *E. sect. Nanechinops* are represented in the analysis and form a monophyletic group along with *E. acantholepis*. This clade is well characterized by involucre bracts with a plumose margin (Fig. 1A; Table 1). Clade II includes the types of two different sections, namely *E. sect. Acantholepis* (*E. acantholepis*) and *sect. Nanechinops* (*E. nanus*), which were described simultaneously (Bunge,

1863), even although Bunge assigned *E. sect. Acantholepis* to Spach because that author had previously described *Echinops* subg. *Acantholepis* (Jaubert & Spach, 1848). In such a case of equal priority, either *E. sect. Acantholepis* or *sect. Nanechinops* can be chosen for the combined section, and we must make a decision (Art. 11.5 of the ICBN; McNeill & al., 2006): **we select here *Echinops* sect. *Acantholepis* and place *Echinops* sect. *Nanechinops* in synonymy under *E. sect. Acantholepis*.**

The close relationship between *E. sect. Acantholepis* and *sect. Chamaechinops* has been suggested on the basis of the likely biennial habit of *E. integrifolius* (Kamelin & Tscherneva, 1971). Bunge (1863) also described both *E. humilis* and *E. integrifolius* as biennial plants. Moreover, Mulkidzhanyan (Bobrov, 1997) stated that “genus *Acantholepis* is a derivative from the genus *Echinops* and seems to have originated from an ancestor of the type *E. integrifolius* and *E. humilis*”. Our results support these assumptions of close affinities between the two sections (lineage A, PP = 1.00, BS = 85%; Fig. 2).

***Echinops* sect. *Psectra*.** — (Clade III, PP = 0.99; Fig. 2) Our phylogenetic trees are not consistent with the segregation of *E. strigosus* in *Psectra strigosa* (L.) Tomšovic (Tomšovic, 1997). The phylogenetic position of this taxon is clearly within *Echinops* in an early-branched lineage (clade III, Fig. 2), as stated in previous work (Garnatje & al., 2005). The relationship between *E. emiliae* and *E. strigosus* is confirmed with significant statistical support, and *E. elbursensis* is added to this group of species.

This group shows an interesting, strongly disjunct distribution. Both *E. elbursensis* and *E. emiliae* are narrow endemics from the Central Alborz mountain range (Iran) and the Antalya mountains (Turkey), respectively (Hedge, 1975; Rechinger, 1979), whereas *E. strigosus* grows below an altitude of 500 m. and is distributed in the north of Africa and the southern parts of the Iberian Peninsula (Valdés, 2002). The so called “Kiermack” disjunctions, between the eastern and western Mediterranean, or even Central Asia and the western Mediterranean (Ribera & Blasco-Zumeta, 1998), have been reported for a number of taxa (Braun-Blanquet & Bolòs, 1957; Davis & Hedge, 1971; Thorne, 1972; Willis, 1996; Oberprieler, 2005; Meerow & al., 2006; Pérez-Collazos & al., 2009). Processes of dispersal and vicariance occurring during the Miocene (15–10 Ma) between the eastern and western Mediterranean are documented for the Asteraceae, in the tribe Anthemideae (Oberprieler, 2005).

***Echinops* sect. *Hololeuce*.** — (Fig. 2) *Echinops hololeucus* was described as constituting the monotypic *E. sect. Hololeuce*, and related to *E. sect. Nanechinops* (in the present classification, *E. sect. Acantholepis*) on the basis of its plumose bracts (Rechinger, 1979). *Echinops hololeucus* appears as isolated within lineage B (Fig. 2), and therefore its affinities should be searched more likely amongst this lineage than with *sect. Acantholepis*, which belongs to lineage A (Fig. 2).

***Echinops* sect. *Hamolepis*.** — (Clade IV, PP = 1.00, BS = 100%; Fig. 2) The African species *E. angustilobus* and *E. hoehnelii* were classified by Fries (1923) in *E. sect. Oligolepis* and *sect. Hamolepis* respectively. The two species are sister in the molecular phylogeny, and consequently the transfer of *E.*

*angustilobus* to sect. *Hamolepis* is required in order to keep the sections monophyletic. *Echinops angustilobus* was the only representative of *E. sect. Oligolepis* having CMT inner involucre bracts, and its new sectional assignment results in two homogeneous sections for bract type (Fig. 3).

***Echinops sect. Oligolepis.*** — (Clade V; PP = 1.00; Fig. 2) This section, characterized by the presence of CCT inner involucre bracts (Rechinger, 1979; Bobrov, 1997; Fig. 1F; Table 1), is well defined in our phylogenetic analysis. *Echinops cornigerus* and *E. kandaharensis*, included only in the ITS analyses, are also located within this clade (data not shown). *Echinops sect. Oligolepis* is practically restricted to the Middle East. In fact, 38 of the 40 species described by Rechinger (1979) for this section in the Iranian flora are endemic.

***Echinops sect. Phaeochaete.*** — (Clade VI, PP = 1.00, BS = 81%; Fig. 2) Although morphological features of *E. cephalotes* and *E. ilicifolius* are clearly attributable to *E. sect. Oligolepis*, these species group with *E. longifolius*, the type of the name of *E. sect. Phaeochaete*. However, this clade VI forms a polytomy with sects. *Hamolepis*, *Oligolepis* and *Ritropsis*, and therefore its merging with *E. sect. Oligolepis*, thus reconciling the molecular and morphological evidences, cannot be discarded.

***Echinops sect. Ritropsis.*** — (Clade VII, PP = 0.96; Fig. 2) *Echinops parviflorus* and *E. nitens*, whose morphological characters are perfectly attributable to the sections to which they were previously assigned (*E. sects. Echinops* and *Oligolepis*, respectively; Fig. 2), are now nested in *E. sect. Ritropsis*. This clade includes *E. spinosissimus*, a polymorphic taxon that has been subject to different taxonomical interpretations (Rechinger, 1943; Feinbrun, 1977; Greuter, 2003). It is mainly distributed in North Africa and the east of the Mediterranean basin, coexisting with several related taxa, like *E. gaillardotii* and *E. glaberrimus* (Rechinger, 1943; Feinbrun, 1977), which are also close phylogenetic relatives (PP = 0.96; Fig. 2). *Echinops tenuisectus* (only on ITS analyses, data not shown) is also included in this *E. spinosissimus* clade. *Echinops spinosissimus* subsp. *fontqueri*, an endemism from the Rif in Morocco (Valdés, 2002) does not group with its presumed conspecifics, which supports its consideration as an independent species, namely *E. fontqueri* Pau. We found the only case of incongruence between combined and plastid analyses for the second specimen of *E. hystrichoides* (see Results; Fig. 2), a species that has been related to *E. spinosissimus* (Tan, 1995). This finding affecting a group with such a taxonomical complexity, which probably results from a recent radiation, may be a product of hybridization.

***Echinops sect. Echinops.*** — (Clade VIII, PP = 1.00; Fig. 2). Clade VIII comprises both types of the names of *E. sect. Echinops* and *E. sect. Terma*. *Echinops sect. Terma* is represented in our study by *E. exaltatus* (the type species) and *E. platylepis*. The two species are both nested within *E. sect. Echinops* representatives and do not group together. Our results agree with Kožuharov (1976) for considering *E. sect. Terma* a synonym of *E. sect. Echinops*. *Echinops sect. Echinops* is characterized by unflowered capitula with 16–25 involucre bracts, the inner ones free or slightly connate at the base (Fig. 1D; Table 1; Rechinger, 1979). Bunge (1863) and Bobrov (1997) considered

an additional section, *E. sect. Ritro* Endl., characterized by external involucre bracts deprived of glandular hairs and leather-like, whereas *E. sect. Echinops* s.str. species have usually glandular external involucre bracts relatively thin. Moreover, species of section *Echinops* s.str. grow in forest edges and shrubby thickets, whereas those of sect. *Ritro* are found in open habitats in steppes and semi-deserts (Bobrov, 1997). The resolution of clade VIII prevents any conclusion as to the suitability of such a classification.

Clade VIII includes a series of eight species previously assigned to *E. sect. Ritropsis* (Fig. 2): *E. albicaulis*, *E. chantavicus*, *E. dasyanthus*, *E. dubjanskyi*, *E. karatavicus*, *E. obliquilobus*, *E. nuratavicus* and *E. tjanschanicus*. These species were also classified in *E. sect. Rythrodes* Bunge (Li, 1987; Bobrov, 1997). Bunge (1863) defined *E. sect. Rythrodes* because of the connation of the inner involucre bracts in a membranous tube, a trait never observed in the type of the name of this section, *E. ritrodes*. This led Rechinger (1979) to transfer this species in *E. sect. Echinops*, and to describe a new section, *E. sect. Ritropsis*, whose type, *E. orientalis*, is characterized by CMT involucre bracts (Table 1). Our results give support to this consideration of *E. sect. Rythrodes* as a synonym of *E. sect. Echinops*. Furthermore, the phylogenetic tree also suggests that other species from *E. sect. Rythrodes* should be placed in *E. sect. Echinops* rather than in *E. sect. Ritropsis*. Morphological evidence reveals that *E. chantavicus*, *E. dasyanthus*, *E. dubjanskyi*, *E. karatavicus* and *E. obliquilobus* have FB (Bobrov, 1997), *E. nuratavicus* and *E. tjanschanicus* SUB (Li, 1987; Bobrov, 1997); and only *E. albicaulis* has CMT (Bobrov, 1997; Fig. 3). Therefore, *E. sect. Rythrodes*, as considered by Bunge (1863) and Bobrov (1997), includes several species that do not present the morphological characters used for defining it. Further, none of them have other frequent characteristics of *E. sect. Ritropsis*, like cornigerous capitula with middle bracts ending in long spines and leaves with large and strong spines. On the contrary, all those characters strongly resemble a general aspect similar to species from section *Echinops*. To sum up, the morphological evidence supports the placement of these species in *E. sect. Echinops*. *Echinops talassicus*, which has until now never been assigned to any section, is included in this group and therefore in *E. sect. Echinops*.

**Clade IX.** — (PP = 0.90; Fig. 2) This clade constitutes one of the few uncertainties of our phylogenetic reconstruction regarding the assignment of species throughout *Echinops* sections. In fact, this is a double uncertainty, because the monophyly of the group in itself is not significantly established, and because there is a trichotomy between clades (IV, V, VI, VII), VIII and IX. If the polytomy is resolved by the grouping of *E. pungens*, *E. onopordum* and *E. transcaucasicus* with clade VIII, these species would remain in *E. sect. Echinops*, as previously stated by morphological data (Hedge, 1975), but if it is resolved in another possible topology, this would mean that these species should probably constitute a new section within the genus. *Echinops polyacanthus* (included only in ITS analyses) groups with *E. pungens* and *E. transcaucasicus* (PP = 0.98 and BS = 81%, data not shown), a result consistent with morphological data (Hedge, 1975).



Our results do not resolve the systematic position of *Echinops onopordum*, an endemic species from southwest Turkey. Nevertheless, they do permit the discarding of a possible relationship of this taxon with the *E. pannosus* group (represented in our phylogeny by *E. emiliae*) suggested by Davis (1956). Its consideration as an isolated species (Hedge, 1975) is still possible.

### Evolution of morphological characters

**Involucral bracts of the unflowered capitula.** — These have been consistently used for sectional characterization, especially regarding their degree of connation (Table 1). The *Echinops* ancestral bract type seems to be that of inner free bracts (FB), present in lineage A, and successively in the grades giving rise to *E. sect. Psectra* and *sect. Hololeuce* at the base of B lineage (Fig. 3). Inner involucral bracts of increasing connection degrees are found later in the tree: bracts slightly united at the base (SUB); connate, forming a membranous tube (CMT); and connate, forming a coriaceous tube (CCT; Fig. 3). Although bract types in themselves do not provide autapomorphies at a sectional level, their distribution throughout *Echinops* sections is quite homogeneous. Each section has only one bract type or at least one clearly dominating type (Fig. 3): FB for *E. sects. Acantholepis, Chamaechinops, Echinops* and *Psectra*, CMT for *E. sects. Hamolepis* and *Ritropsis*, CCT for *E. sect. Oligolepis*. The SUB type is exclusively found in *E. sect. Echinops* (Fig. 3). The association of *sects. Hamolepis, Oligolepis, Phaeochaete* and *Ritropsis* (PP = 0.98, Fig. 2) is supported morphologically; they are characterized by connate inner involucral bracts (of CMT or CCT types) with the only exception of *E. parviflorus* (Rechinger, 1979; Tadesse, 1997).

Some sectional re-locations are quite difficult to explain on morphological bases. This is the case of *E. graecus*, a species with CCT inner bracts previously classified in *E. sect. Oligolepis* (Hedge, 1974; Kožuharov, 1976), which appears in our phylogenetic reconstruction nested in *E. sect. Echinops* (Fig. 2). The observation of unflowered capitula of *E. graecus* at different maturation stages revealed that involucral bracts can be free at floration. We believe that connation of the inner bracts in this and other species may occur during cypselas formation. We agree with Kožuharov (1975) that connation of the bracts should be treated with caution as a differential character, in spite of the fact that it can be consistently found in several groups like *E. sect. Oligolepis* and *sect. Ritropsis*. Nevertheless, since the one-seeded capitulum is the dispersion unit in *Echinops* (Davis, 1956) it is not surprising that evolutionary convergence occurs for this character.

**Pollen types.** — Garnatje & Martín (2007) suggested a close relationship of *E. acantholepis* with other annual *Echinops* species on the basis of pollen type. Both *E. acantholepis* and *E. nanus* have subprolate microechinate pollen without prominent intercolpia (annual-like pollen type), whereas a subprolate microechinate/echinate verrucoid pollen with very prominent intercolpia in the shape of a bridge is present in all the perennial species of *Echinops* examined at this time (perennial-like pollen; Garnatje & Martín, 2007). Recent

pollen morphological work revealed the occurrence of the annual-like pollen type in *E. elbursensis* (I. Sánchez-Jiménez, unpub., Fig. 3), also seen before in *E. strigosus* (Garnatje & Martín, 2007), both from *E. sect. Psectra* (clade III, Fig. 2). Annual pollen type, which seemed at first to be restricted to species with an annual habit (Garnatje & Martín, 2007), is in fact present in *E. acantholepis, E. elbursensis, E. nanus* and *E. strigosus*, and consequently extended to *E. sect. Psectra* in the B lineage (Figs. 2–3). Therefore, annual pollen type can no longer be considered as constituting an autapomorphy of the clade grouping the annuals, but it may represent the ancestral character state for the whole genus.

### CONCLUDING REMARKS

The present study contributes to the establishment of an infrageneric classification of the genus *Echinops*. The sections *Acantholepis, Hamolepis* and *Psectra* were considered before as monotypic, whereas the present phylogenetic study shows they are composed of two or more species. Based on our study, only *E. sect. Hololeuce* remains as monotypic. More work including analysis of other molecular markers is necessary to clarify the phylogenetic relationships of species with doubtful placement. Moreover, sampling should be enlarged to include some more species belonging to the African sections. Although biogeography was not a principal aim of this work, some interesting geographical patterns were detected such as the disjunction within *E. sect. Psectra* and the distributions of *E. sects. Chamaechinops* and *Oligolepis*. Detailed phylogeographical analysis of sections such as *E. sect. Ritropsis* might contribute to elucidate the relationships between the floras of the Mediterranean and the Middle East. The complex taxonomy of the *E. spinosissimus* group could be clarified by means of population studies. A detailed study of pollen morphology and evolution in the genus *Echinops* might also be promising.

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