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*Saxifraga delphinensis* Ravaud Grand Veymont, Vercors. ©A. Baumel

**Title:**

**Evolutionary distinctiveness with incomplete isolation of the narrow endemic alpine plant *Saxifraga delphinensis* Ravaud**

**Authors :**

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## Abstract

The growing evidence for the role of introgression in evolution is challenging for species delimitation, taxonomy, and conservation. Here, we examine the isolation of *Saxifraga delphinensis*, a south-west alpine narrow endemic, from its sympatric congeners *Saxifraga exarata* and *Saxifraga moschata*. Our analyses, based on morphological, molecular and genome-wide data, were extended to taxa from Jura, Massif Central, north-east Iberian mountains and Pyrenees to broaden the phylogenetic scope of the study. Phylogenetic results support the close relationship between (i) *S. delphinensis*, *Saxifraga cebennensis* and one population of *S. exarata* from Queyras and (ii) between *S. lamottei*, *S. giziana* and *S. moschata*. *Saxifraga exarata*, which was extensively sampled, is composed of several clades and is paraphyletic to other taxa. While morphological and genetic data support the evolutionary distinctiveness of *S. delphinensis*, introgression with *S. exarata* is shown by various data and analyses. In addition, ABBA-BABA analyses and plastome phylogeny reveal a possible contribution of the north-east Iberian endemic *Saxifraga vayredana* into the history of *S. delphinensis*. Overall, the results show a complex evolutionary history with frequent hybridization and spanning several mountain ranges in Europe.

## Key words

Endemism, Introgression, ITS, RADseq, Morphometry, Saxifrages, Speciation

## Introduction

Long before the development of molecular markers and omics, the terms "incipient species" (Cuénot 1917) or "semispecies" (Grant 1981) were proposed to refer to cases of incomplete isolation between taxa. Today, molecular and genomic evidence confirms that continued gene flow and recombination between species is widespread in nature (Mallet et al. 2016; Edelman & Mallet 2021). Hybridization and introgression are important for speciation and diversification (Abbot et al. 2013) as well as for adaptation (Suarez-Gonzalez et al. 2018). In plants, hybridization challenges our understanding of grouped biodiversity (i.e., species) and reproductive isolation (Rieseberg et al. 2006; Christie et al. 2022). Hybridization can also have been facilitated in climate refugia during Quaternary climate-driven shifts in species range (Nieto Feliner 2014; Voisin et al. 2023). Lastly, hybridization is clearly of conservation significance when dealing with rare or restricted endemic taxa. In practice, incomplete differentiation and introgression can lead to taxonomic problems and confusion that weaken our knowledge of the biogeography and ecology of species, which also undermines biodiversity conservation strategies.

In the iconic genus *Saxifraga*, Ebersbach et al. (2020) reported that more than 40 papers or book chapters were devoted to hybridization and estimated that 84 hybrid taxa have been described. They also showed that the rate of hybridization varied among mountain regions and within the genus and that several reproductive barriers counteract gene flow. The clade corresponding to the section *Saxifraga* in Tkach et al. (2015), has a high rate of hybridization causing incongruities between morphological and molecular data (Vargas 2000) or between nuclear ribosomal and plastid data (Mas De Xaxars et al. 2015; Carnicero et al. 2023). In this clade, *Saxifraga exarata* Villars, widespread in Alps, and *Saxifraga moschata* Wulf., widespread in Pyrenes and Alps, are known to be interfertile with at least 10 other taxa (Ebersbach et al. 2020). In Pyrenees and north-east Iberian mountains, *S. moschata* hybridizes with several endemic taxa, such as *Saxifraga pubescens* Pourr, *Saxifraga vayredana* Luizet or *Saxifraga geranioides* Host (Mas De Xaxars et al. 2015; Carnicero et al. 2023) but the taxa maintain their genetic integrity (Carnicero et al. 2023). However, in the Alps this issue was never addressed to the relationships among the "Saxifrage du Dauphiné" (*Saxifraga delphinensis* Ravaud) and its widespread congeners. It is a narrow endemic taxon restricted to the calcareous massifs situated at the south-west margin of the European Alps (Fig. 1a). In the 19<sup>th</sup> century, Ravaud described it in Vercors from the Grand Veymont summit and the Grimone Pass in his "Guide du botaniste dans le Dauphiné" (Ravaud 1885), naming it *Saxifraga delphinensis*. Based on a general aspect of the plant but without morphological study, it was renamed *S. exarata* subsp. *delphinensis* Ravaud (Kerguélen) in Kerguélen (1993) and Flora Alpina (Aeschimann et al. 2004), before being finally renamed *S.*

*delphinensis* Ravaud by Garraud (2004) and this name is used in Flora Gallica (Tison and Foucault 2014). Ravaud (1885) underlined the similarity of *S. delphinensis* with two taxa endemic to the Pyrenees *Saxifraga pubescens* Pourr (which shares similar leaves) and *Saxifraga iratiana* Schullz (which shares similar flowers). Until recently, *S. delphinensis* has not been the subject of any molecular study but a recent plastome-based phylogeny (Smyčka et al. 2022) reported its closest relationship not with *S. exarata* nor *S. pubescens* but with *S. vayredana*, a north-east Iberian endemic (Catalanidic mountains). Interestingly, in Smyčka et al. (2022) *S. pubescens* and *Saxifraga moschata* Wulf are the closest taxa of *S. exarata* whereas in Tkach et al. (2015), the sister species of *S. exarata* is *Saxifraga cebennensis* Rouy & Camus, an endemic from the Cévennes range, located at the south-east margin of the Massif Central (Fig. 1a). Similarly, various geographical and ecological isolates have been separated over time from *S. moschata*, such as *Saxifraga lamottei* Luizet (Luizet 1913), a mid-altitude isolate on basaltic siliceous rocks restricted to the Massif Central (Fig. 1a), and *Saxifraga giziana* Mouly & Ferrez (Bouchard 1947; Mouly & Ferrez 2020), a low altitude isolate on limestone from the southern Jura Massif (Fig. 1a). This latter region also harbors low altitude isolates of *Saxifraga rosacea* Moench. and *S. moschata*.

In this study, we obtained morphological, molecular and genome-wide sequencing data to examine the distinctiveness and reproductive isolation between *S. delphinensis*, *S. exarata* and *S. moschata* in south-west Alps where they often occur in sympatry. Acknowledging previous studies (Vargas 2000; Mas De Xaxars et al. 2015; Smyčka et al. 2022) that established phylogenetic links between taxa of different mountain ranges in the section *Saxifraga*, we broaden the scope of the study by including several endemic taxa to contribute to the knowledge of their phylogenetic relationships. We therefore include *S. giziana* from Jura; *S. lamottei* and *S. cebennensis* from Massif Central, *Saxifraga pubescens* and *S. geraniodes* from Eastern Pyrenees; *Saxifraga vayredana* and *Saxifraga genesiana* Vargas from the Catalan Pre-Coastal Range; *Saxifraga moncayensis* Webb from the Moncayo Range and *Saxifraga hariatii* Luizet & Soulié from Western Pyrenees.

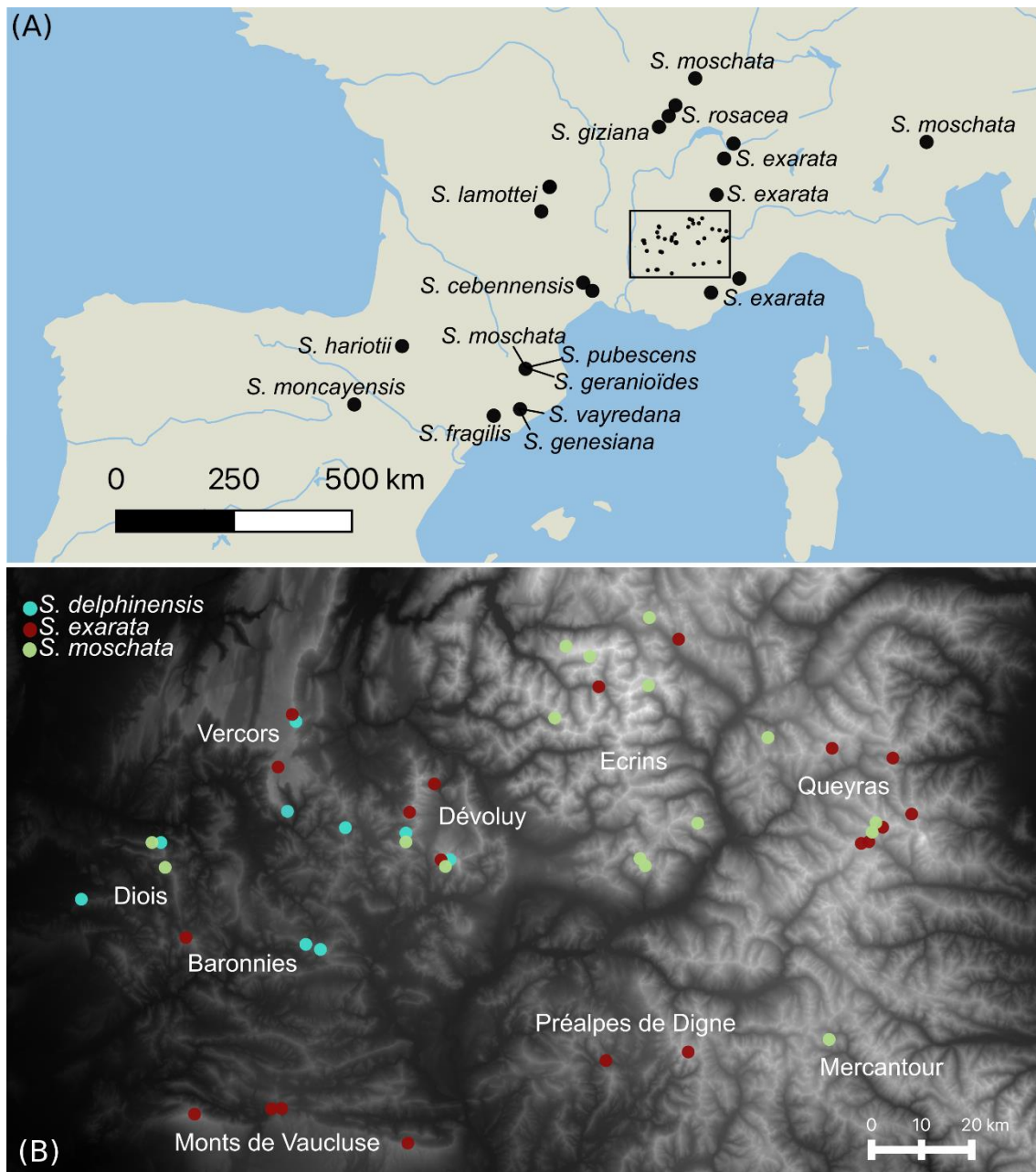
## **Material and Methods**

### **1. Sampling and general strategy**

The sampling strategy was focused on *S. delphinensis* and *S. exarata* and *S. moschata* in the range of *S. delphinensis* in southwestern PreAlps. Apart from this area, *Saxifraga exarata* was more intensively sampled in eastern Queyras (France), because several forms were described by Villars (1779) in the higher parts of Queyras (in high mountains, above the village of Ristolas). In the

locality of Pic de Bure (Devoluy), we sampled more intensively *S. delphinensis* and *S. exarata* because both taxa are abundant on this plateau, sharing a similar ecology and occurring in close sympatry. The three narrow endemics, *S. lamottei*, endemic from Auvergne, *S. cebennensis*, endemic from Cevennes, and *S. giziana* endemic from Jura, were less sampled but some material was obtained from several populations whenever possible. In Jura, samples from *S. moschata* and *S. rosacea* occurring in the same area as *S. giziana* were also included. Two populations of *S. vayredana* were sampled from north-east Iberian mountains (Catalanidic mountains, Catalan Pre-Coastal Range) and two populations of *S. pubescens* in Eastern Pyrenees. Only one locality was sampled for the other taxa, namely *S. geraniodes*, *S. genesiana*, *S. moncayensis* and *S. hariotii*. The geographical coordinates of sampling locations are listed in Sup. Mat. Table 1. A map of the 57 localities sampled is shown in Figure 1.

We generated three data sets: morphometric measurements, rDNA internal transcribed spacer (ITS) sequences and genome-wide genotypes by double digest restriction site associated DNA sequencing (RADseq). The accessions used to generate molecular data are listed in Sup. Mat. Table 1. Morphometric measurements were made on 9 taxa and a subset of localities, and all the data are summarized in Sup. Mat. Table 2. In total, the morphological data, the ITS data and the RADseq data are based on 242, 105 and 168 individuals respectively. Sixty-four specimens, representing various sampled populations, were deposited in the GAP herbarium (voucher list in Sup. Mat. Table 1). ITS haplotypes sequences were deposited in Genbank (accession list in Sup. Mat. Table 5) and the raw RADseq reads were deposited at NCBI SRA database under Bioproject accession PRJNA935954.



**Figure 1:** Map of the 57 sampling sites (A) with a focus on south-west Alps (B) where *S. delphinensis*, *S. exarata*, and *S. moschata* were more intensively sampled.

### DNA extraction

Total DNA was extracted from leaves stored with silica-gel using NucleoSpin Plant II kit (Macherey Nagel) following the kit instructions with slight modifications. From 40 to 50 mg of dry leaves were grinded (Qiagen Tissu Lyser), centrifuged and resuspended into 700  $\mu$ l PL1 + 10 of RNase A for 45 minutes of lysis at 65°C. After filtration, the lysate was collected taking care not to mix the solution and the last 100  $\mu$ l, at the bottom of the tube, were abandoned. This modification made it possible to remove contaminants that caused abnormal 260/230 ratio values, which were measured to assess DNA purity. After this modification, we followed the manufacturer's protocol



and elution of DNA was done with 90 µl of PE solution. DNA concentrations were measured with a Nanodrop, and concentration results were trusted only if 260/230 ratio was greater than 1. Only DNA extracts with at least 20 ng/µl of DNA in 80µl were used for RADseq.

### **ITS sequencing and analysis**

The ITS region (ITS1+5.8S+ITS2) was amplified using the ITSsp5 and ITSsp4 primers of Chen et al. 2016. PCR was performed in a 25 µL reaction volume containing 5 µL 5X buffer (kit GoTaq G2 Flexi DNA polymerase, Promega), 1.5 mM MgCl<sub>2</sub>, 1.25 mM each PCR Nucleotide (Promega, U1420), 0.3 µM each primer, 0.2 units Taq polymerase (GoTaq G2 Flexi DNA Polymerase, M7805, Promega), and 10 ng DNA. Amplifications were carried out in an Eppendorf thermocycler. The following thermal profile was used: initial denaturation at 94 °C for 5 min, followed by 30 cycles of 45 s at 94 °C, 45 s at 55 °C and 1 min at 72 °C, and finishing with 8 min extension at 72 °C. Sanger sequencing was done by Eurofins Genomics; sequences were examined and aligned in Mega-X (Kumar et al. 2018). When the chromatograms revealed polymorphic peaks we tried to identify the ribotypes or we replaced the ambiguous site by the appropriate IUPAC code. Haplotypes were identified with Fabox (Villesen 2007) and a network was built using the haploNet function of R pegas package (Paradis 2010).

### **Plastome phylogeny**

To include plastid data in our study, we used the plastome phylogeny of Smyčka et al. (2022). Briefly, complete circular plastomes from several species of *Saxifraga* sect. *Saxifraga* were retrieved using a genome skimming approach, then dated phylogenies were done with BEAST 2 (Bouckaert et al., 2014) with fossil calibrations in accordance with Ebersbach et al. (2017). We used only the clade including the taxa studied here.

### **RADseq, assembly optimization , data filtering**

Genomic library preparation and sequencing were conducted by Microsynth ecogenics GmbH (Blagach, Switzerland). DNA samples (with a minimum of 1 µg input and on average 3.6 µg of DNA) were digested with the restriction enzymes EcoRI/MseI following heat inactivation according to the manufacturer's protocol (New England Biolabs, NEB). Fragments between 500 and 600 bp were selected by automated gel cutting (Pippin Prep system, sage science<sup>®</sup>), Illumina Y-shaped adaptors were ligated, and ligation products were bead purified. Each library was then individually barcoded by PCR using a dual-indexing strategy. Individually barcoded libraries were pooled and subsequently purified before sequencing. Three runs of paired-end sequencing 2x75 bp on an Illumina NextSeq platform were performed on 60, 60 and 48 samples respectively

leading to an average of 4.2 million raw reads per sample. Demultiplexing and trimming of Illumina adaptor residuals was done by EcoGenics.

Ipyrad was used for reads trimming and assembly (Eaton and Overcast 2020). A pilot study was done on 18 samples to optimize the level of sequence similarity for the clustering threshold (steps 3 and 6 of ipyrad); the assembly was repeated for 7 values of this parameter from 84 to 96%. The beginning and the end of each read were trimmed (5 bp), the minimum length of reads was set on 35 bp and the quality filters were left at the default values. The minimal depth for base calls was set to 6. To limit the presence of repetitive loci we set the maximum depth, above which cluster were removed, to 1000. The parameters helping to limit paralogs were set to default values: maximum number of SNPs in a final locus (20%), the maximum number of indel by locus (4) and the maximum proportion of shared polymorphic sites in a locus (50%). The other parameters were also set to default values. After assembly, to estimate the efficiency of the assembly we first used the following ipyrad summary statistics: the number of RAD loci filtered by the maximum % of shared heterozygous sites in a locus, the sequencing error rate, the number of RAD loci, and the sum of parsimony informative sites. Then we used a custom R script to estimate the inertia of the two first axes of a principal coordinates analysis to roughly estimate the information of the dataset. According to this pilot analysis (Sup. Mat. Table 4) we found that cluster thresholds of 92% or 94% were optimal. Because the number of parsimony informative sites was slightly higher, we proceed to the assembly of the whole data set with a 92% clustering threshold. To account for the impact of the minimum number of individuals by loci (Min\_ind) we produce two assemblies by setting this parameter to 24 (Min\_24) and 48 (Min\_48). Below these values the file sizes were difficult to handle and above these values a large amount of data was lost. We used ITS sequence as a filter to remove reads that could correspond to this sequence. The other parameters of ipyrad were set to be the same as those used during the pilot study.

#### **IQ-TREE maximum-likelihood phylogeny of 166 RADseq genotypes**

The concatenated matrix of RADseq loci was used to infer a phylogenetic tree of all samples by maximum likelihood, using the software IQ-TREE (Nguyen et al. 2015) for a maximum likelihood phylogenetic analysis. The best model of molecular evolution was determined by ModelFinder implemented in IQ-TREE (Kalyaanamoorthy et al. 2017). Branch support was estimated using the ultrafast bootstrap method (1000 iterations; Thi Hoang et al. 2017). Default parameters were applied. This analysis was performed on Min\_24 and Min\_48 data-sets and the two phylogenetic trees were compared with the comparePhylo function of ape R packages (Paradis & Schliep 2019). Tree were drawn, rooted and edited with FigTree (Rambaut 2009).

### **Genetic admixture analyses**

Evidence for introgression were searched by estimating a co-ancestry matrix among all individuals performing RADpainter (Malinsky et al. 2018a) on the Min\_24 datasets. RADpainter, is based on haplotype sequence differences and therefore uses all of the data. Because RAD loci were not mapped on a reference genome, we used the LD script, provided with RADpainter, to reduce, as recommended, the effect of linkage disequilibrium. After this step, we used RADpainter with default settings to compute a co-ancestry matrix among all individuals. The matrix was re-organized with respect to the phylogenetic tree obtained with iqTREE and then used to build a heat map displaying shared ancestry among individuals with a custom R script. Under a scenario of strict divergence, the RADpainter heat map organized in this way will show a contrasting color pattern with squares in warm colors (i.e. clusters of high co-ancestry genotypes) aligned on the diagonal and corresponding to clades. Incomplete lineage sorting will produce cool to warm color gradients around the squares, but these will still be aligned on the diagonal, respecting the phylogenetic structure, whereas introgression will result in warm squares appearing off the diagonal.

Genetic admixture was also estimated with *snmf* (R LEA package, Frichot and François, 2015) which has the merit of relaxing all assumptions of the Hardy-Weinberg structure. We used the Min\_24 data sets that were thinned and filtered with an R custom script to keep one SNP by locus, the one having the minimum allelic frequency, and only SNP with the less frequent allele observed in at least 3 individuals. *Snmf* was run for  $K = 2$  to 6, 400 repetitions, 100 iterations, regularization parameter set to 25 and 20% of the genotypes masked to compute the cross-entropy criterion. Barplots showing ancestry coefficients were obtained with the *compplot* function in *adegenet* R package (Jombart and Ahmed, 2011), with genotypes sorted according to their order in the phylogenetic tree obtained with IQ-TREE. A second genetic admixture analysis was restricted to *S. delphinensis*, *S. exarata*, and *S. moschata* with the same parameters as indicated above, but we filtered individual and SNPs to have an overall missing rate below 20% and *Snmf* was run for 3 groups.

### **SVDQuartet coalescent-based phylogeny**

Concatenation methods such as IQ-TREE may potentially be affected by incomplete lineage sorting and overconfidence in node support. This is a likely issue here because of the recent diversification of the taxa studied (Smyčka et al. 2022). To explicitly address this issue, we perform a coalescent-based phylogenetic inference. To reduce the influence of hybridization and missing data, we selected genotypes to maximize loci coverage and minimize admixture level

(see *Snmf* analysis below ). The ipyrad assembly was repeated with this new set of genotypes and was filtered with vcfTools (Danecek et al. 2011) to retain only biallelic SNPs, a maximum of two SNPs per locus, only SNPs with the minimum frequency allele present in at least 3 individuals and with a missing data rate below 50%. Then we use the vcf2phylo script (Ortiz 2019) to obtain a nexus file that was analyzed with the SVDQuartet method (Chifman and Kubatko 2014), as implemented in PAUP (Swofford 2018) to obtain a coalescent-based phylogenetic tree. The analysis was conducted at the genotype and at the species level, using all possible quartets, the multi-species coalescent and 200 replicates for the bootstrap.

### **Test of introgression**

Introgression was tested with the Dsuite software (Malinsky et al. 2021) implementing ABBA-BABA tests and related statistics. The interpretations of ABBA-BABA tests are difficult because (i) the number of possible trios (P1, P2, P3) increase with the number of populations or species analyzed, (ii) it is not always possible to use a phylogenetic tree to eliminate non-relevant tests (iii) and the D statistics can be correlated among the branches of a phylogeny thus causing possible confusion between recent and ancient gene flow. To overcome these difficulties, we performed two analyses.

The objective of the first analysis was to confirm by a test the significant prevalence of introgression in the whole data set (Min\_24). We conducted ABBA-BABA tests at the level of the RADpainter clusters of genotypes. This analysis was not guided by a phylogenetic tree and we summarize the results by analyzing the distribution of significant D statistics. We filtered the Min\_24 data set with vcfTools to keep one SNP per locus and a minimum of 12 individuals per SNP. Then we used the genotypes of four taxa that were grouped as a basal lineage in the maximum likelihood tree as the outgroup. Dtrios was conducted according to the BBAA pattern to organize the P1 and P2 taxa. Significant Dstat scores indicate a phylogenetic conflict due to an excess of ABBA pattern with P2 as receiver and P3 as donor. To avoid the risk of false positives we considered only Zcores above 3. We reported these results as violin plots of Dstat distributions (for Zscores > 3) to show overall patterns of the contribution, as donor or receiver, of each of the three main taxa sampled in the study: *S. delphinensis*, *S. exarata* and *S. moschata*.

The second analysis examined the significance of introgression. For this we re-used the SNPs dataset obtained for the SVDQuartet phylogeny in which admixed individuals were deleted. The genotypes were assigned to their species and the analysis was guided by the species-level phylogeny. We developed the f-branch method which assigns hypotheses of gene flow to internal or external branches of the phylogeny (Malinsky et al., 2018b; Malinsky et al. 2021) and

therefore facilitates the interpretation by reducing the number of possible donor or receptor lineages. The f-branch metric, derived from f4-ratios, reflects excessive sharing of alleles between the P3 species and the descendants of a branch. The graphical output assigns gene flow between the branches of the phylogeny.

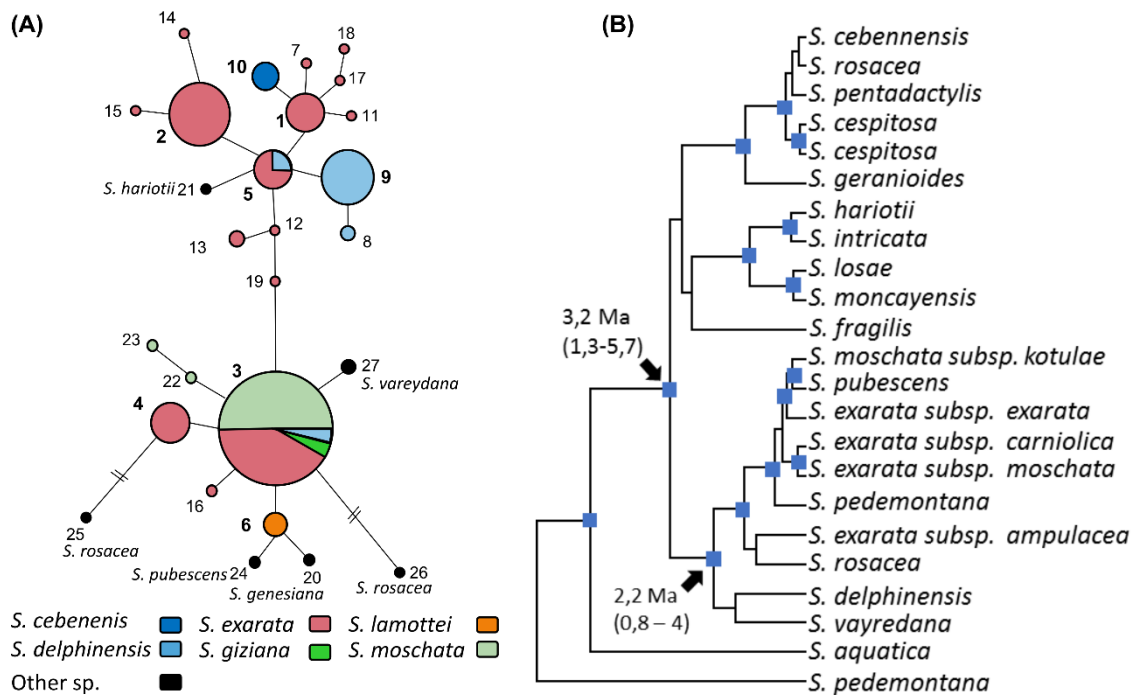
### **Morphometry**

To investigate morphological distinctiveness of study taxa, we estimated or measured vegetative characters. The cushion compactness was estimated from 1 (open and loose) to 5 (compact and domed cushion). Based on leaf pictures, hairiness was estimated from 1 (total or nearly total absence of trichomes) to 4 (dense hairiness); leaf furrows were estimated as following: 1 (absence), 2 (incomplete furrow) and 3 (complete furrow from the basis to the tip of the leaf). All characters were coded by the same person. Three leaves collected per individual were scanned in the lab in order to count the blade segments and to capture the leaf shape with positioning 12 pseudo-landmarks on each leaf (R geomorph package, Adams and Otárola-Castillo 2013) (Sup. Mat. Fig. 1a). Generalized Procrustes analysis was performed, and the leaf values were averaged to obtain an estimated leaf shape for each individual. A total of 242 individuals were scored for 10 taxa (Sup. Mat. Table 3). Multivariate analyses were conducted in two steps. First, the principal components of leaf shapes variation were retrieved by a discriminant analysis with the taxa as the discriminant factor (R ade4 package, Dray and Dufour 2007). Secondly, we performed a Hill and Smith analysis ( R ade4 package) to analyze the variation of all traits: hairiness, leaf furrow, cushion compactness, leaf shapes principal components and blade segments number.

## Results

### rDNA ITS network versus plastid phylogeny

The alignment of 105 ITS sequences (686 bp) revealed 59 variable sites and 27 haplotypes that were present together in a network (Fig. 2a). The distribution of ITS haplotypes among the samples, the taxa and their localities are shown in Sup. Mat Tables 5 and 6. *Saxifraga exarata* is present in almost all the network and has the main central haplotypes 1, 2, 3 and 5. By contrast *S. moschata* is represented by only three haplotypes, the most frequent being shared with other taxa and two singletons. The four endemics present different patterns. *S. delphinensis* and *S. cebennensis* have evolved original haplotypes but *S. delphinensis* shares haplotypes 3 and 5 with *S. exarata*, *S. moschata*, *S. giziana* (haplotype N°3) and with *S. exarata* (haplotype N°5) respectively. These haplotypes are observed in several locations (Sup. Mat Table 6). By contrast *S. giziana* and *S. lamottei* have only one haplotype each. *Saxifraga giziana* has the haplotype 3 shared with 3 other taxa (*S. exarata*, *S. moschata*, *S. delphinensis*) and *S. lamottei* has the haplotype 6 deriving from haplotype 3.



**Figure 2:** Nuclear ribosomal and plastid data report different histories for *Saxifraga*. (A) Parsimony network of ITS haplotypes, the size of the circles is proportional to the number of samples and the colors indicate the taxa. The strokes on lines indicate longer branches that were shortened. (B) Plastome-based time tree adapted from Smyčka et al. (2022). Blue squares indicate bootstrap values above 95%. In (B) taxa names are those used by Smyčka et al. (2022), *S. exarata* subsp. *moschata* corresponds to *S. moschata* here.

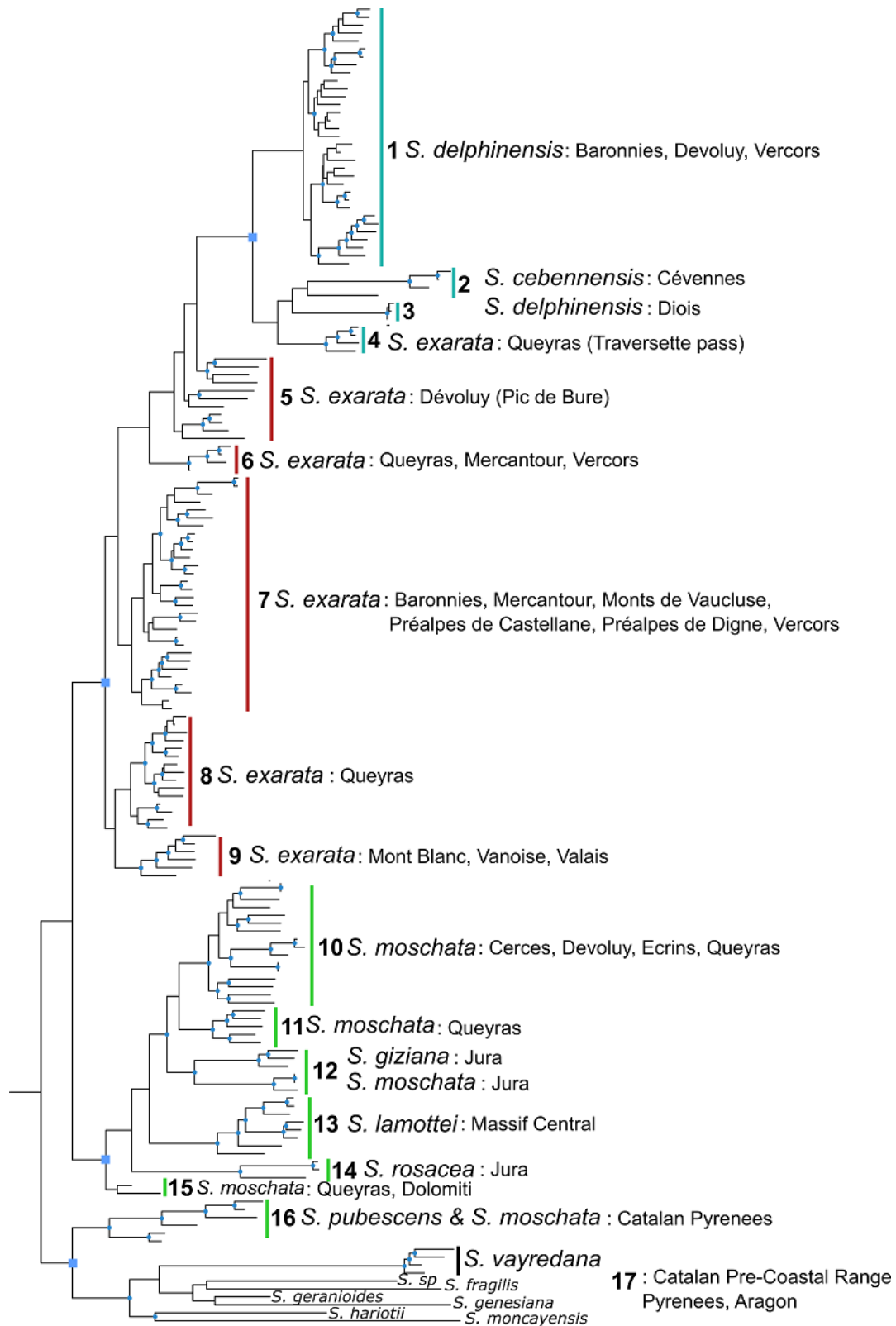
The plastome-based time tree (Fig. 2B) shows that the taxa studied here are forming two clades diverging from 3.2 Ma (1.3 – 5.7). It shows also the close relationship between *S. moschata* and *S. exarata*, but that *S. cebennensis* and *S. delphinensis* belong to different clades; *S. cebennensis* being related to *S. rosacea* and *S. pentadactylis* while *S. delphinensis* is at the base of the second clade which includes *S. exarata* and *S. moschata*. The sister species of *S. delphinensis* could be *S. vareydana* but their node is not robust (posterior 50%).

### **IQ-TREE maximum-likelihood phylogeny and admixture analyses**

After filtering, the mean number of reads was of 3.6 M reads per individual. Two samples having less than one million raw reads were removed (too few loci after the assembly) and we performed loci assemblies on 166 individuals. After the assembly, the Min\_24 data set retained 32,398 loci with 73% missing data and 58,138 parsimony informative sites. The mean number of loci per individual was of 9291 loci, the mean individual heterozygosity of 2.2%, and the mean sequencing error rate of 0,7% (Sup. Mat. Table 7). The Min\_48 had 10931 loci, with 58% missing data and 22,519 parsimony informative sites. As expected, the mean number of loci per individual was lower (5039 loci) but the mean individual heterozygosity was identical to that of the Min\_24 assembly (Sup. Mat. Table 7).

The IQ-TREE analysis performed on the Min\_24 and Min\_48 alignments revealed very similar trees (Sup. Res. Fig. S1). The main differences produced by changing the minimum number of individuals per locus concerned a few genotypes but didn't change the overall phylogenetic structure. All of the following results are thus based on the Min\_24 data set (summary stats of this assembly are given in Sup. Mat. Table 7). The IQ-TREE phylogeny of 166 samples (Fig. 3; Sup. Res. Fig. S2) reveals three main robust clades (ultrafast bootstrap values > 95%). *Saxifraga delphinensis*, *S. cebennensis* and *S. exarata* genotypes form the first clade, *S. moschata*, *S. giziana*, *S. lamottei* and *S. rosacea* the second clade and the genotypes from Aragon and Catalan Pyrenees, including *S. pubescens*, *S. vareydana* and *S. moschata*, the third clade.

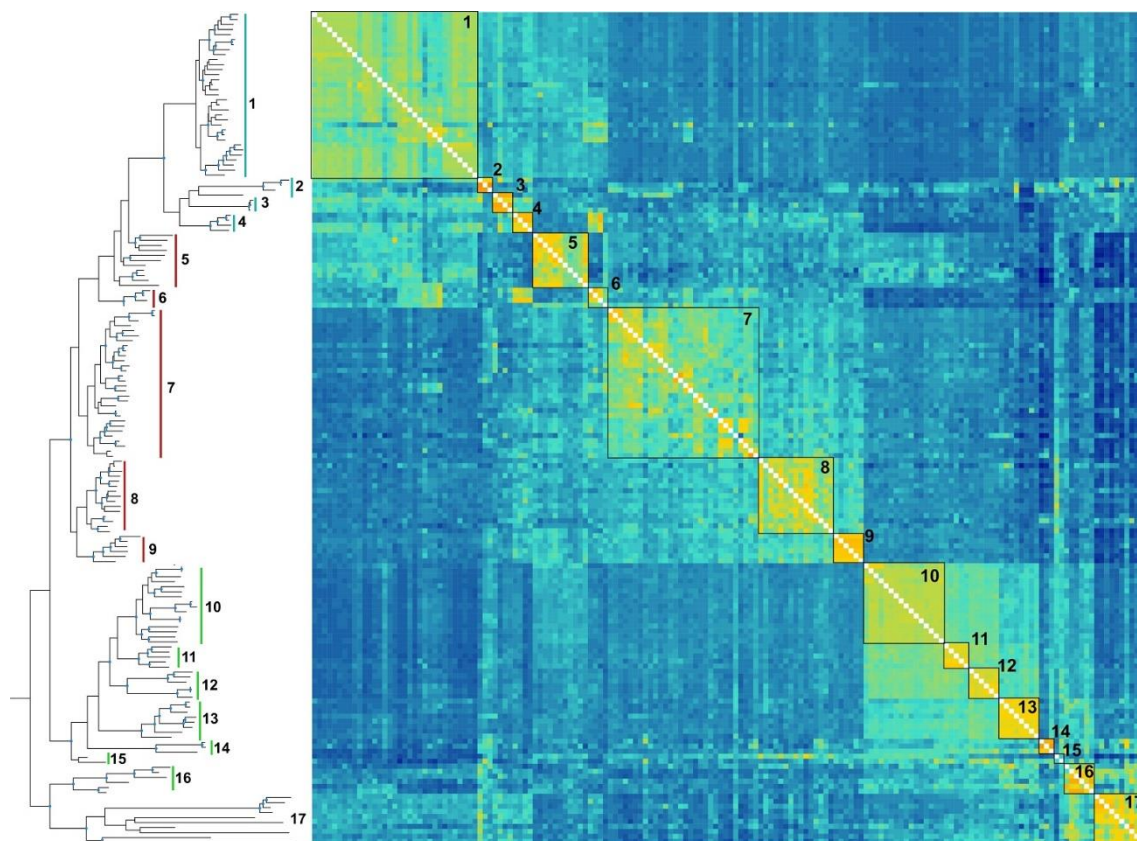
Interestingly, although *S. delphinensis* and *S. cebennensis* are not well supported, and *S. delphinensis* is not monophyletic, these two taxa form a robust clade with *S. exarata* genotypes from the Traversette Pass (Queyras) that were morphologically different from other *S. exarata* (see morphometry).



**Figure 3:** Genome-wide phylogenetic relationships among 166 *Saxifraga* individuals revealed by an IQ-TREE maximum likelihood tree based on a concatenated alignment of 264,183 bp (58,138 parsimony informative sites, 73% of missing data) and a TVM+F+I+G4 model. Blue squares are indicating ultrafast bootstrap values > 95%. Numbers correspond to groups of high co-ancestry (RADpainter; Fig. 4). The tree with each genotype ID is in Sup. Res. Fig. S2.



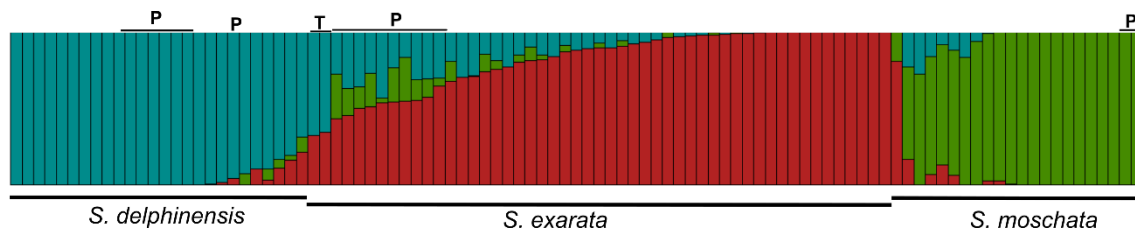
When compared, the IQ-TREE phylogeny and the RADpainter co-ancestry matrix match for 17 groups (Fig. 3 and 4). Their geographic distribution is summarized in the Figure 3. Within these groups, genotypes have high co-ancestry (warm colors along the diagonal, Fig. 4) but the overall pattern of RADpainter reveal the presence of incomplete differentiation and probable gene flow among the groups, indicating a loose phylogenetic structure. For example, there is clear gradient of decreasing co-ancestry from the group “1” to group “9” that is congruent with the paraphyletic relationships between *S. exarata*, *S. delphinensis* and *S. cebennensis*. Interestingly, the *S. exarata* genotypes from Pic de Bure site (“5”) have clear co-ancestry with *S. delphinensis* (“1”) and *S. moschata* (“10”). This pattern is already noted in the *Snmf* admixture analysis (see below). *Saxifraga moschata*, *S. lamottei* and *S. giziana* are closely related as shown by their high genetic co-ancestry and they are less isolated from *S. exarata* than from *S. delphinensis*.



**Figure 4:** RADpainter co-ancestry matrix of 166 *Saxifraga* genotypes. The heat map was ordered according to the IQ-TREE phylogeny (left and Fig. 3) obtained on the same dataset. Genetic similarity between pairs of genotypes increases from blue to orange values. The black squares with numbers were reported for comparison with the phylogeny.

As with the RADpainter heat map, the genetic admixture analysis by *Snmf* performed on the 166 genotypes (Fig. S2, 566 SNPs, 33% missing data) shows a general correspondence between the genetic and phylogenetic structures but without complete differentiation between clades or

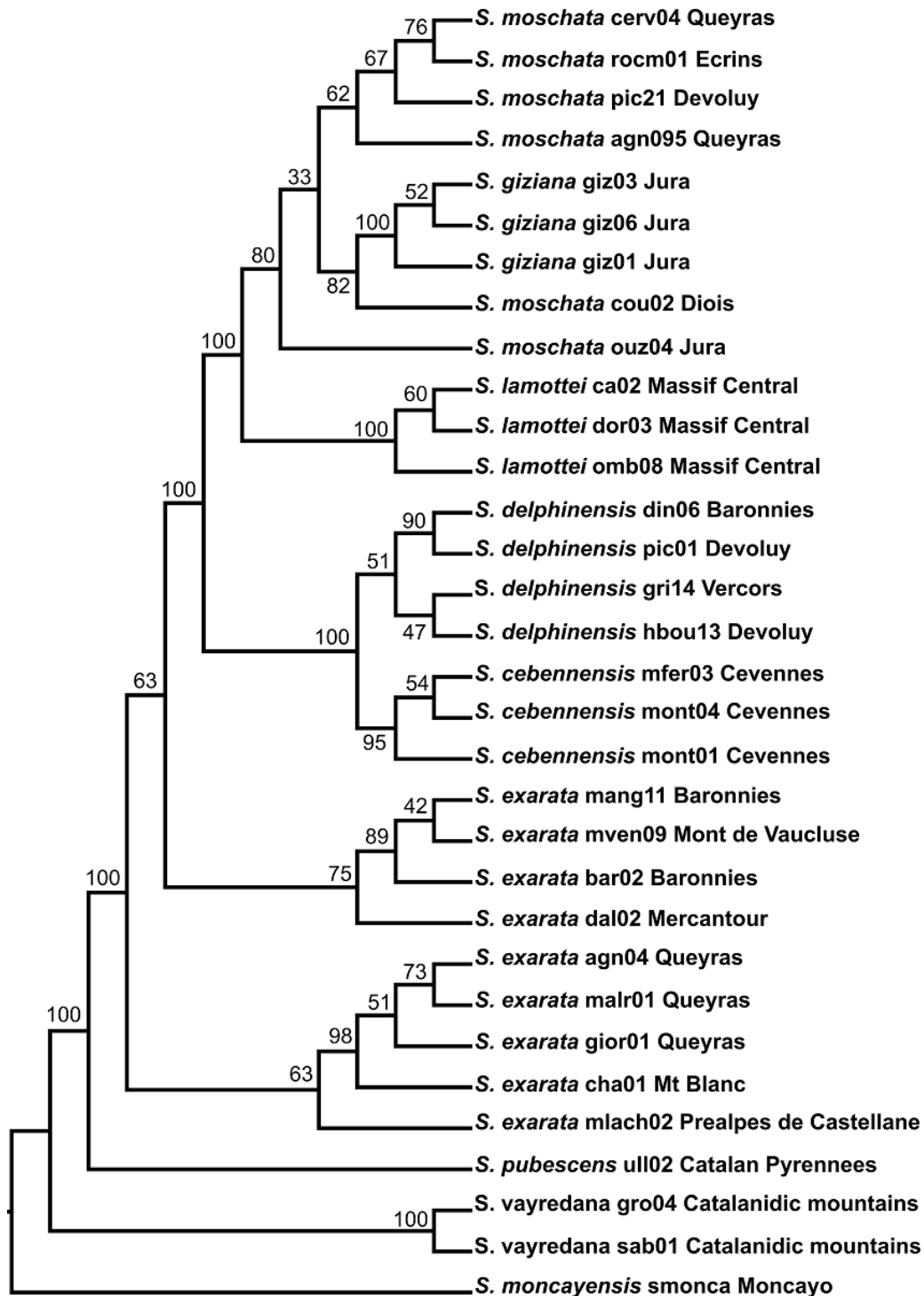
between taxa. The *Snmf* analysis (Fig. 5, 560 SNPs, 12% missing data) confirmed the clustering of the *S. delphinensis*, *S. exarata* and *S. moschata* into 3 genetic groups and the genetic admixture involving mostly *S. exarata* and *S. delphinensis* and to a lesser degree *S. moschata*. On the plateau of Pic de Bure, 10 individuals (group “5”, “P” Fig. 5) were identified in the field as *S. exarata* but they are genetically admixed with genetic *S. delphinensis* and with *S. moschata* to a lesser extent. The *S. delphinensis* and *S. moschata* from this locality were almost non-admixed. Genetic admixture is also supported in localities where the taxa do not occur in sympatry. In such cases it involves a contribution of the genetic group corresponding mainly to *S. delphinensis* sometimes distant from hundreds of kilometers away. This is very clear for *S. exarata* genotypes sampled from the Traversette Pass (“T” Fig. 5), and in other populations of *S. exarata* or *S. moschata* from the inner Alps, although to a lesser extent.



**Figure 5:** Admixture analysis (*Snmf*) performed on 99 RAD genotypes of 560 independent SNPs and 12% of missing data. Genotypes are ordered and colored according to their membership in 3 genetic clusters corresponding to *S. delphinensis*, *S. exarata* and *S. moschata* respectively. “P” and “T” indicate *S. exarata* genotypes from Pic de Bure and Traversette Pass respectively.

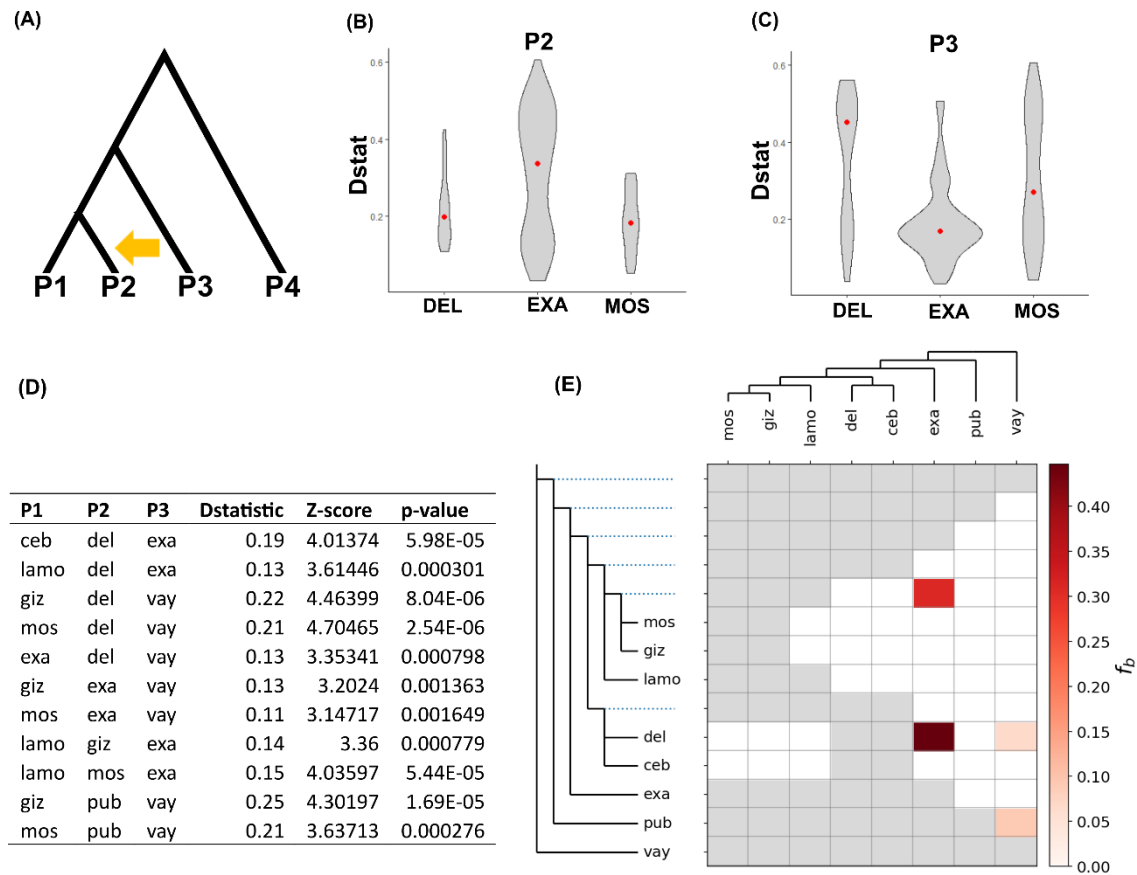
### SVDQuartet coalescent-based phylogeny

This analysis was designed to reduce the influence of incomplete lineage sorting and genetic admixture in the phylogenetic inference. It was performed on 7625 biallelic SNPs with the minimum frequency allele present in at least 3 individuals. The SVDQuartet phylogeny, supported by 76% of the quartets (Fig. 6), revealed a different phylogenetic structure than the IQ-TREE phylogeny. Indeed, a robust clade unites *S. delphinensis* and *S. cebennensis* in one clade and *S. moschata*, *S. lamottei* and *S. giziana* in a second. This clade derives from *S. exarata* which is again paraphyletic.



**Figure 6:** Coalescent-based phylogeny by SVDQuartet method of 32 *Saxifraga* genotypes. Exhaustive quartet sampling was conducted in a matrix of 7625 biallelic SNPs. The bootstrap values near the nodes were obtained with 200 replicates. Branch lengths are not meaningful.

## Tests of introgression

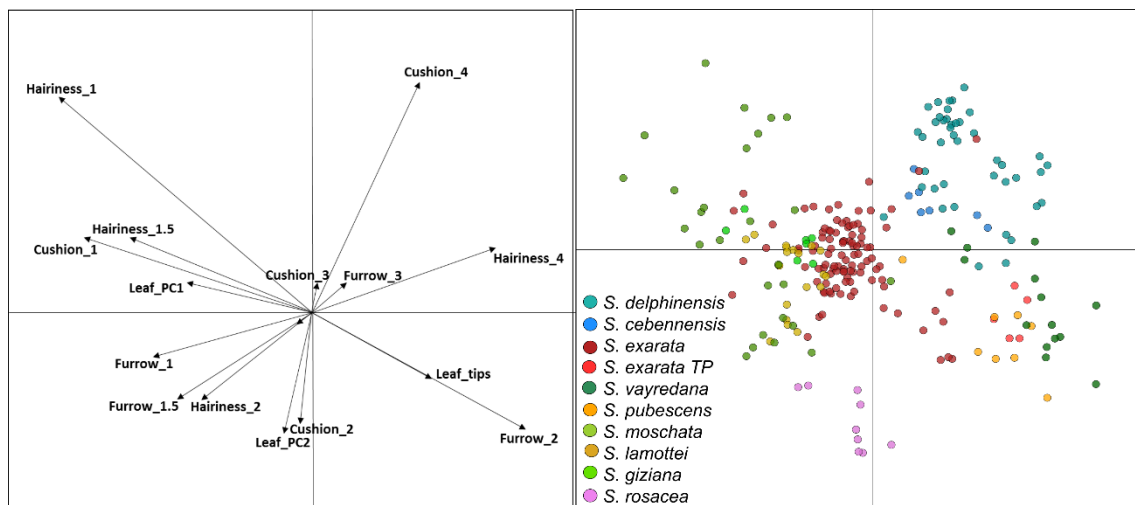


**Figure 7:** Results of Dsuite introgression tests. (A) to (C) ABBA-BABA tests performed on 166 genotypes assigned to 17 groups (Fig. 4), only Dstat with Zscores > 3 were kept. (A) Positive Dstat scores indicate an excess of ABBA pattern and an introgression from P3 to P2. (B) and (C) Violin plots of the Dstat distribution according to P2 or P3 for the three main species sampled here, *S. delphinensis* (DEL), *S. exarata* (EXA) and *S. moschata* (MOS). (D) to (E) Dsuite analyses guided by a coalescent-based species phylogenetic tree and performed on individuals with low genetic admixture. (D) Results of ABBA-BABA tests with z-scores > 3. (E) Results of f-branch analysis. The phylogenetic tree is shown on x and y axes, colored squares indicate significant excess of SNPs sharing caused by gene flow. White or grey squares represent non-significant values or trios incompatible with the phylogeny respectively.

ABBA-BABA tests, performed on the 166 genotypes dataset assigned to the 17 groups (Fig. 4) and without phylogenetic guide tree (Sup. Mat. Table 8; Fig. 7B,C), revealed 313 out of 680 trios with Zcores greater than 3 (all pval \*\*\*, Sup. Mat. Table 8) with a Dstat ranging from 3% to 60% and a median of 22%, supporting a high excess of ABBA pattern in the data set. Such result indicates that introgression, rather than incomplete lineage sorting, is at the source of the incomplete differentiation and phylogenetic conflicts revealed previously. The median Dstat is higher when *S. exarata* is the receiver (P2, 30%, Fig. 7B) and lower when it is the donor (P3, 19%, Fig. 7C). The second analysis (Fig. 7 D,E) was performed on the 32 genotypes used for the SVDQuartet phylogeny. The genotypes were assigned to their taxa and the analysis was guided

by the SVDQuartet phylogeny. The ABBA-BABA tests revealed 11 out of 56 trios with Zscores greater than 3 (all pval \*\*, Fig. 7D, Sup. Mat. Table 9) with Dstat ranging from 11% to 25%, thus indicating significant excess of ABBA pattern. The detected introgressions involve (i) *S. delphinensis* as a receiver (P2) with *S. exarata* or *S. vayredana* as a donor (P3), (ii) *S. exarata* as a receiver with *S. vayredana* as a donor, (iii) *S. moschata* and *S. giziana* as receivers with *S. exarata* as a donor, and (iv) *S. pubescens* as a receiver and *S. vayredana* as a donor. The f-branch metric (Fig. 7E) confirmed gene flow between *S. exarata* and *S. delphinensis*, *S. vayredana* and *S. delphinensis* and between *S. exarata* and the ancestor of *S. moschata* and *S. giziana*.

### Morphological differences with overlap



**Figure 8:** Multivariate Hill & Smith analysis of vegetative polymorphism of *Saxifraga* based on principal components of leaf shapes (Leaf\_PC), number of leaf blade segments (Leaf\_tips), absence of leaf furrow (Furrow 1), incomplete leaf furrow (Furrow 1.5), complete leaf furrow (Furrow 2), multiple leaf furrow (Furrow3), leaf hairiness from nearly absent (Hairiness 1) to very dense (Hairiness 2) and cushion form from loose and open (Cushion 1) to compact and domed cushion (Cushion 4).

The ten taxa show largely overlapping vegetative phenotypes (Fig. 8). The main axis is opposing, left to right, phenotypes with absent or incomplete furrow to phenotypes with complete furrow, strong hairiness, and 3 to 5 or more leaf blade segments. The second axis is explained by compact, dense cushions with stocky leaves versus flat, loose cushions with slender leaves.

The leaf furrow is always present and complete in *S. delphinensis*, *S. cebennensis*, *S. exarata*, *S. pubescens* and *S. vayredana*. It is rarely present or it is incomplete in *S. moschata*, *S. giziana* and *S. lamottei*. The leaf hairiness is very dense in *S. delphinensis*, *S. cebennensis*, *S. pubescens*, *S. vayredana* and the Traversette Pass (“TP”) populations of *S. exarata*. In contrast it is either sparse or absent in *S. moschata* and *S. lamottei*. Note that *S. giziana* differs from *S. moschata* through

its stronger hairiness (not seen in Fig. 8, see Sup. Mat Table 3). For *S. exarata*, the leaf hairiness can be either dense or sparse. The cushion is very compact and domed in most *S. delphinensis*; compact but not domed in *S. cebennensis*, often loose and flat in *S. exarata*. Both loose and compact cushions were observed in *S. moschata*. Note that it is not possible to distinguish *S. vayredana* from *S. pubescens* using the morphological traits measured in this study, but both have a high number of leaf blade segments and slender leaves. These traits are also characteristic of *S. exarata* phenotypes from the Traversette Pass, which are well differentiated from other *S. exarata*.

## Discussion

With 400-500 species, *Saxifraga* displays its greatest diversity in mountain areas and has one of the highest endemism rates in the Alps (Aeschimann 2011). According to Ebersbach et al. (2017) and Folk et al. (2021), *Saxifraga*, and related genera, originated at the end of the Cretaceous period in rare cold habitats, probably restricted to arctic or alpine North America and then colonized Europe during the late Eocene (*ca* 33 Ma). Appearing in the early Miocene (*ca* 19 Ma), the *Saxifraga* sect. *Saxifraga* evolved with a constant rate of diversification, mainly driven by geographic isolation and bedrock shifts (Smyčka et al., 2022). The lineage studied here originated at the end of the Pliocene (3.8 million years ago; Fig. 2, Smyčka et al., 2022) and now forms several species, sometimes rare and endemic, in the European mountains.

We examined the distinctiveness of *S. delphinensis*, based on morphological, molecular, and genome-wide sequencing data. This narrow endemic is considered in the literature as morphologically well differentiated from closely related species occurring in the same region (Tison and Foucault 2014). Apart from the high-altitude areas in the Vercors and Devoluy, its distribution mainly corresponds to unglaciated areas during the last glacial maximum (Seguinot et al., 2018). This historical fact reinforces the hypothesis of a differentiation acquired over a long period. Our results support an origin probably earlier than the Würm and Riss glaciations. Indeed, the differentiation of *S. delphinensis* is significant with a characteristic leaf and cushion morphology, a plastid lineage deeply rooted in the Pleistocene, the existence of original and specific ITS haplotypes, and the grouping of almost all its RAD genotypes into one clade. However, despite this evolutionary distinctiveness, our various analyses further reveal a gradient of genetic admixture between *S. exarata* and *S. delphinensis* that supports fertility of F1 hybrids allowing for backcrosses. The introgression between *S. delphinensis* and *S. exarata* is particularly frequent in high-altitude sites in the Vercors (Grand Veymont, Royou) or in the Devoluy (Pic de Bure). Morphologically, this introgression is difficult to discern but some hybrids resemble *S.*

*delphinensis* (strong hairiness, stocky leaf shape and compact cushion) without having a dense, domed cushion. The final rebuttal to the complete isolation of *S. delphinensis* came from the Diois area (Eysahut and Saou populations) where introgression causes *S. delphinensis* to be polyphyletic in the RAD phylogeny or the ITS network. The genotypes from Eysahut (Diois) population are situated at an altitude of only 700 meters of elevation, on the extreme western margin of the Alps. At this site, *S. delphinensis* shows introgression from *S. exarata* with which it also shares an ITS haplotype. As far as we know, the closest *S. exarata* population is situated approximately 20 kilometres away. In Saou, *S. delphinensis*, living in sympatry with *S. moschata*, shows clear signs of introgression from *S. moschata*, with a shared ITS haplotype. This result is rather surprising since the introgression between *S. moschata* and *S. delphinensis* was not expected because of their very rather distinct morphologies in the field.

Our analysis also detected hybrids between *S. exarata* and *S. moschata* but at a lesser extent, despite their frequent sympatry. Recently Carnicero et al. (2023) also reported genetic admixture and nuclear-plastid mismatches for Pyrenean taxa, among which *S. moschata* and *S. pubescens*, but they concluded that the consequence of introgression was limited due to the partial sterility of hybrids and the absence of allopolyploidy that could fix hybrid genomes. By contrast to *S. moschata* from the Pyrenees (Mas de Xaxars et al. 2015), our study revealed a reduced ITS haplotypes diversity for *S. moschata* in south-west Alps and for its relatives *S. giziana* and *S. lamottei*. This indicates a recent diversification maintained by the geographical or ecological isolation. In the Jura Massif, the very low altitude (c. 350 to 500 m), favouring an earlier reproduction period than in the higher massifs, probably contributed to the genetic isolation in the periphery of *S. moschata* range. The present relative isolation of these populations combined with their low pollen-dispersal capacity or pollen limitation (García-Camacho and Totland 2009) indicates that introgression took place very early in their divergence and explain why *S. moschata* and *S. giziana* maintain their morphological differences (Mouly et al., 2020).

### **Limits and perspectives**

A limit of our study was the lack of chromosome counts and genome size estimations, which are known to be very variable in these taxa and more particularly in *S. exarata* (Vargas 2000) and *S. moschata* (Mas de Xaxars et al., 2015; Carnicero et al., 2023). Polyploidy is an important factor of evolution and speciation in plants and the separation of diploids from polyploids helps to optimize bioinformatics and phylogenomics by reducing the effect of paralogy (Karbstein et al., 2020; Carnicero et al., 2023). Paralogs, if they are present in the data, can lead to false phylogenetic relationships. However, we found the main phylogenetic clades rather robust in the

face of changing of assembly parameters or when we repeat the assembly and analysis by limiting admixture. Even if we cannot reject the possible effect of paralogy, we are confident that our phylogenetic results are valuable in the context of studying *S. exarata*, *S. moschata* and their relatives in the western Alps. In the phylogenies shown here, some relationships are robust and allow to confirm some differences among taxa. For example, the complete leaf furrow is a reliable characteristic to differentiate *S. moschata* from *S. exarata* and we observed that hybrids have partial leaf furrow. Leaf furrow, which is easier to observe on fresh material, can help to correct several confusions between these taxa. For example, *Saxifraga exarata* still appears in the Pyrenees in the GBIF database (<https://www.gbif.org> consulted 2/02/2023) or the POWO database (<https://powo.science.kew.org> consulted 14/03/2023) because *S. moschata* was formerly a subspecies of *S. exarata*.

The incomplete isolation taxa shown here is challenging to reconcile taxonomy with evolutionary theory. It challenges also our understanding of speciation and how species maintain their genetic integrity. Almost a century ago, Lotsy (1925) illustrated the syngameon concept with *Saxifraga* and *Betula* case studies. This concept has recently enjoyed renewed interest (Cannon and Petit 2020) because it emphasizes the role of adaptive introgression at ecological margins (Buck et al. 2022). The question of whether the syngameon is a relevant issue for *S. exarata* and its relatives remains to be answered. In parallel with this theoretical aspect, our results suggest that *S. exarata* probably encompass unknown diverging lineages that need to be studied more thoroughly. We confirmed the presence of at least two different forms of *S. exarata* in the Queyras, which were already suggested by Villars (1779). The population from the Traversette Pass is morphologically differentiated from *S. exarata*. According to RADseq based phylogeny, it forms a new lineage close to *S. delphinensis* and *S. cebennensis* but not completely isolated from *S. exarata* as shown by its shared co-ancestry or genetic admixtures with other *S. exarata*.

Last but not least, we must acknowledge that Ravaud proposed a close relationship of *S. delphinensis* with Pyrenean taxa based on morphological similarities (Ravaud, 1885). At first glance, this hypothesis is not supported by RADseq phylogeny or ITS network, but we cannot exclude an early influence of taxa from the Pyrenees or north Iberian mountains in the history of *S. delphinensis*. First, the plastome phylogeny (Smyčka et al., 2022) suggests a close relationship with the Iberian endemic *S. vayredana*, which is confirmed by ABBA-BABA tests and f-branch analysis of RADseq data. Second, *S. delphinensis* forms a well-supported clade with *S. cebennensis* and with the *S. exarata* population from the Col de la Traversette (Queyras) and the morphological characteristics of this clade (three to five leaf tips, strong leaf pilosity and complete leaf furrow) are also observed in certain Iberian taxa such as *S. pubescens* or *S.*



*vayredana*. Such influence is quite consistent with the hypothesis of Ravaud (1885). Future research will aim to establish the sequence of divergence and hybridization events that shaped this story during Quaternary range shifts through several mountain ranges in Europe.

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**Authors contribution**: AB, CR, LG, SA, CD, AM, MV and MC contributed to the sampling and to design of the study. AB performed the bioinformatics of RADseq and morphometric analyses. AB, KC, CM, JS and LA performed molecular data analyses. AB wrote the manuscript with the help of CR, SL, LG, SA, CD, AM and MV and all authors approved the manuscript.

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**Availability of data and materials**: Molecular data are available online (NCBI). Morphometrical and occurrence data are available in supplementary material.

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## Supplementary information

### 1. Supplementary material:

Table 1: Accessions for molecular data

Table 2: Data summary

Table 3: Morphological data

Table 4: Summary of ipyrad pilot assembly analyses

Table 5: Distribution of rDNA ITS haplotypes among *Saxifraga* samples

Table 6: Distribution of rDNA ITS haplotypes among taxa and sites

Table 7: ipyrad assembly stats

Table 8: Dstats of ABBA-BABA tests

Table 9: Dstats of ABBA-BABA tests performed between six *Saxifraga* species, with genotypes filtered to reduce the influence of admixture. The tests were guided by the phylogeny.

Figure 1: Morphometry of leaves

### 2. Supplementary results:

**Figure S1:** Phylogenetic relationships of 166 *Saxifraga* genotypes. IQ-TREE maximum likelihood trees based on a concatenated alignment of RADseq loci obtained by ipyrad assembly with a minimum of 24 individuals by locus (Min\_24) or a minimum of 48 individuals by locus (Min\_48).

**Figure S2:** Phylogenetic relationships versus genetic admixture of 166 *Saxifraga* genotypes. Maximum likelihood tree obtained with IqTree and based on a concatenated alignment of RAD locus (264,183 bp, 58,138 parsimony informative sites, 73% of missing data) and a TVM+F+I+G4 model. Blue squares are indicating bootstrap values above 95%. Numbers were added for comparison with RADpainter coancestry matrix. Admixture plots were obtained with SnpSift method on a filtered data set of 566 independent SNPs having a missing data rate of 33%. K=2 to K=6 groups are shown.

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## Figures and Table captions

**Figure 1:** Map of the 57 sampling sites (A) with a focus on south-west Alps (B) where *S. delphinensis*, *S. exarata*, and *S. moschata* were more intensively sampled.

**Figure 2:** Nuclear ribosomal and plastid data report different histories for *Saxifraga*. (A) Parsimony network of ITS haplotypes, the size of the circles is proportional to the number of samples and the colors indicate the taxa. The strokes on lines indicate longer branches that were shortened. (B) Plastome-based time tree adapted from Smyčka et al. (2022). Blue squares indicate bootstrap values above 95%. In (B) taxa names are those used by Smyčka et al. (2022), *S. exarata* subsp. *moschata* corresponds to *S. moschata* here.

**Figure 3:** Genome-wide phylogenetic relationships among 166 *Saxifraga* individuals revealed by an IQ-TREE maximum likelihood tree based on a concatenated alignment of 264,183 bp (58,138 parsimony informative sites, 73% of missing data) and a TVM+F+I+G4 model. Blue squares are indicating ultrafast bootstrap values > 95%. Numbers correspond to groups of high co-ancestry (RADpainter; Fig. 4). The tree with each genotype ID is in Sup. Res. Fig. S2.

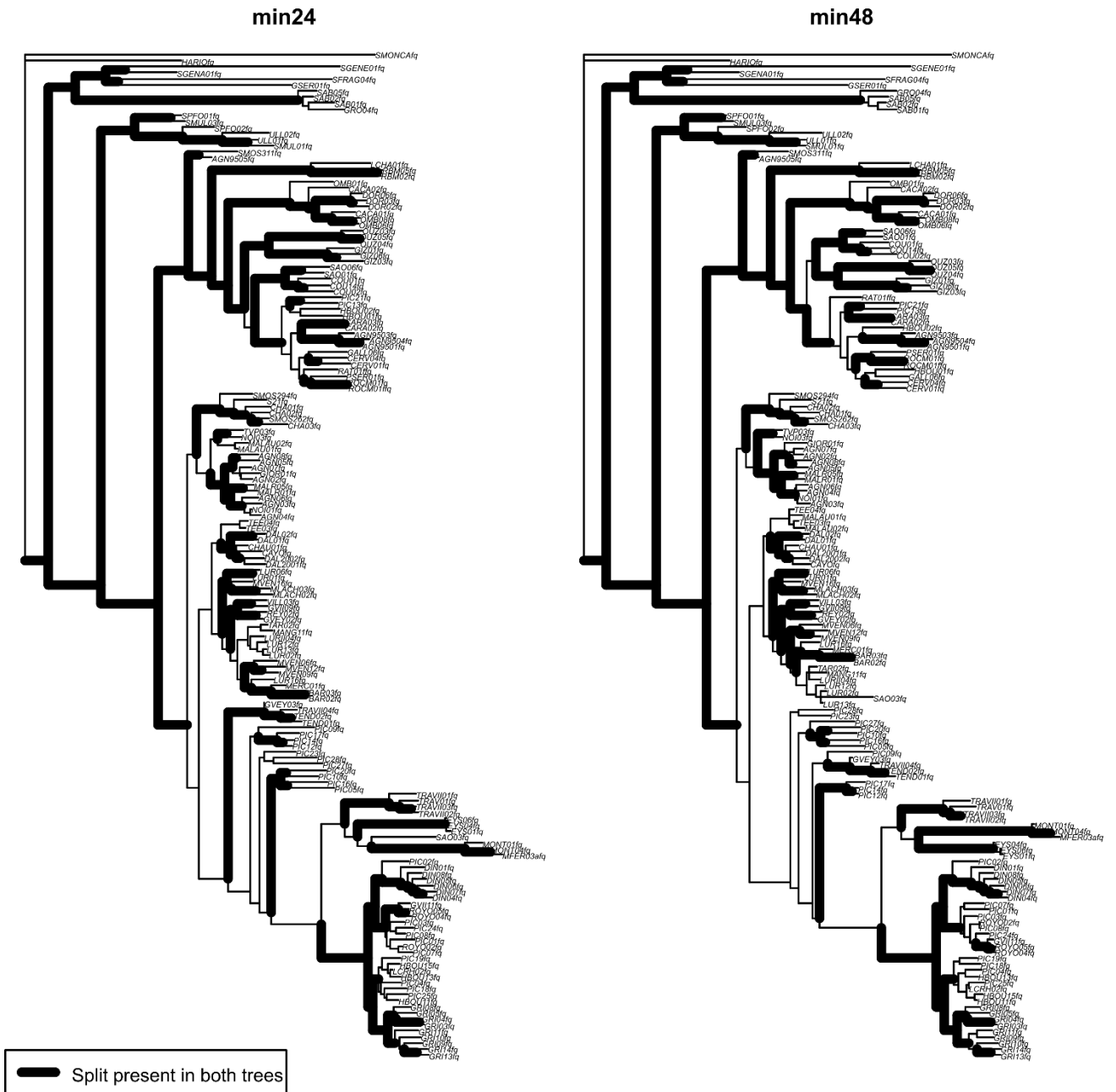
**Figure 4:** RADpainter co-ancestry matrix of 166 *Saxifraga* genotypes. The heat map was ordered according to the IQ-TREE phylogeny (left and Fig. 3) obtained on the same dataset. Genetic similarity between pairs of genotypes increases from blue to orange values. The black squares with numbers were reported for comparison with the phylogeny.

**Figure 5:** Admixture analysis (*Snmf*) performed on 99 RAD genotypes of 560 independent SNPs and 12% of missing data. Genotypes are ordered and colored according to their membership in 3 genetic clusters corresponding to *S. delphinensis*, *S. exarata* and *S. moschata* respectively. "P" and "T" indicate *S. exarata* genotypes from Pic de Bure and Traversette Pass respectively.

**Figure 6:** Coalescent-based phylogeny by SVDQuartet method of 32 *Saxifraga* genotypes. Exhaustive quartet sampling was conducted in a matrix of 7625 biallelic SNPs. The bootstrap values near the nodes were obtained with 200 replicates. Branch lengths are not meaningful.

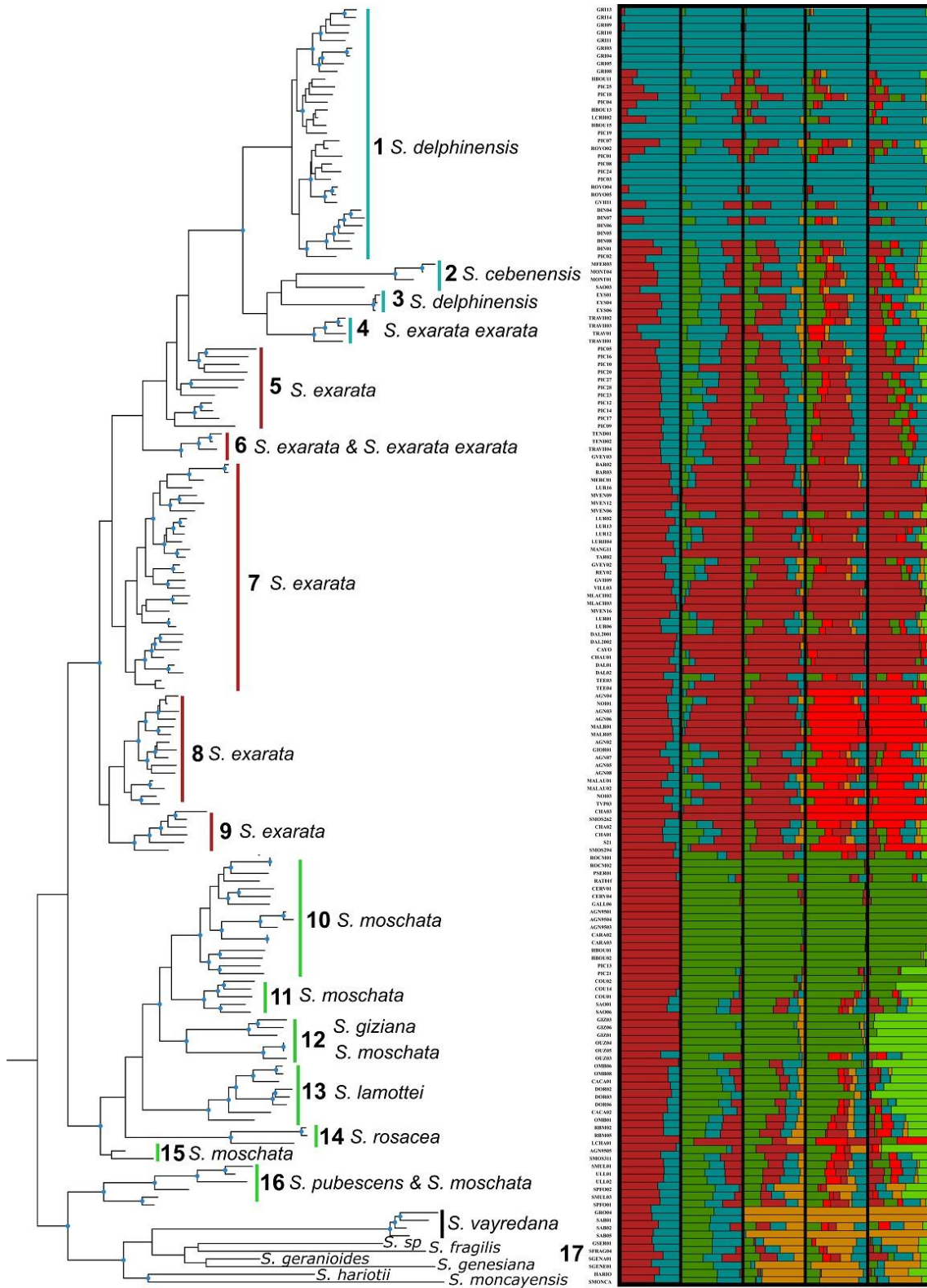
**Figure 7:** Results of Dsuite introgression tests. (A) to (C) ABBA-BABA tests performed on 166 genotypes assigned to 17 groups (Fig. 4), only Dstat with Zscores > 3 were kept. (A) Positive Dstat scores indicate an excess of ABBA pattern and an introgression from P3 to P2. (B) and (C) Violin plots of the Dstat distribution according to P2 or P3 for the three main species sampled here, *S. delphinensis* (DEL), *S. exarata* (EXA) and *S. moschata* (MOS). (D) to (E) Dsuite analyses guided by a coalescent-based species phylogenetic tree and performed on individuals with low genetic admixture. (D) Results of ABBA-BABA tests with z-scores > 3. (E) Results of f-branch analysis. The phylogenetic tree is shown on x and y axes, colored squares indicate significant excess of SNPs sharing caused by gene flow. White or grey squares represent non-significant values or trios incompatible with the phylogeny respectively.

**Figure 8:** Multivariate Hill & Smith analysis of vegetative polymorphism of *Saxifraga* based on principal components of leaf shapes (Leaf\_PC), number of leaf blade segments (Leaf\_tips), absence of leaf furrow (Furrow 1), incomplete leaf furrow (Furrow 1.5), complete leaf furrow (Furrow 2), multiple leaf furrow (Furrow3), leaf hairiness from nearly absent (Hairiness 1) to very dense (Hairiness 2) and cushion form from loose and open (Cushion 1) to compact and domed cushion (Cushion 4).



**Figure S1:** Phylogenetic relationships of 166 *Saxifraga* genotypes. IQ-TREE maximum likelihood trees based on a concatenated alignment of RAD loci obtained by ipyrad assembly with a minimum of 24 individuals by locus (Min\_24) or a minimum of 48 individuals by locus (Min\_48).





**Figure S2:** Phylogenetic relationships versus genetic admixture of 166 *Saxifraga* genotypes. Maximum likelihood tree obtained with IqTree and based on a concatenated alignment of RAD locus (264,183 bp, 58,138 parsimony informative sites, 73% of missing data) and a TVM+F+I+G4 model. Blue squares are indicating bootstrap values above 95%. Numbers were added for comparison with RADpainter coancestry matrix (Fig. 4). Admixture plots were obtained with *Snmf* method on a filtered data set of 566 independent SNPs having a missing data rate of 33%. K=2 to K=6 groups are shown.