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An integrative taxonomic study of *Santolina* (Asteraceae) from southern France and north-eastern Spain reveals new endemic taxa

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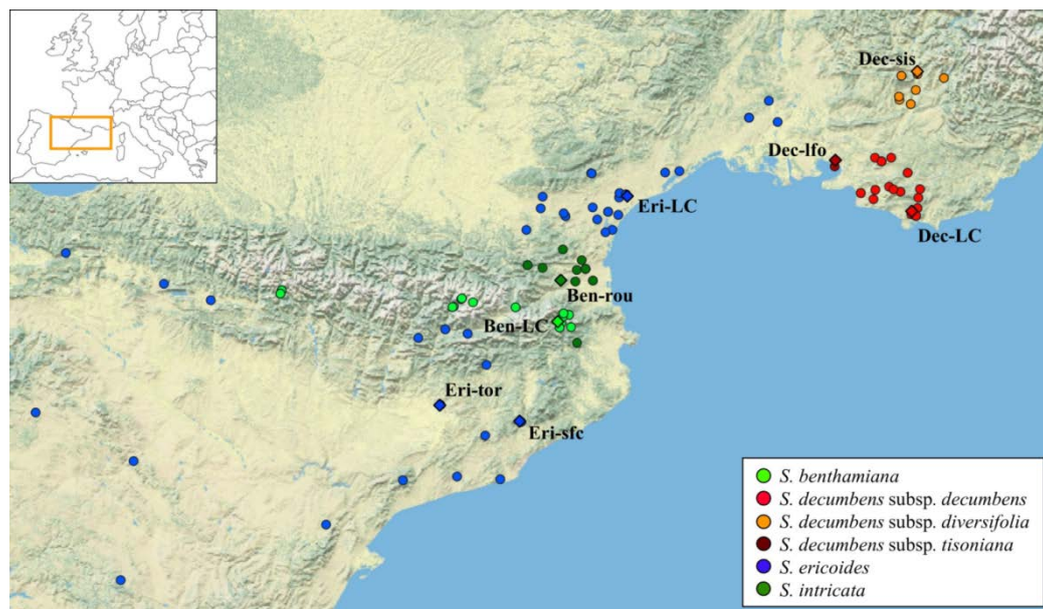
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Abstract: *Santolina* is a clear example of a genus lying in an alpha-taxonomic status, with species accepted only based on qualitative morphological descriptions. In particular, taxonomic issues still need to be resolved for *Santolina* populations from southern France and north-eastern Spain, so that we carried out an integrative taxonomic study involving morphometrics, cypsela morphometrics, niche overlap, and phylogenetic analysis based on six plastid markers (*trnH-psbA*, *trnL-trnF*, *trnQ-rps16*, *rps15-ycf1*, *psbM-trnD*, and *trnS-trnG*). Our results revealed that the current taxonomic circumscription is not adequate. In particular, the *Santolina* populations at the foothills of eastern Pyrenees, previously included in the variability of *S. benthamiana*, have to be considered as a distinct species, namely *S. intricata*. In addition, despite their high phylogenetic relatedness, *S. benthamiana* s.str. and *S. ericoides* can be still considered as distinct species due to clear morphological and ecological differentiation. Finally, we demonstrated that three different subspecies can be recognized in *S. decumbens*, a species endemic to Provence. For one of these subspecies, due to its extremely restricted distribution range, conservation issues are pointed out.

Graphical Abstract

An integrative taxonomic approach involving morphometrics, niche analysis, and molecular systematics with plastid markers (*trnH-psbA*, *trnL-trnF*, *trnQ-rps16*, *rps15-ycf1*, *psbM-trnD*, and *trnS-trnG*) revealed new endemic taxa of *Santolina* in southern France and north-eastern Spain. In particular, *Santolina intricata*, previous considered as synonym of *S. benthamiana*, has to be considered as a distinct species, endemic to eastern Pyrenees. Then, for *Santolina decumbens*, a species endemic to Provence, three subspecies can be recognized. Finally, despite a high phylogenetic relatedness, *S. ericoides* and *S. benthamiana* can be still considered as distinct species due to both morphological and ecological differentiation.



Keywords: Mediterranean Basin, Anthemideae, morphometrics, image analysis, molecular systematics, niche overlap

1. Introduction

In plant taxonomy, the study of morphology is the most traditional and basic approach to infer relationships among taxa (Radford, 1986). Species delimitation is thus based on the observation of qualitative or quantitative diagnostic characters showing scarce or no overlap among the studied taxonomic units (Wiens, 2007). However, morphology alone can be deceptive and can lead scholars to infer conclusions that do not reflect the actual relationships among taxa (Gaudeul et al., 2018; Bateman, 2018; Liu et al., 2022). Despite this, a great number of species is still accepted nowadays based only on qualitative morphological description (Rouhan & Gaudeul, 2014). Turrill (1938) defined this condition as “alpha taxonomy” and hypothesized that future scholars would have experimented new methods to infer taxonomic conclusions. Indeed, cytogenetics, molecular systematics, and also methods based on ecological niche analysis are now commonly used in systematics and taxonomy (Raxworthy et al., 2007; Rouhan & Gaudeul, 2014). In addition, also morphology, the most traditional approach, experimented a renovation thanks to the implementation of advanced statistical tools and classification methods based on machine learning, the so-called morphometry (Rohlf & Marcus, 1993; Moffat et al., 2015). In order to overcome the limits of a taxonomic approach based on a single method, Dayrat (2005) defined the “integrative taxonomy” as “the science that aims to delimit the units of life’s diversity from multiple and complementary perspectives”. Since then, an integrative approach was successfully used in a plenty of studies to untangle the complex relationships within several groups of plants (e.g., Herrando-Moraria et al., 2017; Andriamihaja et al., 2022; Liu et al., 2022,

Tiburtini et al., 2022) and animals (e.g., Vacher 2017; Venkatraman 2019; Sakaguchi 2020).

The *Santolina chamaecyparissus* L. species complex is an example of a group lying in an alpha-taxonomic status, for which an integrated systematic approach is being currently carried out (Giacò et al., 2022; De Giorgi et al., 2022). Except for the cytotoxic studies conducted by Marchi & D'Amato (1973), Marchi et al. (1979), and Valdés-Bermejo & Antúnez (1981), the taxonomic circumscription of species within this complex is mostly based on qualitative morphological observations (Arrigoni, 1982; Tison & de Foucault, 2014; Carbajal et al., 2019). This complex includes 13 subshrub species from the western Mediterranean Basin. In a recent paper, the two traditionally recognized species occurring in Corsica and Sardinia, namely *S. insularis* (Gennari ex Fiori) Arrigoni and *S. corsica* Jord. & Fourr., were investigated. Using different lines of evidence, the authors were able to conclude that, despite the different ploidy level (tetraploid and hexaploid), these two putative species are actually included into the wide morphological variation of a single species (De Giorgi et al. 2022). In a nomenclatural revision of the *S. chamaecyparissus* complex, Giacò et al. (2021) pointed out several taxonomic issues concerning *Santolina* populations from southern France and north-eastern Spain. Indeed, 15 names were validly published for that area. Of all these names, just three are currently accepted, whereas the others are considered as heterotypic synonyms. However, based on the study of type materials, Giacò et al. (2021) concluded that the taxonomic position of some of these synonyms is unclear. The three accepted species are: *S. decumbens* Mill., endemic to Provence (south-eastern France); *S. ericoides* Poir., widely distributed in Occitanie (southern France) and eastern Spain; *S. benthamiana* Jord. & Fourr., endemic to the eastern Pyrenees (France and Spain) (Figure 1). According to Tison et al. (2014) and Carbajal et al. (2019), the distribution ranges of the two latter species may overlap on both sides of the Spanish and French Pyrenees. This article is protected by copyright. All rights reserved.

Moreover, although *S. ericoides* and *S. benthamiana* are respectively recorded for lower and higher altitudes, their altitudinal ranges also overlap. In these putative contact areas, populations with a somewhat intermediate morphology are recorded (Carbajal et al., 2019). Giacò et al. (2021) also pointed out that four names, currently provisionally considered as synonyms of *S. benthamiana*, are based on type specimens that actually show also some morphological features that are otherwise typical of *S. ericoides*. A recent karyomorphological study (Giacò et al., 2022) demonstrated that all the *Santolina* populations native to southern France and north-eastern Spain are diploid with $2n = 18$ chromosomes. Therefore, the current taxonomic delimitation of *Santolina* species from southern France and north-eastern Spain is based exclusively on qualitative morphological observations (Tison & de Foucault, 2014; Tison et al., 2014). The aim of the present study is to test the current three-species taxonomic hypothesis, through an integrated approach already successfully applied in *Santolina* (De Giorgi et al., 2022), involving morphometric analysis of adult plants and cypselae, niche similarity, and phylogenetic analysis (plastid markers *trnH-psbA*, *trnL-trnF*, *trnQ-rps16*, *rps15-ycf1*, *psbM-trnD*, and *trnS-trnG*).

2. Material and methods

2.1 Collection of data

The populations analysed in this study (Table 1) are the same sampled by Giacò et al. (2022) in southern France and north-eastern Spain: the populations from type localities of *S. benthamiana*, *S. decumbens*, and *S. ericoides* (Figure 1); a population collected at the foothills of the French Pyrenees, from an area where populations morphologically intermediate between *S. benthamiana* and *S. ericoides* are reported (Giacò et al., 2021) (Ben-rou in Table 1); a population of *S. decumbens* located at the northernmost portion of

its species range (Dec-sis in Table 1); an isolated population of *S. decumbens* located at the westernmost portion of its species range (Dec-lfo in Table 1); two populations of *S. ericoides* from north-eastern Spain. For morphometrics, 20 flowering individuals were collected for each population. A subset of three individuals was used for the molecular phylogenetic analysis. A set of 100 cypselae for each population was used for the cypselae morphometric analysis. Finally, for the niche analysis, occurrence points were obtained by online databases (<http://flore.silene.eu> and <http://www.anthos.es>, both accessed on the 7th July 2022), herbarium specimens georeferenced with sufficient precision, personal communications, and field investigations.

2.2 Molecular phylogenetic analysis

The same markers used by De Giorgi et al. (2022) were screened for variability (ITS region and the plastid markers *trnH-psbA*, *trnL-trnF*, *trnQ-rps16*, *rps15-ycf1*, *psbM-trnD*, and *trnS-trnG*). As outgroups, *Achillea millefolium* L. from NCBI and two populations of *Santolina villosa* Mill. (the same sampled in the karyo-morphological study by Giacò et al., 2022) were used. For *A. millefolium*, we extracted the abovementioned markers from the plastid genome available on NCBI (accession n° MT500583), while for the two populations of *S. villosa* they were *de novo* amplified and sequenced. DNA was extracted using the kit Exgene™ Plant SV Mini (GeneAll Biotechnology, Singapore). Amplification was carried out with a Polymerase Chain Reaction (PCR) using the primers and amplification conditions listed in Table S1. The following reagents were added in 0.2 mL Eppendorf tubes: 12.5 µL of Kodaq 2X PCR MasterMix (Applied Biological Materials, British Columbia, Canada), 1 µL of forward primer at 10 µM, 1 µL of reverse primer at 10 µM, and the DNA template (0.5–1.5 µL); then, tubes were filled with distilled water up to the volume of 25 µL. PCR products underwent quality checks and purification. The final product (5 µL) was then diluted

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with 8 μ L of distilled water, and, finally, underwent capillary electrophoresis with a 3130 Genetic Analyzer (Applied Biosystems, Massachusetts, USA). The electropherograms were edited with the software Sequence Scanner 2 (Applied Biosystems, Massachusetts, USA), while sequences were edited and aligned with the software Bioedit v. 7.2.5 (Hall, 1999) and Clustal W (Thompson et al., 1994). The sequences were submitted to DDJB (accession numbers available in Table S2). The nuclear markers were not further analysed because of the total absence of variation among sequences. The nucleotide evolution models were computed separately for each of the six plastid markers by using jModelTest ver. 2.1.10 (Darriba et al., 2012) with the corrected Akaike information criterion (Akaike, 1974). A Bayesian analysis was then carried out by using the software MrBayes ver. 3.2.6 (Ronquist et al., 2012), with default values for 2,000,000 generations, sampling chains every 2,000 generations. Convergence and effective sample sizes (ESS) were investigated by using Tracer v.1.7 (Rambaut et al., 2018). The first 10% of the samples was discarded as burn-in. The majority-rule consensus tree was observed in FigTree v1.4.3 (<http://tree.bio.ed.ac.uk/software/figtree/>).

2.3 *Cypselia morphometric analysis*

This approach is based on image analysis and its aim is to detect differences on the external size and shape of diaspores of different populations/groups of populations. It was successfully used in several studies to detect differences among taxonomically critical species groups providing data useful for taxa delimitation (Sarigu et al., 2019; De Giorgi et al., 2022; Tiburtini et al., 2022). A set of 100 cypselae for each population was acquired with a flatbed scanner (Epson Perfection V550) with a digital resolution of 1200 dpi. Before image acquisitions, the scanner was calibrated as suggested by Shahin and Symons (2003). One hundred cypselae were then placed on the scanner. To avoid possible interference of environmental light, images were firstly acquired covering the

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scanner with a white box, and then with a black box. These images were processed using the software package ImageJ v1.53a (Available for free online: <http://rsb.info.nih.gov/ij> (Accessed on 27 June 2022)).

For each cypsela, to extract and analyzed 20 colorimetric and 26 morphometric features and 78 variables of the Elliptic Fourier Descriptors (EFDs), the same procedure described in De Giorgi et al. (2022) and in Tiburtini et al. (2022) was used. In this study, only the morphometric variables were used, since the colorimetric and elliptic Fourier descriptors were found to be scarcely discriminant among populations.

A Pearson correlation test was conducted on the 26 morphometric variables, and highly correlated variables ($r > 0.9$) were discarded. The final dataset was composed of 12 variables. After standardization of data, a Linear Discriminant Analysis (LDA) was conducted using SPSS 16.0 (Statistical Package for Social Science, IBM Corp., Armonk, NY, USA).

2.4 Morphometric analysis

For each individual, 33 characters (29 quantitative and 4 qualitative) were measured (Table 2). Measures were taken on dried material using a digital caliper or imageJ, depending on the character (Table 2). For the latter case, 1200 dpi scans were acquired using a flatbed scanner (Epson perfection 2480 photo). The degree of tomentosity of leaves and stems (fs_hair, ss_hair, fsl_hair, and ssl_hair in Table 2) was obtained after acquisition of HD images of a selected area of leaf or stem using a digital camera (Canon PowerShot S45) mounted on a WILD Heerbrugg M420 stereomicroscope. The degree of tomentosity was then calculated dividing the area covered by tomentum by the total selected area. The character ss_hair (tomentosity of the sterile stem) was categorized as follows: 0–5% (hairless or almost hairless), 6–30% (slightly pubescent), 31–60%

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(pubescent), 61–90% (tomentose), 91–100% (densely tomentose). After measurements, the specimens were conserved at the Herbarium of the Botanic Museum of the Pisa University (PI; herbarium acronyms follow Thiers 2022). Digital images of all the herbarium specimens used in this study can be found at JACQ Virtual Herbaria: <https://www.jacq.org/> (the list of voucher specimens is reported in Table 1).

After standardization of data, a Principal Coordinate Analysis (PCoA) based on Gower distance was carried out to explore the overall morphological variability of populations. The Random Forest (RF) classification method was used to test the correct classification of groups. In particular, we tested with RF the current taxonomic hypothesis and other alternative grouping hypotheses derived by the morphometric and phylogenetic results. RF was successfully used in other taxonomic studies that used a morphometric approach (Moffat et al., 2015; De Giorgi et al., 2022). The main difference and advantage with respect to other classification methods (e.g., LDA) is that qualitative variables can also be included. In addition, it is a non-parametric method, and the classification model is robust even when correlated variables are used (Strobl et al., 2008; Touw et al., 2013). The RF analysis was conducted in R environment using the package “randomForest” (version 4.6-14, Liaw & Wiener, 2002). The morphometric dataset was randomly and equally split in two subsets (training and testing). One hundred iterations of the function “tuneRF” were ran to calculate the optimal number of characters to use at each node. The algorithm was calibrated in order to take into account the covariation among characters. Then, a forest of 800 decision trees was built and a confusion matrix based on the results of RF was obtained. This process was reiterated 100 times randomly splitting each time the training and testing subsets. Finally, a confusion matrix reporting the mean values of classification was calculated.

Univariate analyses comparing populations were conducted on each quantitative character. Homoscedasticity was checked on each character using the Bartlett test. When

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$p > 0.05$, the analysis of variance (ANOVA) was conducted followed by the Tukey-Kramer post-hoc test. Otherwise, Welch t tests were performed for each population pair, using the Hochberg correction to reduce the family-wise error rate. When a significant difference was detected ($p < 0.05$), the index of effect size Cohen's d (Cohen, 1988) was calculated. For characters with unequal variance, an adjusted Cohen's d index was calculated (Aoki, 2020). The Cohen's d index is a measure of the standardized difference between two means (Cohen, 1988). According to Sawilowsky (2009), significant differences with $d < 0.2$ should be considered negligible; instead, when $d > 0.8$, differences are considered as large, and very large if $d > 1.2$. For the purposes of this study, in order to exclude characters with high overlap between population pairs, significant morphological differences were considered relevant only when $d > 1.2$, i.e. when the two means are distant as 1.2 standard deviations.

2.5 Niche analysis

We quantified niche overlap among morphotypes by using the PCA-based method developed by Broennimann et al. (2012). The climatic niche is defined in a multivariate space that is built from real occurrences of taxa. Nineteen bioclimatic variables for current (2000–2016) time slices at about 1×1 km spatial resolution were downloaded from CHELSA v.1.2 dataset (Karger et al. 2017, www.chelsa-climate.org, accessed on the 7th July 2022). We performed Principal Component Analysis (PCA) using “ade4” package in R (Dray & Dufour, 2007) to reduce the transferability issue (Petipierre et al. 2017). The first two axes of PCA were used as new variables. To measure niche overlap, we used the Schoener's D index (1970), which ranges from 0 (no overlap) to 1 (full overlap). We used the niche similarity test to test whether the environmental niche occupied by a morphotype is more similar to the one occupied by another morphotype than would expect at random (Warren et al., 2008). The niche similarity test compares the

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set of environmental conditions occupied by two taxa taking into account the background environmental conditions that are available in both distributional ranges. In brief, the observed niche overlap was compared with the overlap measured between the niche of one morphotype and the niche obtained by randomly sampling (100 times) occurrence points in the background area of the other morphotype. The randomized niche was obtained by randomly shifting the entire observed density of occurrences in one range (the background). We repeated the randomization procedure 100 times and we used a 5-km background area calculated around the points of occurrence. Significant results suggest that the ecological niches of morphotypes are either more or less similar than expected by chance. The analyses were conducted in R (R Core Team, 2019) using the “ecospat” package.

2.6 Distribution of circumscribed taxa

We studied the distribution of the taxa circumscribed at the end of our study by surveying the following online herbaria: AIX, ANG, AUR, BESA, CBPF, CLF, GAP, LIP, LY, LYJB, MARS, MHNM, MPU, NCY, P, SLA, VIL, and VTA (Thiers, 2022). The specimens were re-identified according to the new identification key, georeferenced, and used to build a distribution map.

3. RESULTS

3.1 Molecular phylogenetic analysis

The ITS region was excluded from further analysis since it did not provide any informative site. The marker *trnQ-rps16* (14) shows the highest number of informative sites, followed by *rps15-ycf1* and *trnH-psbA* (9) (Table 3). Conversely, the marker *trnS-trnG* shows the lowest number of informative sites (2). The concatenated matrix was 3485 bp

long and the total number of informative sites is 45 (1.2%). Runs of the Bayesian analysis were mutually converging ($SD < 0.01$) and ESS were all $\gg 200$. The phylogenetic tree based on Bayesian inference is reported in Figure 2. The studied taxa are monophyletic with respect to the selected outgroups. Two well supported clades ($p = 1$) can be observed: one is partly unresolved and consists of the typical *Santolina benthamiana* and the three populations of *S. ericoides*; the second clade consists of *S. benthamiana* from Le Roumenga (Ben-rou), that is sister to the three *S. decumbens* populations.

3.2 *Cypselia* morphometric analysis

Of the 26 morphometric variables considered, 12 not correlated variables were retained (Table S3). The LDA applied to the current taxonomic hypothesis (Table 4) returned a percentage of correct classification of 85.3%. *Santolina decumbens* is well classified (97%), whereas *S. benthamiana* and *S. ericoides* show lower values. Then, we applied the LDA considering each population as a distinct group (Table 5). In this case, the value of correct classification was 53.1%. However, the typical *S. benthamiana* (Ben-LC) is well classified (95.9%), followed by *S. decumbens* from La Fare-les-Oliviers (Dec-lfo) with 86%. All the other populations show lower values and are largely misclassified.

3.3 Morphometric analysis

The first two axes of the PCoA account for 58.68% of the overall morphological variability (Figure 3). The three populations of *S. ericoides* almost completely overlap. The three populations of *S. decumbens* do not overlap, although they are located very close in the multivariate space. The population of *S. benthamiana* from the type locality (Ben-LC) is distinct with respect to *S. ericoides* and *S. decumbens*, whereas the

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population of *S. benthamiana* from Le Roumenga (Ben-rou) overlaps in part with the typical *S. benthamiana*, with *S. decumbens* from Sisteron, and to a lesser extent with the typical *S. decumbens*. RF applied to the current taxonomic hypothesis (Table 6) shows a percentage of mean overall correct a priori classification of 95.2%. However, while *Santolina ericoides* and *S. decumbens* are highly correctly classified by the algorithm (100% and 97.1%, respectively), *S. benthamiana* shows lower values (88.3%) and is mostly misclassified with *S. decumbens*.

We tested alternative grouping hypothesis that can be inferred by the phylogenetic results. If we consider *S. benthamiana* from the type locality (Ben-LC) and *S. ericoides* as belonging to the same group and *S. benthamiana* from Le Roumenga (Ben-rou) as a distinct group, the mean overall value of correct a priori classification drops to 88.5 % (Table 7). The group composed by *S. benthamiana* s.str. + *S. ericoides* is well classified (99.3%), as well as *S. decumbens* (97.8%). However, *S. benthamiana* from Le Roumenga is misidentified with other groups in 31.7% of cases.

Finally, we also tested an alternative grouping hypothesis based on the results of the PCoA (Table 8). Under this hypothesis, each population is considered as a distinct group, with the exception of *S. ericoides*. The percentage of mean correct a priori classification is 91.4%. The typical *S. benthamiana* is well classified (95.6%) and only scarcely misclassified with *S. benthamiana* from Le Roumenga (2.1%) and with *S. ericoides* (2.4%). *Santolina benthamiana* from Le Roumenga (76.5% of mean correct classification) is misclassified with *S. decumbens* from Sisteron (14.1%), with the typical *S. decumbens* (5.9%), and with the typical *S. benthamiana* (3.5%). The three populations of *S. decumbens* are well classified, except the population from Sisteron, that is misclassified for a 7.5% with *S. benthamiana* from Le Roumenga. On the contrary, *Santolina ericoides* is correctly classified with a value of 100%.

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In Table 9, the mean value \pm standard deviation of each character is reported for each population, whereas in Table 10, the number of significant scarcely overlapping characters (Tukey-Kramer or Welch t test with p-value < 0.05 and Cohen's $d > 1.2$) is reported.

The lowest number of scarcely overlapping significant characters was recorded among the three populations of *S. ericoides*, which differ by 2–4 characters (the lowest Cohen's d value is 1.26, while the highest is 1.88). *Santolina benthamiana* from Le Roumenga and the typical *S. benthamiana* differ by 5 scarcely overlapping significant characters (Cohen's d ranging from 1.31 to 3.74), whereas *S. benthamiana* from le Roumenga and *S. decumbens* from Sisteron differ by 6 (Cohen's d ranging from 1.24 to 1.73). The three populations of *S. decumbens* differ each other by 7–10 scarcely overlapping significant characters (Cohen's d ranging from 1.25 to 4.01). The highest number of scarcely overlapping significant characters (17) was recorded between the typical *S. benthamiana* and the typical *S. decumbens* (Cohen's d ranging from 1.3 to 8.45).

3.4 Niche Analysis

The occurrence points used for this analysis were categorized based on the morphotypes detected by morphometric results (Table S4). The niche overlap between each pair of morphotype ranged from no (0.0) to moderate (0.38) overlap (Table 11). The highest value was recorded between the typical *S. benthamiana* and the Ben-rou morphotype. Moderate niche overlap was also detected between *S. ericoides* and *S. decumbens* s.str., Dec-Sis morphotype and Dec-lfo morphotype, and Ben-rou morphotype. The lowest overlap values were detected between the morphotypes of *S. decumbens* and those of *S. benthamiana*, and between Dec-lfo and Dec-sis morphotypes. The significant values in niche similarity test suggested that the ecological niche of *S. ericoides* was generally

more similar to that of Ben-rou, Dec-lfo, and *S. decumbens* s.str. morphotypes than expected by chance, given their environmental backgrounds. In the other cases, the lack of significance in the similarity test suggests differences in optimal niche positions without niche shift.

4. Discussion

As for the *Santolina* populations from Corsica and Sardinia studied by De Giorgi et al. (2022), the ITS region was uninformative. This result suggests a high overall phylogenetic relatedness among all species of the *S. chamaecyparissus* complex and a relatively recent diversification in the Mediterranean context, as already hypothesized by Oberprieler (2005) for this genus. Nevertheless, the six plastid markers allowed to infer the phylogenetic relationships among the diploid taxa native to southern France and north-eastern Spain, that are monophyletic with respect to the polyploid *S. villosa*. This first result fully confirms the taxonomic distinction of *S. villosa* and *S. ericoides* that, before Carbajal et al. (2019) and Giacò et al. (2021), were for a long time included into the variability of a single species, namely *S. villosa* (Greuter, 2008; Tison & de Foucault, 2014; Tison et al., 2014). Based on our results, three main issues need to be discussed more in detail: the taxonomic position of the populations at the foothills of the Pyrenees; the taxonomic distinction of *S. benthamiana* and *S. ericoides*; the evaluation of the infraspecific variability found within *S. decumbens*.

4.1 The taxonomic position of the populations at the foothills of the Pyrenees

The morphology of the populations at the foothills of the Pyrenees was interpreted by Giacò et al. (2021) as intermediate between *S. benthamiana* and *S. ericoides*. Based on the geographic position and qualitative morphological observations, these populations were hypothesized to possibly represent a hybrid zone between these two species

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(Carbajal et al., 2019; Giacò et al., 2021). However, our results do not support this hypothesis. According to the molecular phylogenetic analysis, the population from Le Roumenga (Ben-rou) falls in a different clade with respect to both *S. benthamiana* and *S. ericoides*. Instead, it is sister to *S. decumbens*. The relatedness between *S. benthamiana* from Le Roumenga (Ben-rou) and *S. decumbens* is also supported by the moderate values of niche overlap detected (Table 11). On morphological grounds, the population from Le Roumenga shows a high variability (PCoA in Figure 2). Indeed, RF suggests that individuals of this population can be confused with *S. decumbens* from Sisteron, with the typical *S. decumbens*, and with *S. benthamiana* s.str. (Table 10). However, thanks to the univariate analyses, we detected that the population from Le Roumenga shows a peculiar combination of diagnostic characters. With respect to the typical *S. benthamiana* and *S. decumbens* from Sisteron, plants from Le Roumenga show, respectively, a higher degree of tomentosity and a different leaf morphology (Table 10). Based on these evidence, the populations at the foothills of the eastern Pyrenees can be considered as a distinct species, for which the name *Santolina intricata* Jord. & Fourr. is available. This name was published by Jordan & Fourreau (1869) based on material from Caudies (Pyrenées Orientales, France), just 25 km far from Le Roumenga. Carbajal et al. (2019) hypothesized that such kind of populations (here referred to *S. intricata*) may occur also in the Spanish side of the Pyrenees. We were indeed able to detect a population from the Spain whose morphology fits well with *S. intricata* (Figure 4 and Table S5), albeit more field and herbarium investigations are needed to have a more detailed picture of the distribution of this species in north-eastern Spain. Compared to other Mediterranean mountain chains, the Pyrenees host a lower percentage of endemic species, and the area of occurrence of *S. intricata* is one of the poorest (Gómez et al., 2017). According to the latter authors, besides biogeographic, climatic, and ecological reasons, this can be due to the exclusion of putative endemic species whose taxonomy was unresolved. Therefore, This article is protected by copyright. All rights reserved.

the recognition of *S. intricata*, whose ‘intricate’ morphology has tricked previous taxonomists, is an important enrichment for the flora of the eastern Pyrenees.

4.2 The taxonomic distinction of *Santolina benthamiana* and *S. ericoides*

Our phylogenetic analysis did not resolve the clade that includes *S. benthamiana* and *S. ericoides*. Traditionally, these two species have been considered as well distinct due to the different leaf morphology (Tison & de Foucault, 2014; Tison et al., 2014; Carbajal et al., 2019). Our morphometric univariate analyses confirm previous observations (Table 10), and the RF never failed to correctly classify these two species. *Santolina benthamiana* and *S. ericoides* are distinct also concerning cypselae morphometry. Indeed, the cypselae of *S. benthamiana* were correctly classified by the LDA with high percentages of correct classification, and only the topotypical population of *S. ericoides* was slightly misclassified (3.4%) with *S. benthamiana*. Finally, the relatedness between these two species is supported also by climatic niche analysis that suggested they have a moderate niche overlap (Table 11). Based on this evidence, we think that *S. benthamiana* and *S. ericoides* could be still considered as distinct species. Regarding the high affinity detected by our molecular phylogenetic analysis, we can speculate that *S. benthamiana* is a species that just very recently differentiated from the widely distributed *S. ericoides*. The adaptation to a distinct environment at, on average, higher elevations (0–1800 m a.s.l. for *S. ericoides* and (600)1500–2000 for *S. benthamiana*) (Tison & de Foucault, 2014; Carbajal et al., 2019) may have contributed to the acquisition of a different morphology. A similar pattern was detected by Fréville et al. (1998) for *Centaurea stoebe* L. (= *C. maculosa* Lam. subsp. *maculosa*; Asteraceae), a widespread Mediterranean species, and *C. corymbosa* Pourr., a narrow endemic species occurring in Occitanie, southern France. According to these authors, the latter species may have differentiated from the former after a colonization event, and the new local environmental conditions

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may have contributed to the morphological differentiation. Another possible evolutionary interpretation may involve a chloroplast capture during a hybridization event and subsequent backcrosses. In this perspective, a phylogeny of the whole genus *Santolina* using NGS techniques is currently ongoing and will likely provide further data to better understand the relationships among these two related species.

4.3 *Intraspecific variability within Santolina decumbens*

The monophyly of *S. decumbens* is well supported by our phylogenetic analysis (Figure 2). According to Tison & de Foucault (2014) and Tison et al. (2014), this species can be easily distinguished by other taxa for the high degree of tomentosity on stems and leaves. However, we detected that the three studied populations, located at the extremes of the distribution of this species, significantly differ from each other on morphological grounds. Indeed, when considered as distinct groups, RF correctly classified these populations. Thanks to the univariate analyses, we detected that each population can be distinguished by a combination of diagnostic characters. Indeed, the toptypical population, located at the southernmost portion of the species range, is the most tomentose, showing white stems and leaves, the fertile branches are short as well as the leaves. The population from Sisteron, located at the northernmost portion of the species range, shows less tomentose fertile stems and leaves, which are also longer. Finally, the population from La Fare-les-Oliviers, located at the westernmost portion of the species range, when compared to the toptypical population shows less tomentose fertile stems and leaves, the leaves are shorter, and cypselae can be easily distinguished, too (Table 10). According to the niche analysis, these three morphotypes occur in different climatic conditions, suggesting that the observed morphological differences may be due to difference in environmental conditions. Based on our results, we believe that it is preferable to treat these groups as distinct subspecies (Hamilton & Richard, 1992). For

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the “Sisteron morphotype” a name at species level is available, i.e. *Santolina diversifolia* Jord. & Fourr., that can be recombined at subspecific level. Concerning the “La Fare-les-Oliviers morphotype”, no previously published name is available, so that it is described here as a subspecies new to science. Serious conservation issues must be pointed out for this latter taxon. Indeed, it occurs in a restricted area around Bouches-du-Rhône. This area is strongly anthropized and the natural environments are crucially affected by urbanization, industrial activities, and agriculture (Tatoni et al., 2004). Approximately in the same area, *Teucrium pseudochamaepitys* L. (Lamiaceae) occurs. It is another southern French narrow endemic species that, according to Lhotte et al. (2014), is strongly threatened by urbanization and habitat fragmentation. Another case of geographically close rare species is *Arenaria provincialis* Chater & G.Halliday (Caryophyllaceae), which is endemic to the hills around Marseille, and it is “near threatened” according to IUCN (Véla et al., 2008). Based on our field observations and the literature concerning endangered species occurring approximately in the same area, a conservation status assessment, possibly involving analyses of fitness and reproductive biology, is urgent for this subspecies.

In conclusion, the integrative taxonomic approach carried out in this study allowed to reveal a biological diversity in *Santolina* diploids from southern France and north-eastern Spain that was not detected by previous qualitative taxonomic investigations. The *Santolina* populations at the foothills of Pyrenees that were previously interpreted as “intermediate” between *S. benthamiana* and *S. ericoides* belong actually to a distinct species (i.e. *S. intricata*). According to our herbarium research, this species occurs at both sides of French and Spanish Pyrenees (Figure 4). For *S. decumbens*, a species endemic to Provence, three different isolated lineages (Figure 4) worth of the subspecies rank were detected. Finally, concerning *S. benthamiana* and *S. ericoides*, despite a high phylogenetic relatedness, they can preserve the status of species thanks to clear

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morphological and ecological differentiation. The patterns detected in this study are in accordance with the rich literature available for recently diversified plant species in the Mediterranean Basin (De Castro et al., 2015; Bateman, 2018; Liu et al., 2022; Terlević et al., 2022). Indeed, the Mediterranean region is one of the most important hotspots for biodiversity, and the factors that have driven plant speciation are many and are related to its complex climatic and geologic history (Greuter, 1991; Cañadas et al., 2014). In the case of *Santolina*, the current diversity can be explained by both biological and geological/climatic phenomena. Concerning biological factors, the role of polyploidy is one of the main factors that has driven the diversification of lineages currently occurring in Corsica, Sardinia, and the Iberian Peninsula (Rivero-Guerra, 2008a; Rivero-Guerra, 2008b; Giacò et al., 2021). However, this is not the case of the taxa studied here, since they are all diploid. Previous scholars (Arrigoni, 1979; Torricelli et al., 2000) suggested that diploid *Santolina* species may be interpreted as schizoendemics (Siljak-Yakovlev & Peruzzi, 2012), i.e. taxa originated by the fragmentation of a common, more widespread, ancestor. A phylogenomic study involving NGS techniques is currently ongoing on the whole genus, in order to understand the general patterns of diversification and to address biogeographical questions.

Taxonomic treatment

Santolina benthamiana Jord. & Fourr., Icon. Fl. Eur. 2: 10. 1869 \equiv *Santolina pectinata* Benth., Cat. Pl. Pyrénées: 117. 1826, nom. illeg., non Lag. 1816 \equiv *Santolina chamaecyparissus* subsp. *pecten* Rouy, Fl. France 8: 224. 1903 \equiv *Santolina chamaecyparissus* var. *pectinata* Fiori in Fiori & al., Fl. Italia 3(1): 270. 1903 – Lectotype designated by Giacò et al. (2021: 191): France, Occitanie, Pyrénées-Orientales, Prats-de-Mollo, inter Arles et Prats-de-Mollo, 11 Jun 1825, *G. Bentham* 573 (K barcode K000929471!).

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= *Santolina chamaecyparissus* subsp. *pecten* var. *ruscinonensis* Rouy, Fl. France 8: 225. 1903 – Lectotype designated by Giacò et al. (2021: 193): France, Occitanie, Pyrénées-Orientales, Prats-de-Mollo, Jun 1846, A. *Irat s.n.* (LY barcode LY0003145!; isolectotypes: LY barcode LY0341058!, P barcodes P00752629!, P00752630!, P00752631!, P00752632!, P00752633!, P00752636!, P00752673!, P00752674! & P03315514!).

= *Santolina chamaecyparissus* subsp. *pecten* var. *hispanica* Rouy, Fl. France 8: 225. 1903 – Lectotype by Giacò et al. (2021: 193): Spain, Aragon, Broto, Jul 1873, A. *Autheman s.n.* (LY barcode LY0715656!).

Distribution: Spanish and French central-eastern Pyrenees (Fig. 4; Table S5).

Santolina decumbens Mill., Gard. Dict., ed. 8: *Santolina* no. 3. 1768 – Neotype designated by Giacò et al. (2021: 194): France, Provence-Alpes-Côte d’Azur, Mont Caume, 23 Jun 1972, L. *Mercurin s.n.* (P barcode P00113904!).

subsp. ***decumbens***

= *Santolina incana* Lam., Fl. Franç. 2: 43. 1779 ≡ *S. chamaecyparissus* var. *incana* (Lam.) DC., Prodr. 6: 35. 1838 – Neotype designated by Giacò et al. (2021: 196): France, Occitanie, Aude, Narbonne [likely cultivated], s.d., *s.coll.* (P barcode P06898681!).

= *Santolina villosissima* Poir. in Lamarck, Encycl. 6: 505. 1805 ≡ *Santolina chamaecyparissus* var. *villosissima* (Poir.) DC., Prodr. 6: 35. 1838 – Neotype designated by Giacò et al. (2021: 200): France, Provence-Alpes-Côte d’Azur, Montagne Sainte-Victoire, 29 Jun 1906, J.-J. *Paulet s.n.* (LY barcode LY0340523!).

Distribution: southern France (Provence), from Marseille eastwards to Toulon and northwards up to Montagne de Sainte-Victoire (Fig. 4; Table S5).

subsp. *diversifolia* (Jord. & Fourn.) Giacò & Peruzzi, **comb. nov.**

Basionym: *S. diversifolia* Jord. & Fourn., Icon. Fl. Eur. 2: 9. 1869 \equiv *S. chamaecyparissus* var. *incana* subvar. *diversifolia* (Jord. & Fourn.) Rouy, Fl. France 8: 222. 1903 \equiv *Santolina chamaecyparissus* var. *diversifolia* (Jord. & Fourn.) Guinea in Anales Inst. Bot. Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. 2021: 195): France, Provence-Alpes-Côte d’Azur, Montagne de Lure, 5 Jun 1864, A. Jordan *s.n.* (LY barcode LY0826364; isoelectotypes: LY barcodes LY0826365, and LY0826366).

Distribution: southern France, Alpes-de-Haute-Provence (Fig. 4; Table S5).

subsp. *tisoniana* Giacò & Peruzzi, **subsp. nov.**

It is similar to *S. decumbens* s.str., but the fertile stems are not white-tomentose. With respect to *S. decumbens* subsp. *diversifolia*, all vegetative parts are distinctly shorter (fertile stems < 15 cm long, leaves of the sterile and fertile stems < 15 mm long). The leaf morphology reminds of *S. ericoides*, but the leaves of the sterile stems are densely tomentose, whereas in *S. ericoides* are almost glabrous. Named after Jean-Marc Tison, respected scholar of the French flora.

Holotype (here designated): France, Provence-Alpes-Côte d’Azur, La Fare-les-Oliviers, small spots around the aerodrome (WGS84: 43.539610 N, 5.172029 E), about 50 m, 28 June 2020, A. Giacò & L. Peruzzi *s.n.* (PI barcode PI043100; isotypes: PI barcodes PI043099, PI043101, PI043102, PI043103, PI043104, PI043105, PI043106).

Distribution: southern France (Provence), between Berre-L'Étang and Lançon-Provence (Fig. 4; Table S5).

Santolina ericoides Poir. in Lam., Encycl. 6: 504. 1805 ≡ *Santolina chamaecyparissus* var. *squarrosa* subvar. *ericoides* (Poir.) Rouy, Fl. France 8: 223. 1903 ≡ *Santolina chamaecyparissus* subsp. *ericoides* (Poir.) Sennen & Elias in Bol. Soc. Ibér. Ci. Nat. 28(1–2): 32. 1929 – Lectotype designated by Ferrer-Gallego et al. (2021: 235): [illustration] “ABROTONUM foemina, Dodonaei” in Daléchamps, Hist. Gen. Pl.: 938. 1586.

= *Santolina brevifolia* Jord. & Fourr., Icon. Fl. Eur. 2: 11. 1869 ≡ *Santolina chamaecyparissus* var. *incana* subvar. *brevifolia* (Jord. & Fourr.) Rouy, Fl. France 8: 222. 1903 ≡ *Santolina chamaecyparissus* var. *brevifolia* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. (2021: 191): [illustration] “*Santolina brevifolia* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 234. 1869.

= *Santolina chamaecyparissus* var. *mariolensis* O.Bolòs & Vigo in Collect. Bot. (Barcelona) 17(1): 90. 1987 – Holotype: Spain, Valencian Community, Serra de Mariola, 520 m, s.d., A. Bolòs & O. Bolòs s.n. (BC barcode BC-150851!).

= *Santolina chamaecyparissus* var. *virens* Willk., Prodr. Fl. Hispan. 2(1): 80. 1865 – Lectotype designated by Giacò et al. (2021: 194): Spain, Madrid, Alcalá de Henares, dans champs pirreux incultes, 27 Jun 1854, J. Gay s.n. (COI barcode COI00035925!).

= *Santolina glabrescens* Jord. & Fourr., Icon. Fl. Eur. 2: 11. 1869 ≡ *Santolina chamaecyparissus* var. *squarrosa* subvar. *laxa* Rouy, Fl. France 8: 223. 1903 ≡ *Santolina chamaecyparissus* var. *glabrescens* (Jord. & Fourr.) Guinea in Anales Inst. Bot.

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Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. (2021: 195): [illustration] “*Santolina glabrescens* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 233. 1869.

= *Santolina homophylla* Jord. & Fourr., Icon. Fl. Eur. 2: 8. 1869 ≡ *Santolina chamaecyparissus* var. *incana* subvar. *homophylla* (Jord. & Fourr.) Rouy, Fl. France 8: 223. 1903 ≡ *Santolina chamaecyparissus* var. *homophylla* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 41. 1970 – Lectotype designated by Giacò et al. (2021: 196): [illustration] “*Santolina homophylla* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 223. 1869.

= *Santolina microcephala* Jord. & Fourr., Icon. Fl. Eur. 2: 11. 1869 – Lectotype designated by Giacò et al. (2021: 197): Spain, Castilla La Mancha, Hellin, Collines a Hellin, 24 May 1850, *J. Gay s.n.* (LY barcode LY0826375!).

Distribution: widespread from north-eastern Spain up to Nîmes in southern France (Fig. 4; Table S5).

***Santolina intricata* Jord. & Fourr.**, Icon. Fl. Eur. 2: 11. 1869 ≡ *Santolina chamaecyparissus* var. *intricata* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. (2021: 196): [illustration] “*Santolina intricata* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 232. 1869.

= *Santolina brevicaulis* Jord. & Fourr., Icon. Fl. Eur. 2: 10. 1869 ≡ *Santolina chamaecyparissus* var. *incana* subvar. *brevicaulis* (Jord. & Fourr.) Rouy, Fl. France 8: 222. 1903 ≡ *Santolina chamaecyparissus* var. *brevicaulis* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 41. 1970 – Lectotype designated by Giacò et al. (2021:

191): [illustration] “*Santolina brevicaulis* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 230. 1869.

= *Santolina rigidula* Jord. & Fourr., Icon. Fl. Eur. 2: 10. 1869 ≡ *Santolina chamaecyparissus* var. *rigidula* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. (2021: 198): France, Occitanie, Pyrénées-Orientales, Perpignan, Estagel, Jun 1838, *s.coll.* (LY barcode LY0826378!; isolectotypes: LY barcodes LY0341579! & LY0341580!).

= *Santolina valida* Jord. & Fourr., Icon. Fl. Eur. 2: 9. 1869 ≡ *Santolina chamaecyparissus* var. *incana* subvar. *valida* (Jord. & Fourr.) Rouy, Fl. France 8: 223. 1903 ≡ *Santolina chamaecyparissus* var. *valida* (Jord. & Fourr.) Guinea in Anales Inst. Bot. Cavanilles 27: 42. 1970 – Lectotype designated by Giacò et al. (2021: 199): [illustration] “*Santolina valida* Jord. et Fourr.” in Jordan & Fourreau, Icon. Fl. Eur. 2: t. 224. 1869.

These four names were published simultaneously at species level by Jordan & Fourreau (1869) for the distribution area typical of the “Le Roumenga” morphotype. We opt here for *Santolina intricata* as having priority over the competing *S. brevicaulis*, *S. rigidula*, and *S. valida* (Art. 11.5 of the ICN; Turland et al. 2019).

Distribution: southern France, Pyrénées-Orientales and Girona province in Spain (Fig. 4; Table S5).

Identification key for diploid *Santolina* native to France and north-eastern Spain

1. Leaves of the sterile stems, except for the ones at the tips, green and almost glabrous (tomentose at least on the leaf axis)

.....

..... 2

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1. Leaves of the sterile stems tomentose, grey or white
..... 3
2. Leaves of the sterile stems with 0.8–2.2 mm spaced out segments, (1.5)2–8 mm long
..... *S. benthamiana*
2. Leaves of the sterile stems with 0.2–0.8 mm spaced out segments, usually shorter than
2 mm..... *S. ericoides*
3. Leaves of the sterile stems with 25–65, 0.8–2 mm spaced out, segments. Segments of
the fertile stem leaves 1.5–3 mm long and 0.5–1.5 mm spaced-out
..... *S. intricata*
3. Leaves of the sterile stems with 50–80, < 1 mm spaced out, appressed segments.
Segments of the fertile stem leaves 0.5–2 mm long and 0.0–1.0 mm spaced-out
..... 4 (*S. decumbens*)
4. Fertile stems white, tomentose as (or almost as) the sterile stems *S.*
decumbens subsp. *decumbens*
4. Fertile stems green, clearly less tomentose than the sterile stems
..... 5
5. Longest fertile stems 12–30 cm long. Longest leaves of the sterile stems 15–35 mm
long *S.*
decumbens subsp. *diversifolia*
5. Longest fertile stems 5–15 cm long. Longest leaves of the sterile stems 8–15 mm long
..... *S.*
decumbens subsp. *tisoniana*

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Table 1. Sampled *Santolina* populations from southern France and north-eastern Spain and related voucher information. LC = type locality (*locus classicus*). The taxonomic assignment of populations follows Giacò et al. (2021).

Species	Population	Acronym	Vouchers
<i>S. benthamiana</i>	France, Occitanie, Prats-de-Mollo-la-Preste [WGS84: 42.407222 N, 2.523055 E]	Ben-LC	A. Giacò and L. Peruzzi, 29 June 2020, PI 043080–043098
<i>S. benthamiana</i>	France, Occitanie, Montalba-le-Château, Le Roumenga [WGS84: 42.699054 N, 2.552235 E]	Ben-rou	A. Giacò and L. Peruzzi, 28 June 2020, PI 043079, PI 057098–057114
<i>S. decumbens</i>	France, Provence-Alpes-Côte d'Azur, Mont Caume [WGS84: 43.184768 N, 5.908187 E]	Dec-LC	A. Giacò and L. Peruzzi, 27 June 2020, PI 043107–043118
<i>S. decumbens</i>	France, Provence-Alpes-Côte d'Azur, Sisteron [WGS84: 44.153341 N, 5.953744 E]	Dec-sis	A. Giacò and L. Peruzzi, 11 July 2021, PI 053348–053364
<i>S. decumbens</i>	France, Provence-Alpes-Côte d'Azur, La Fare-les-Oliviers [WGS84: 43.539610 N, 5.172029]	Dec-lfo	A. Giacò and L. Peruzzi, 28 June 2020, PI 043099–043106
<i>S. ericoides</i>	France, Occitanie, Béziers [WGS84: 43.28959 N 3.18539 E]	Eri-LC	A. Giacò and L. Peruzzi, 28 June 2020, PI 036086–036100
<i>S. ericoides</i>	Spain, Barcelona province, Sant Feliu de Codines [WGS84: 41.692294 N, 2.174761 E]	Eri-sfc	L. Sáez, 7 July 2020, PI 043077, PI 057135–057154
<i>S. ericoides</i>	Spain, Lleida province, Torà [WGS84: 41.814325 N, 1.404588 E]	Eri-tor	L. Sáez, 13 July 2020, PI 043076, PI 057115–057134

Table 2. Morphometric characters and their description for *Santolina* native to France and south-eastern Spain. QC = quantitative continuous, QD = quantitative discrete, CO = ordered factor, and Y/N = bimodal.

Code	Description of the Character	Type	Tool
Vegetative parts			
fs_len	Length of the fertile stem (cm)	QC	Ruler
br_ratio	Ratio between the terminal ramification of the fertile stem and fs_len	QC	Ruler
dist_cap	Distance between the terminal leaf on the stem and the floral head (mm)	QC	Caliper
fs_n_br	Number of branches of the fertile stem	QD	
fs_n_nodes	Number of nodes of the fertile stem	QD	
ss_len	Length of the sterile stem (cm)	QC	Ruler
ss_n_nodes	Number of nodes of the sterile stem	QD	
ss_bicolor	Sterile stems densely tomentose on the upper portion, less tomentose at the bottom	Y/N	
ss_hair	Tomentosity of the sterile stem	CO	ImageJ

fs_hair	Degree of tomentosity of the fertile stem (%)	QC	ImageJ
fsl_n_seg	Number of segments on the fertile stem leaf (the longest)	QD	
ssl_n_seg	Number of segments on the sterile stem leaf (the longest)	QD	
ssl_len	Length of the sterile stem leaf (mm)	QC	ImageJ
ssl_pet_len	Length of the petiole of the sterile stem leaf (mm)	QC	ImageJ
ssl_seg_len	Length of the segment of the sterile stem leaf (mm)	QC	ImageJ
ssl_seg_dist	Distance between the segments of the sterile stem leaf (mm)	QC	ImageJ
fsl_len	Length of the fertile stem leaf (mm)	QC	ImageJ
fsl_pet_len	Length of the petiole of the fertile stem leaf (mm)	QC	ImageJ
fsl_seg_len	Length of the segment of the fertile stem leaf (mm)	QC	ImageJ
fsl_seg_dist	Distance between the segments of the fertile stem leaf (mm)	QC	ImageJ
ssl_hair	Degree of tomentosity of the sterile stem leaf segment (%)	QC	ImageJ
fsl_hair	Degree of tomentosity of the fertile stem leaf segment (%)	QC	ImageJ
Floral head			
cap_diam	Diameter of the floral head involucre (mm)	QC	Caliper
fl_len	Length of the floral tube (mm)	QC	ImageJ
fl_tooth_len	Length of the floral tooth (mm)	QC	ImageJ
sq_ext_len	Length of the external involucre bract (mm)	QC	ImageJ
sq_ext_wid	Width of the external involucre bract (mm)	QC	ImageJ
sq_int_len	Length of the internal involucre bract (mm)	QC	ImageJ
sq_int_wid	Width of the internal involucre bract (mm)	QC	ImageJ
sq_if_len	Length of the inter-floral bract (mm)	QC	ImageJ
sq_if_wid	Width of the inter-floral bract (mm)	QC	ImageJ
sq_if_n_hair	Tomentosity of the inter-floral bract (hairless/slightly pubescent/pubescent/hairy/densely hairy)	QD	ImageJ
sq_ext_hair	Tomentosity of the external involucre bract (hairless/only on the margin/everywhere)	CO	ImageJ
sq_int_hair	Tomentosity of the internal involucre bract (hairless/only on the margin/everywhere)	CO	ImageJ

Table 3. Length, consensus length, and number of informative sites of the six markers in the studied *Santolina* populations. Lengths refers to the ingroup.

Marker	Length (bp)	Consensus length (bp)	Informative Sites	Inferred DNA evolution model
<i>psbM-trnD</i>	714–715	721	7	GTR
<i>rps15-ycf1</i>	500–510	559	9	F81
<i>trnF-trnL</i>	381	386	4	HKY
<i>trnH-psbA</i>	420–429	463	9	HKY+I
<i>trnQ-rps16</i>	885–895	931	14	F81+G
<i>trnS-trnG</i>	420–425	425	2	F81

Table 4. Cypsela morphometric analysis. Results of the LDA applied to the current taxonomic hypothesis for *Santolina* native to France and south-eastern Spain. Values are percentages.

	<i>S. benthamiana</i>	<i>S. decumbens</i>	<i>S. ericoides</i>
<i>S. benthamiana</i>	85.5	7.8	6.7
<i>S. decumbens</i>	1.3	97	1.7
<i>S. ericoides</i>	6.8	19.8	73.4

Table 5. Cypsela morphometric analysis for *Santolina* native to France and south-eastern Spain. Results of the LDA applied considering each population as a distinct group. Values are percentages. Ben-LC = *S. benthamiana* from type locality, Ben-rou = *S. benthamiana* from Le Roumenga, Dec-LC = *S. decumbens* from type locality, Dec-lfo = *S. decumbens* from La Fare-les-Oliviers, Dec-sis = *S. decumbens* from Sisteron, Eri-LC = *S. ericoides* from type locality, Eri-fsc = *S. ericoides* from Sant Feliu de Codines, Eri-tor = *S. ericoides* from Torà.

	Ben-LC	Ben-rou	Dec-LC	Dec-lfo	Dec-sis	Eri-LC	Eri-sfc	Eri-tor
Ben-LC	95.9	0	0.9	0	2.9	0.2	0	0.1
Ben-rou	0.3	35.8	12	5.7	8.3	13.3	8.3	16.3
Dec-LC	5.5	13.9	41.7	0.5	22.3	9.6	2	4.4
Dec-lfo	0	1.4	0	86	3.3	0.1	2.3	6.9
Dec-sis	0.2	13.1	23.2	4.4	35.7	13	6.8	3.8
Eri-LC	3.4	12.5	5.8	0.7	12.1	34.6	23	7.9

Eri-sfc	0	11.9	2.5	2.1	5.4	24.3	43.6	10.2
Eri-tor	0	6.5	6.8	12.1	8.9	5.1	8.8	51.7

Table 6. Morphometric analyses for *Santolina* native to France and south-eastern Spain. Results of the Random Forest applied to the current taxonomic hypothesis. Values are percentages.

	<i>S. benthamiana</i>	<i>S. decumbens</i>	<i>S. ericoides</i>
<i>S. benthamiana</i>	88.3	11.3	0.3
<i>S. decumbens</i>	2.9	97.1	0
<i>S. ericoides</i>	0	0	100

Table 7. Morphometric analysis for *Santolina* native to France and south-eastern Spain. Results of the Random Forest applied at the alternative grouping hypothesis inferred by the molecular phylogenetic analysis. Values are percentages. Ben-Eri = the typical population of *S. benthamiana* + the three populations of *S. ericoides*; Ben-rou = *S. benthamiana* from Le Roumenga; Dec = the three populations of *S. decumbens*.

	Ben-Eri	Ben-rou	Dec
Ben-Eri	99.3	0.7	0
Ben-rou	10.3	68.3	21.4
Dec	0.5	1.7	97.8

Table 8. Morphometric analysis for *Santolina* native to France and south-eastern Spain. Results of the Random Forest applied at the taxonomic hypothesis inferred by the results of the PCoA (Figure 3). Values are percentages. Ben-LC = *S. benthamiana* from type locality, Ben-rou = *S. benthamiana* from Le Roumenga, Dec-LC = *S. decumbens* from type locality, Dec-lfo = *S. decumbens* from La Fare-les-Oliviers, Dec-sis = *S. decumbens* from Sisteron, Eri = the three populations of *S. ericoides*.

	Ben-LC	Ben-rou	Dec-LC	Dec-lfo	Dec-sis	Eri
Ben-LC	95.6	2.1	0	0	0	2.4
Ben-rou	3.5	76.5	5.9	0	14.1	0
Dec-LC	0	0.4	96.6	1.6	1.4	0
Dec-lfo	0	0	0	99.2	0.8	0
Dec-sis	0	7.5	5.9	0.2	86.1	0.3
Eri	0	0	0	0	0	100

Table 9. Mean \pm standard deviation values of the quantitative morphological characters for each *Santolina* population. Character codes follow Table 2. Ben-LC = *S. benthamiana* from type locality, Ben-rou = *S. benthamiana* from Le Roumenga, Dec-LC = *S. decumbens* from type locality, Dec-lfo = *S. decumbens* from La Fare-les-Oliviers, Dec-sis = *S. decumbens* from Sisteron, Eri-LC = *S. ericoides* from type locality, Eri-fsc = *S. ericoides* from Sant Feliu de Codines, Eri-tor = *S. ericoides* from Torà.

Character	Ben-LC	Ben-rou	Dec-LC	Dec-lfo	Dec-sis	Eri-LC	Eri-sfc	Eri-tor
fs_len (cm)	22.9 \pm 9.2	21.7 \pm 6.3	10.8 \pm 2.6	8.6 \pm 2.9	18.1 \pm 4.4	18.6 \pm 6.7	18.6 \pm 4.1	20.8 \pm 5.1
dist_cap_lf (mm)	44.2 \pm 26.7	35 \pm 12.1	15.3 \pm 7.4	10.6 \pm 5.4	23.7 \pm 11.8	20.4 \pm 10.4	29.8 \pm 15.7	33.4 \pm 16.5
ss_len (cm)	11.8 \pm 5.2	16.5 \pm 6.4	6.3 \pm 2.5	7.7 \pm 2.6	10.1 \pm 3.0	13.7 \pm 6.8	15.3 \pm 4.2	17.9 \pm 7.5
cap_diam (mm)	6.6 \pm 1.3	6.8 \pm 1.1	6.9 \pm 0.7	6.4 \pm 1.0	7.0 \pm 0.9	6.2 \pm 1.3	7.2 \pm 0.8	7.0 \pm 0.9
sq_ext_len (mm)	2.8 \pm 0.6	3.1 \pm 0.5	2.8 \pm 0.4	2.7 \pm 0.4	3.2 \pm 0.4	2.8 \pm 0.3	2.8 \pm 0.4	3.0 \pm 0.4
sq_ext_wid (mm)	1.1 \pm 0.2	1.1 \pm 0.2	1.1 \pm 0.2	1.1 \pm 0.2	1.2 \pm 0.2	1.2 \pm 0.2	1.0 \pm 0.2	1.0 \pm 0.2
sq_int_len (mm)	2.8 \pm 0.5	3.1 \pm 0.4	3.1 \pm 0.4	2.9 \pm 0.3	3.4 \pm 0.4	3.0 \pm 0.4	3.2 \pm 0.3	3.1 \pm 0.4
sq_int_wid (mm)	1.2 \pm 0.2	1.2 \pm 0.2	1.2 \pm 0.2	1.1 \pm 0.2	1.3 \pm 0.2	1.3 \pm 0.2	1.3 \pm 0.2	1.3 \pm 0.1

sq_if_len (mm)	3.0 ± 0.3	3.0 ± 0.4	3.0 ± 0.3	2.8 ± 0.3	3.1 ± 0.2	2.9 ± 0.3	3.0 ± 0.2	2.7 ± 0.2
sq_if_wid (mm)	1.1 ± 0.2	1.0 ± 0.2	1.0 ± 0.2	1.0 ± 0.2	1.1 ± 0.2	1.1 ± 0.2	1.2 ± 0.2	1.1 ± 0.2
fl_len (mm)	3.4 ± 0.4	3.2 ± 0.4	3.0 ± 0.4	3.5 ± 0.5	3.3 ± 0.5	3.4 ± 0.6	3.5 ± 0.6	3.7 ± 0.4
fl_tooth_len (mm)	0.8 ± 0.1	0.7 ± 0.1	0.7 ± 0.1	0.7 ± 0.1	0.7 ± 0.1	0.5 ± 0.1	0.6 ± 0.1	0.6 ± 0.1
ssl_len (mm)	26.8 ± 9.9	21.3 ± 5.9	14.6 ± 3.8	11.4 ± 2.2	23.4 ± 4.1	11 ± 3.1	11.4 ± 2.1	10.6 ± 3.5
ssl_pet_len (mm)	5.7 ± 2.1	4.0 ± 2.3	2.9 ± 1.2	1.3 ± 0.7	3.7 ± 1.8	1.0 ± 0.8	1.6 ± 0.8	0.8 ± 0.4
ssl_seg_len (mm)	4.2 ± 1.5	2.9 ± 1.0	0.9 ± 0.2	0.8 ± 0.2	2.0 ± 0.4	0.9 ± 0.4	1.5 ± 0.4	1.4 ± 0.5
ssl_seg_dist (mm)	1.6 ± 0.7	1.1 ± 0.3	0.3 ± 0.3	0.3 ± 0.2	0.6 ± 0.3	0.5 ± 0.2	0.5 ± 0.3	0.6 ± 0.2
fsl_len (mm)	21.6 ± 8.9	18.9 ± 5.8	12.3 ± 3.2	8.0 ± 1.9	15.4 ± 3.1	10.7 ± 3.8	12.1 ± 2.2	10.3 ± 3.2
fsl_pet_len (mm)	6.4 ± 3.3	3.4 ± 2.4	2.1 ± 0.9	1.1 ± 0.6	2.3 ± 1.2	0.9 ± 0.6	1.8 ± 0.8	0.8 ± 0.5
fsl_seg_len (mm)	2.6 ± 0.9	2.3 ± 0.7	0.8 ± 0.3	0.6 ± 0.1	1.5 ± 0.4	0.9 ± 0.3	1.3 ± 0.4	1.4 ± 0.2
fsl_seg_dist (mm)	1.6 ± 0.9	1.0 ± 0.3	0.3 ± 0.3	0.3 ± 0.1	0.6 ± 0.2	0.6 ± 0.3	0.6 ± 0.2	0.6 ± 0.2
fs_n_nodes	21.0 ± 3.3	22.0 ± 4.3	19.8 ± 3.5	17.4 ± 4	19.4 ± 3.6	24.9 ± 6.6	26.2 ± 3.7	22.4 ± 4
ss_n_nodes	20.7 ± 4.7	23 ± 6.5	14.8 ± 3.4	20.0 ± 4.1	16.8 ± 2.7	24.6 ± 7.7	28.0 ± 5.9	26.8 ± 7.3
ssl_n_seg	29.7 ± 8.8	41.5 ± 8.9	60.5 ± 12.4	61.8 ± 13.7	60.7 ± 13.3	46.7 ± 10.6	35.4 ± 6.6	33.2 ± 4.3
fsl_n_seg	19.4 ± 5.4	31.8 ± 8	40.4 ± 9.4	38.8 ± 9.1	39 ± 9.7	40.5 ± 9.2	31.9 ± 8.3	31.6 ± 6.2
ssl_hair (%)	47.0 ± 22.0	76.1 ± 7.1	90.8 ± 7.5	61.4 ± 14.3	79.4 ± 9.9	23.8 ± 14.5	22.9 ± 14.0	13.6 ± 18.8
fsl_hair (%)	19.6 ± 18.5	67.3 ± 15.1	90.0 ± 6.2	29.1 ± 19.8	44.0 ± 18.8	15.2 ± 11.7	18.1 ± 10.9	11.3 ± 11.5
fs_hair (%)	15.2 ± 8.2	60.9 ± 14.8	90.3 ± 9.2	60.2 ± 14.0	62.9 ± 16.3	31.8 ± 12.2	31.8 ± 18.9	36.0 ± 21.2

Table 10. Pairwise comparisons of the morphological characters for which we found statistically significant differences between population pairs in *Santolina* native to France and south-eastern Spain. In the lower triangle of the table, the characters that are significantly different with Cohen's $d > 1.2$ are reported; in the upper triangle, the number of these characters is reported. In the following list, in the brackets, the number of times that a character occurs in the table is reported. "a" = fs_len (12), "b" = br_ratio (0), "c" = dist_cap_lf (10), "d" = ss_len (12), "e" = cap_diam (0), "f" = sq_ext_len (2), "g" = sq_ext_wid (1), "h" = sq_int_len (4), "i" = sq_int_wid (0), "j" = sq_if_len (3), "k" = sq_if_wid (0), "l" = fl_len (2), "m" = fl_tooth_len (8), "n" = ssl_len (15), "o" = ssl_pet_len (18), "p" = ssl_seg_len (19), "q" = ssl_seg_dist (12), "r" = fsl_len (16), "s" = fsl_pet_len (15), "t" = fsl_seg_len (22), "u" = fsl_seg_dist (15), "v" = fs_n_br (1), "w" = fs_n_nodes (5), "x" = ss_n_nodes (12), "y" = ssl_n_seg (17), "z" = fsl_n_seg (7), "A" = ssl_hair (24), "B" = fsl_hair (17), "C" = fs_hair (22). Populations acronyms are: Ben-LC = *S. benthamiana* from type locality, Ben-rou = *S. benthamiana* from Le Roumenga, Dec-LC = *S. decumbens* from type locality, Dec-lfo = *S. decumbens* from La Fare-les-Oliviers, Dec-sis = *S. decumbens* from Sisteron, Eri-LC = *S. ericoides* from type locality, Eri-sfc = *S. ericoides* from Sant Feliu de Codines, Eri-tor = *S. ericoides* from Torà.

	Ben-LC	Ben-rou	Dec-LC	Dec-lfo	Dec-sis	Eri - LC	Eri - sfc	Eri - tor
Be n- LC	-	5	17	13	11	13	13	13
Be n- rou	y,z,A,B,C	-	14	14	6	13	10	12
De c- LC	a,c,d,n,o,p,q,r, s,t,u,x,y,z,A,B ,C	a,c,d,n,p,q,r ,t,u,x,y,A,B ,C	-	7	8	9	11	13
De c- lfo	a,c,n,o,p,q,r,s, t,u,y,z,C	a,c,d,n,o,p, q,r,s,t,u,y,A ,B	o,r,s,x,A,B ,C	-	10	8	13	9
De c- sis	f,h,o,p,q,s,t,u, x,y,z,A,B,C	d,q,t,u,y,B	a,d,n,p,t,A ,B,C	a,c,h,n,o,p, r,t,u,A	-	12	11	11
Eri - LC	c,m,n,o,p,q,r,s ,t,u,y,z,A,C	c,m,n,o,p,q, r,s,t,A,B,C	a,d,m,o,s, x,A,B,C	a,m,t,u,w,y ,A,C	f,m,n,o,p, r,s,t,x,A,B ,C	-	4	2

Eri -sfc	h,m,n,o,p,q,r,s ,t,u,w,x,z,A,C	n,o,p,q,r,t,u ,A,B,C	a,d,o,p,t,w ,x,y,A,B,C	a,c,d,h,p,r, t,u,w,x,y,A ,C	d,g,n,o,r, w,x,y,A,B ,C	m, p,t, y	-	2
Eri -tor	j,m,n,o,p,q,r,s ,t,u,z,A,C	l,n,o,p,q,r,s, t,u,A,B,C	a,c,d,l,o,p, s,t,x,y,A,B ,C	a,c,d,p,t,u, y,A,C	d,j,n,o,r,s, x,y,A,B,C	t,y	j,s	-

Table 11. Results of niche similarity tests for *Santolina* native to France and south-eastern Spain. Environmental spaces were obtained from the distribution of the morphotypes following the taxonomic hypothesis inferred by the results of the PCoA (Figure 3). Backgrounds were defined by applying 5 km buffer zones around the occurrence points. The asterisks indicate a significant difference: * $P \leq 0.05$; ^{ns} $P \geq 0.05$. Ben-LC = *S. benthamiana* s.str., Ben-rou = “Le Roumenga” morphotype, Dec-LC = *S. decumbens* s.str., Dec-lfo = “La Fare-les-Oliviers” morphotype, Dec-sis = “Sisteron” morphotype, Eri = *S. ericoides*.

Morphotype 1	Morphotype 2	D
Ben-LC	Ben-rou	0.379 ^{ns/ns}
Ben-LC	Dec-LC	0.000 ^{ns/ns}
Ben-LC	Dec-lfo	0.000 ^{ns/ns}
Ben-LC	Dec-sis	0.000 ^{ns/ns}
Ben-LC	Eri	0.167 ^{ns/ns}
Ben-rou	Dec-LC	0.048 ^{ns/ns}
Ben-rou	Dec-lfo	0.184 ^{*/ns}
Ben-rou	Dec-sis	0.178 ^{ns/ns}
Ben-rou	Eri	0.212 ^{ns/*}
Dec-LC	Dec-lfo	0.017 ^{ns/ns}

Dec-LC	Dec-sis	0.013 ^{ns/ns}
Dec-LC	Eri	0.204 ^{ns/*}
Dec-lfo	Dec-sis	0.000 ^{ns/ns}
Dec-lfo	Eri	0.023 ^{ns/*}
Dec-sis	Eri	0.190 ^{ns/*}

Figures

Figure 1. The three *Santolina* species currently recorded for southern France and north-eastern Spain from their type populations. *Santolina benthamiana* from Prats-de-Mollo-la Preste, 29 June 2020 (A); *S. ericoides* from Béziers, 28 June 2020 (B); *S. decumbens* from Monte Caume, 27 June 2020 (C). All photos by L. Peruzzi.

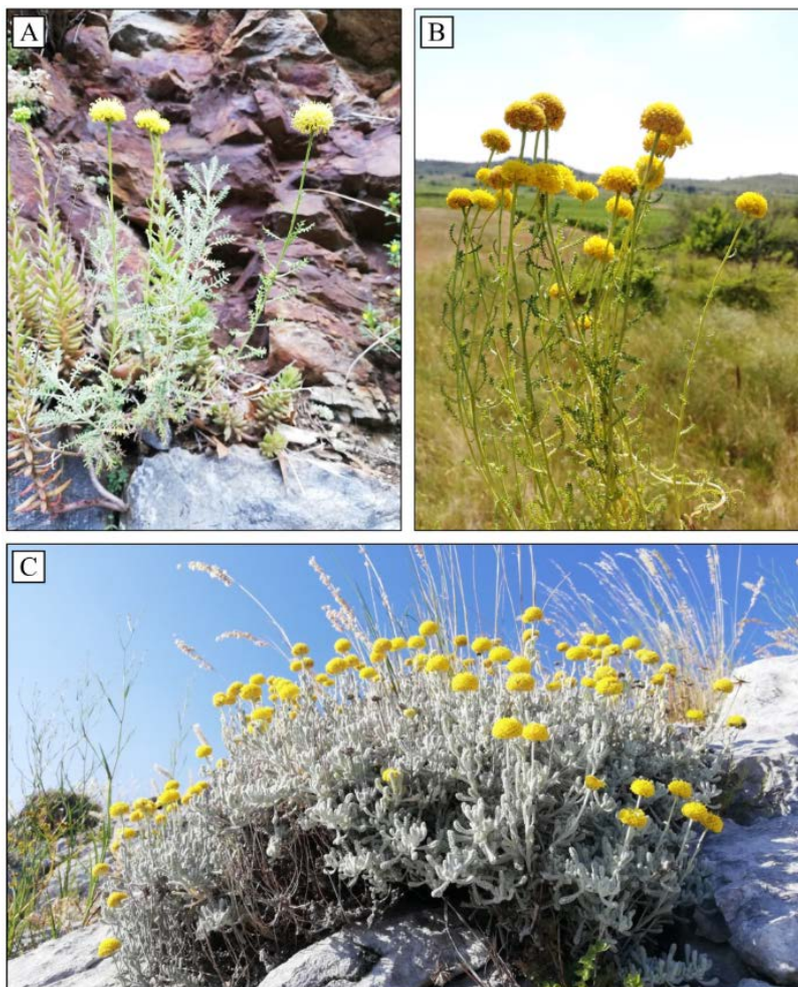


Figure 2. Bayesian consensus tree of plastid dataset for *Santolina* native to France and north-eastern Spain. LC = type locality.

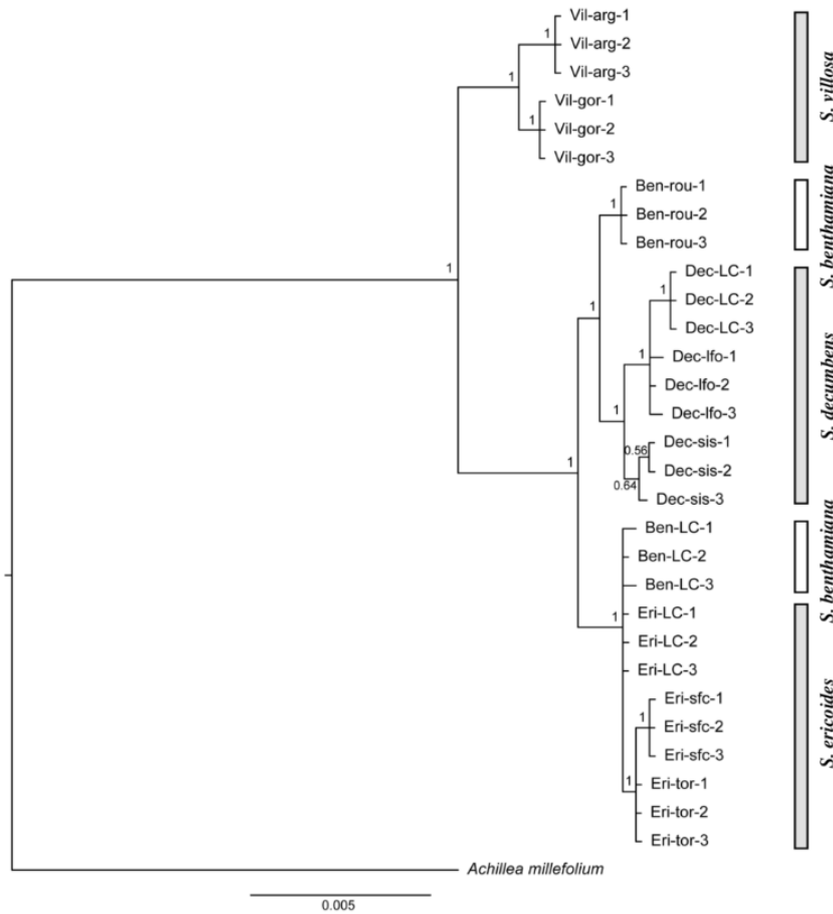


Figure 3. Morphometric analysis for *Santolina* native to France and south-eastern Spain. Scatter plot of the first two axes of the PCoA based on Gower distance. The taxonomic assignment of populations follows Giacò et al. (2021).

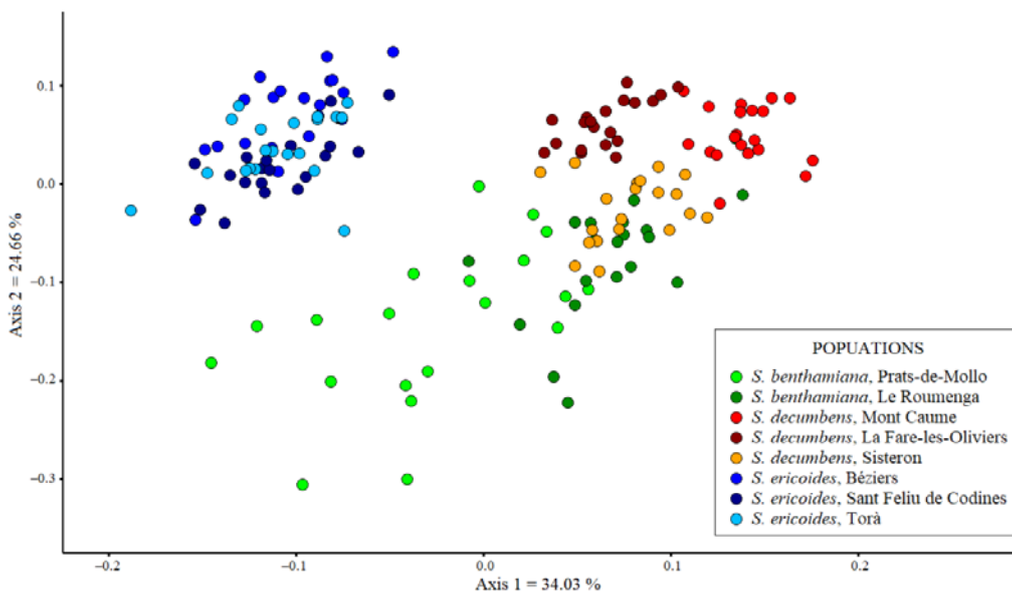


Figure 4. Distribution of the studied *Santolina* taxa native to France and south-eastern Spain, based on 163 herbarium specimens seen (Table S5). Diamonds correspond to the sampled populations: Ben-LC = Prats-de-Mollo-la-Preste, Ben-rou = Le Roumenga, Dec-LC = Mont Caume, Dec-lfo = La Fare-les-Oliviers, Dec-sis = Sisteron, Eri-LC = Béziers, Eri-sfc = Sant Feliu de Codines, Eri-tor = Torà. The taxonomic assignment of populations follows the final treatment presented in this study.

