Multiple markers, niche modelling, and bioregions analyses to evaluate the genetic diversity of a plant species complex

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Abstract

Background: The classification of closely related plants is not straightforward. These morphologically similar taxa frequently maintain their inter-hybridization potential and share ancestral polymorphisms as a consequence of their recent divergence. Under the biological species concept, they may thus not be considered separate species. The Petunia integrifolia complex is especially interesting because, in addition to the features mentioned above, its taxa share a pollinator, and their geographical ranges show multiple overlaps. Here, we combined plastid genome sequences, nuclear microsatellites, AFLP markers, ecological niche modelling, and bioregions analysis to investigate the genetic variability between the different taxa of the P. integrifolia complex in a comprehensive sample covering the entire geographical range of the complex.

Results: Results from molecular markers did not fully align with the current taxonomic classification. Niche modelling and bioregions analyses revealed that taxa were associated with different ecological constraints, indicating that the habitat plays an important role in preserving species boundaries. For three taxa, our analyses showed a mostly conserved, non-overlapping geographical distribution over time. However, for two taxa, niche modelling found an overlapping distribution over time; these taxa were also associated with the same bioregions.

Conclusions: cpDNA markers were better able to discriminate between Petunia taxa than SSRs and AFLPs. Overall, our results suggest that the P. integrifolia complex represents a continuum of individuals from distant and historically isolated populations, which share some morphological traits, but are established in four different evolutionary lineages.

Keywords: Adaptive radiation, Hybridization, Pampas, Phylogeography, Solanaceae, Speciation

Background

The delimitation of taxa at lower taxonomic levels is one of the greatest challenges of systematic biology [1], mainly due to the presence of highly conserved morphological traits [2] or morphological variation that does not follow traditionally defined species boundaries [3]. There are thus major open questions around the identity of species, the way to define them, and even the usefulness of the species concept in the context of conservation biology.

Closely related and young taxa frequently show morphological similarities and low levels of genetic differentiation due their evolutionary proximity. In this context, a species can be defined following different approaches: (1) tree-based methods define a species as a historical lineage according to phylogenetic concepts [4, 5]; (2) non-tree-based methods have been employed to uncover genetic structure associated with population isolation vs. species boundaries [6], which portrays aspects of population genetics that usefully elucidate the emergence of the species [7]; finally, (3) morphological distinctiveness can be used as a criterion to distinguish species from one another [8]. Importantly, to understand speciation in an evolutionary context, it is necessary to identify the factors that triggered the divergence process [9, 10]. In the case of the adaptive radiation of plants, the influence of different pollinators may be significant [11, 12]. Adaptive radiation
drives the evolution of different forms in response to different environmental conditions or habitats, thus often resulting in high levels of morphological or taxonomical diversity, combined with low genetic differentiation between related taxa [13]. On the other hand, many morphs or ecotypes are associated with geographically distant habitats, where reproductive isolation is a result of the physical separation, rather than arising from biological constraints. In such cases, the partial or full reduction of the gene flow between newly established and ancestral populations could allow the independent evolution and differentiation of their gene pools over time, without necessarily leading to reproductive incompatibility [10]. As a consequence, the sustained potential for inter-hybridization prevents the use of the biological species concept to define these taxonomical entities.

The plant genus Petunia Juss (Solanaceae) is recognised worldwide thanks to its prominent member Petunia hybrida (Hook.) Vilm, a widely cultivated interspecific hybrid considered to be a “supermodel” plant [14]. Petunia hybrida resulted from crosses between the white-flowered P. axillaris (Lam.) Britton, Sterns & Poggenb and P. interior T. Ando & Hashim as its main purple ancestor [15]. Petunia comprises 14 diploid species [16], predominantly native to southern Brazil, with the exception of two taxa (P. occidentalis R. E. Fr. and P. axillaris ssp. subandina T. Ando) that are found in the pre-Andean mountains in northern Argentina [17]. Two centres of diversity have been described for this genus [16]. Both of these represent transitional areas between tropical and subtropical formations and form a boundary for the distribution of many plant species: they represent a northern limit for many Austral-Antarctic elements (corresponding to the lowland grasslands of the Pampas region in Uruguay, the Argentinean province of La Pampa, and the Brazilian state of Rio Grande do Sul) and a southern limit for predominantly tropical elements (corresponding to the open highland fields of the Brazilian states of Rio Grande do Sul and Santa Catarina). Transitional areas concentrate a significant fraction of adaptive variation and are therefore an ideal setting for the study of speciation [18].

Phylogenetic analyses based exclusively or predominantly on plastid DNA (cpDNA) markers have revealed two major clades of Petunia species associated with elevation, a highland (> 500 m) and a lowland (< 500 m) clade [19–21]. On the other hand, studies using mostly nuclear markers have found two clades associated with corolla morphology (short vs. long corolla tube; Additional file 1: Figure S1F) [22–25]. Phylogeographic studies of Petunia species sharing morphological, ecological or geographical traits have confirmed an extremely recent divergence of Petunia species and are indicative of a diversification following colonization processes related to climate changes that took place during the Pleistocene [21, 26–31]. Moreover, the occurrence of artificial [32] and natural [26, 29, 33] hybrids between Petunia species suggest that intrinsic barriers to interspecific hybridization do either not exist, or else are rather weak. Where these barriers do exist, they are mainly prezygotic, and interspecific Petunia crosses yield viable diploid offspring, at least under controlled conditions [14]. Despite the apparent morphological and ecological diversity found within the genus Petunia, molecular studies have shown very low levels, or even an absence, of genetic diversity between species, a hallmark of recent adaptive radiation [19, 20, 22, 24].

The taxa of the Petunia integrifolia (Hook.) Schinz & Thellung are delimited by few specific morphological traits (Additional file 1: Figure S1 A-E; Additional file 2: Box S1; Additional file 3: Box 2). All its taxa are bee-pollinated, and the ranges of several of them are geographically close, with some overlaps [34]. Specimens with an intermediary morphology have been observed between taxa of this complex; however, they have not been characterised as hybrids using morphological or molecular traits [35]. It thus remains unknown whether these intermediary morphological traits represent intra-population variation or result from hybridization. In general, the Petunia integrifolia complex is characterised by a purple and infundibuliform corolla, violaceous pollen, and a stigma that is located between the anthers of the didynamous stamens. Due to the many morphological similarities across the P. integrifolia complex, there is little agreement on the number and taxonomic status of its taxa [16, 34, 36] (Additional file 2: Box S1).

Here, we define the P. integrifolia complex to include all taxa that share morphological traits with P. integrifolia, as well as those that have been classified as subspecies of P. integrifolia at least once: P. bajeensis T. Ando & Hashim, P. inflata R. E. Fr., P. integrifolia Ssp. depauperata R. E. Fr. (hereafter, P. depauperata), P. integrifolia Ssp. integrifolia (hereafter P. integrifolia), and P. interior (for a morphological overview see Additional file 1: Figure S1A–E and Additional file 3: Box S2). A number of authors also consider P. littoralis L. B. Sm. & Downs and P. riograndensis T. Ando & Hashim to be valid species in this group. However, it has been suggested to synonymise P. littoralis as P. depauperata, as a result of their morphological similarities and geographical distribution [36]. For the same reasons, P. riograndensis has also been synonymised as P. integrifolia [16]. The grouping of these pairs of taxa (P. littoralis + P. depauperata and P. riograndensis + P. integrifolia) is fully supported by the phylogeographic analysis of molecular data [27, 28], thus justifying the synonymy; however, using a phylogenetic approach, a different scenario has been proposed [25]. In this work, we consider these taxa as synonymous, as
suggested previously [16]. Indeed, our analyses did not find any differences to justify their separation. Phylogenetic analyses of the *P. integrifolia* complex have repeatedly returned incongruent results, and a comprehensive phylogeographic study evaluating the level of differentiation between the remaining species, *P. bajeensis*, *P. inflata*, and *P. interior*, is lacking. The patterns of geographic distribution are highly variable across the taxa of the *P. integrifolia* complex. They include the restricted and narrow distribution of *P. bajeensis*, a pattern of wide distribution across specific biogeographic provinces, as seen in *P. depauperata*, and the general wide distribution of *P. integrifolia* (Fig. 1). Generally, the taxa of the *P. integrifolia* complex are distributed across areas characterised by ecological differences. Paired with geographical and historical barriers, these ecological differences have promoted the establishment of distribution boundaries: *P. bajeensis* and *P. integrifolia* are found in the Pampas, *P. depauperata* is endemic to the South Atlantic Plain Coast, where it grows in salty and nutritionally poor soils (also in the Pampas), and *P. inflata* and *P. interior* are native to the Paraná province [34, 37, 38].

Here, we evaluated the genetic variability of the *P. integrifolia* complex, based on a comprehensive sample of taxa and individuals per taxon, which covers most of the geographical range and the complete contact zone for all taxa. We used a molecular approach based on non-coding cpDNA markers and polymorphic nuclear markers (SSR and AFLP), in tandem with ecological niche modelling and bioregions analyses. We sought to answer the following questions: (1) Does the current taxonomy of the *P. integrifolia* complex match the molecular diversity?; (2) Are the different markers equally effective in assigning individuals to morphological taxa?; and (3) Can ecological niche modelling explain the current species distribution as a result of ancient or current isolation or contact?

**Methods**

**Plant material**

We studied five taxa belonging to the *P. integrifolia* complex. Taxa were identified according to Stehmann et al. [16] (for morphological characterization and taxon distinction see Additional file 3: Box S2). Our sample was collected mainly in southern Brazil. It covers most of the taxa’s distribution, while focusing on the main area of convergence of the *P. integrifolia* complex, where all five taxa are found. The geographical coordinates of all collection sites were obtained using a global positioning system (GPS) unit. We deposited vouchers at the BHCB (Universidade Federal de Minas Gerais, Belo Horizonte, Minas Gerais, Brazil) or ICN (Universidade Federal do Rio Grande do Sul, Porto Alegre, RS, Brazil) herbaria (see Additional file 4: Table S1).

This work was conducted under Brazilian Federal Government permit MP 2.186/16 for the access of plant genetic information to develop evolutionary or taxonomic studies. No specific collection permits were required because none of the analysed taxa are federally listed as endangered or protected, and because no collection sites were located within protected areas.

**cpDNA sequencing and analysis**

Genomic DNA was extracted from dried young leaves following the CTAB protocol [39]. We analysed 26 individuals from three populations of *P. bajeensis*; 308
individuals from 69 populations of *P. depauperata*; 39 individuals from ten populations of *P. inflata*; 84 individuals from 15 populations of *P. interior*; and 114 individuals from 38 populations of *P. integrifolia*, making for a total of 571 individuals from 135 collection sites. The intergenic regions of cpDNA *trnG-trnS* and *psbA-trnH* were amplified, as previously described for *Petunia* species [26], using universal primers as described for each spacer [40, 41]. PCR products were purified using PEG 20% (polyethylene glycol; Sigma-Aldrich Co., St. Louis, MO, USA) [42] and sequenced in a MegaBACE1000 (GE Healthcare Bio Sciences Corp., Piscataway, NY, USA) automatic sequencer according to the manufacturer’s instructions and the DYEnamicET Terminator Sequencing Premix Kit (GE Healthcare). Some published sequences were also included in this work [15, 28]. All sequences were deposited in GenBank (see Additional file 4: Table S1). For each marker, both forward and reverse strands were checked using the CHROMAS 2.0 software (Technelysium, Helensvale, Australia), aligned with MEGA 6 [43] using the CLUSTAL W algorithm, and manually edited when necessary. Poly-A/T regions and small inversions are generally highly variable and homoplastic [44, 45] and were thus not considered in our analyses. Contiguous insertions/deletions (indels) of more than one base pair (bp) were treated as single mutational events [46].

The haploid plastid genome does not normally undergo recombination. This uniparental transmission means that it is inherited as a unit, and the two cpDNA intergenic spacers were therefore concatenated and treated as a single sequence in all analyses. The numbers of variable and informative sites in the manually edited alignment were obtained from the output of MEGA 6. Haplotypes were identified using DNASP 5.10.01 [47], and the relationships between haplotypes were estimated using the Median-Joining method [48] implemented in the NETWORK 4.612 software [49]. Indices of haplotypic (h) and nucleotide (n) diversities [50] and inter- and intra-taxonomic genetic variation by means of analysis of molecular variance (AMOVA) were obtained using ARLEQUIN 3.5.1.2 [51].

The dated haplotype phylogenetic trees were estimated by Bayesian inference as implemented in BEAST 1.8.4 [52]. Two independent runs were employed consisting of $1 \times 10^{8}$ Markov chain Monte Carlo (MCMC) iterations, sampling every 1000 generations under the HKY (Hasegawa, Kishino, and Yano) nucleotide substitution model with four gamma categories as obtained in JM O (Hasegawa, Kishino, and Yano) nucleotide substitution model with four gamma categories as obtained in JM O [53], according to the Akaike information criterion (AIC) [53]. We used a lognormal relaxed clock, Yule process as tree prior, and the prior for nucleotide substitution rates was as following: a gamma distribution prior with a shape parameter 1.6 and scale parameter $1.6 \times 10^{-9}$ as a prior, assuming an offset value of $1 \times 10^{-9}$ s/s/y. This rate was obtained by compiling rates of plastid markers for shrubs and herbaceous plants with a generation time of up to three years [27]. TRACER 1.6 [54] was used to check for convergence of MCMC and adequately effective sample sizes (ESS > 200) after discarding the first 10% of generations as burn-in. The final joint sample was used to estimate the maximum clade credibility tree using the TREEANNOTATOR software, part of the BEAST package, setting 0.5 as limit of posterior probability. Statistical support for the clades was established by assessing the Bayesian posterior probability (PP) with node heights summarized to reflect the posterior median. The FigTree 1.4.0 software package [55] was used to draw and edit the phylogenetic tree. As an outgroup, we used haplotypes of *Calibrachoa excellens* (R.E. Fr.) Wijmsma (Additional file 4: Table S1).

The demographic patterns of each taxon were assessed separately through a Bayesian Skyline Plot (BSP) [56], which considers historical changes in the effective size, as implemented in BEAST software. The priors for this analysis were the same as those used in the haplotype phylogenetic analysis, as previously described and including the sequences of all individuals in the analysis. The BSP was reconstructed using TRACER 1.6.

### Microsatellite genotyping and analysis

To estimate the genetic variability based on simple sequence repeats (SSR), we randomly selected individuals of all taxa of the *P. integrifolia* complex among those included into the cpDNA analysis. We sampled one population of *P. bajensis* (B1), three populations of *P. depauperata* (D1, D25, and D30), *P. inflata* (I1, I3, and I6), and *P. interior* (T3, T6, and T8), and five populations of *P. integrifolia* (G6, G7, G8, G24, and G31), for a total of 108 individuals (Additional file 4: Table S1). These populations were selected to cover the entire geographical distribution of these taxa [15, 16, 28].

We amplified seven microsatellite loci originally isolated in *P. depauperata* [57], according to Segatto et al. [15]; the individuals used in that work formed part of the sample described above. We estimated allele number (A), allele richness (AR), gene diversity (GD), inbreeding coefficient (FIS), and exclusive alleles (EX) using FSTAT [58]. We also evaluated the observed heterozygosity (H0), expected heterozygosity (He), polymorphic information content (PIC), the proportion of null alleles (NUL), and the deviation from the Hardy-Weinberg equilibrium using CERVUS 3.0.3 [59].

Intra- and interspecific genetic differentiation was assessed by AMOVA using ARLEQUIN. Population subdivision was estimated through F-statistic analogues ($R_{ST}$), and the statistical significance was determined based on $2 \times 10^{4}$ permutations.
Bayesian clustering analysis, as implemented in STRUCTURE 2.3.3, was used to establish population structure and individual ancestry [60]. As gene flow was expected among the natural populations, an admixture model using correlated allele frequencies was included [61]. We approximated the optimum number of groups (K) by varying K from 1 to 10, to exceed the number of different possible taxa (five), and ran the analysis ten times for each K. Each run was performed using 2.5 × 10⁶ burn-in periods and 10⁶ MCMC repetitions after burn-in. The optimal number of genetic clusters was determined using the ΔK method [62], which favours the model with the greatest second-order rate of change in lnPr (X|K), as implemented in the online STRUCTURE HARVESTER [63]. The results for the best K value were summarized using CLUMPP 1.1.2 based on the average pairwise similarity of individual assignments across runs with the Full-Search method, weighted by the number of individuals in each population, and G' statistics [64]. DISTRUCT 1.1 [65] was used to visualize the STRUCTURE results after processing with CLUMPP.

AFLP profiles and analysis

For the amplified fragment length polymorphism (AFLP) analyses, we randomly selected seven to 16 individuals per taxon among the individuals previously analysed for cpDNA, for a total of 51 individuals covering the entire geographic distribution of the P. integrifolia complex. AFLP analyses were carried out with EcoRI and MseI (Trul) enzymes (New England Biolabs, Hitchin, UK) and following standard protocols [66]. Initially, nine selective primer combinations of 5'-fluorophore-labelled EcoRI primers and unlabelled MseI primers were tested. The best three pairs (Fam-EcoRI-AAC/MseI-CAC, Hex-EcoRI-AAG/MseI-CAC, and Ned-EcoRI-AGC/MseI-CAG) were retained because they generated clear bands and high variability in preliminary tests for these taxa. The amplification products were subjected to capillary electrophoresis in an ABI 3500 XL automatic sequencer (Applied Biosystems, Foster City, USA). Fragments for each primer combination were processed and analysed through the GENEMAPPER 4.1 AFLP® software (Applied Biosystems), and individual ancestry [60]. As gene flow was expected among the natural populations, an admixture model using correlated allele frequencies was included [61]. We approximated the optimum number of groups (K) by varying K from 1 to 10, to exceed the number of different possible taxa (five), and ran the analysis ten times for each K. Each run was performed using 2.5 × 10⁶ burn-in periods and 10⁶ MCMC repetitions after burn-in. The optimal number of genetic clusters was determined using the ΔK method [62], which favours the model with the greatest second-order rate of change in lnPr (X|K), as implemented in the online STRUCTURE HARVESTER [63]. The results for the best K value were summarized using CLUMPP 1.1.2 based on the average pairwise similarity of individual assignments across runs with the Full-Search method, weighted by the number of individuals in each population, and G' statistics [64]. DISTRUCT 1.1 [65] was used to visualize the STRUCTURE results after processing with CLUMPP.

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Bayesian inference (BI) analysis was carried out in MrBayes 3.2 [69], to obtain a majority consensus tree, with F81-like as the evolutionary model for restriction sites [70] and equal rates as obtained in JMODELTEST, according AIC criterion, and using the coding bias option ‘iset = nobasencesites’. We ran a ten million generation chain, sampled at the length of every 1000 trees. The convergence was verified using TRACER 1.6 [54] after removing 10% of the sampling as burn-in.

Niche modelling and biogeographic region analysis

Georeferencing for the collection sites of all taxa in the P. integrifolia complex was acquired through direct field observation, SpeciesLink [71], and the Global Biodiversity Information Facility [72]. Records obtained from the databases were verified manually for incongruences, and only those matching species distribution were kept (Additional file 5: Table S2). Only records with global positioning system coordinates and detailed localisation were used. This did not compromise the analysis because it still allowed a complete coverage of the taxa’s distribution. Explanatory variables included a set of 19 bioclimatic RASTER layers at a 30 arc-second resolution (ca. 1 km² at the equator) from the WorldClim website version 1.4 [73]. We carried out ecological niche modelling (ENM) for the current species distribution under three contrasting past climate conditions [Last Interglacial (LIG) - ~120,000–140,000 years before present (yBP); Last Glacial Maximum (LGM) - 21,000 yBP, and Mid-Holocene (MH) - 6000 yBP] using a model of maximum entropy (MAXENT 3.3.3) [74]. The grid layers were cut so as to include the entire geographical distribution for all taxa and extracted through the RASTER package [75] implemented in R software [76]. Pearson's correlation between the variables was calculated using the RASTER package, and multicollinearity was minimized by selecting 10 bioclimatic variables pairs showing an R > 0.75 and presenting the lowest percentage of importance to the model in a preliminary run [77, 78]. We used this approach in order to identify areas with high suitability that could have harboured the taxa through the Pleistocene climate changes and ecological variables that could explain the geographical differentiation among taxa.
To understand the role of the biogeographical regions in the evolution of the species, we used the web application INFOMAP BIOREGIONS [79] to identify taxon-specific bioregions from our species distribution data, for discussion about terminology and concepts see Vilhena and Antonelli [80]. A species tree was used as input for the online INFOMAP BIOREGIONS software [81] to reconstruct the ancestral range for the *P. integrifolia* complex through Fitch’s method of maximum parsimony [82]. The WUSCHEL gene sequences from Segatto et al. [25] were used to construct the phylogenetic tree of the *Petunia* species, according to the classification of Stehmann et al. [16]. We used these markers because the phylogenetic tree obtained based on WUSCHEL sequences was the best supported among *Petunia* phylogenies and it is in agreement with previous works using nuclear and plastid markers [24]. We used BEAST 1.8.4 [52] applying the evolutionary model GTR with four gamma categories (obtained from JMODELTEST according AIC criterion), Yule speciation model, and a lognormal relaxed molecular clock to obtain the tree. Two runs of $10^8$ generations were carried out, sampling every 1000 generations, and the first 10% were discarded as burn-in. Markov chain convergence was ensured by values of ESS > 200, and the resulting plots were checked in TRACER. TREEANNOTATOR was used to select the maximum clade credibility tree. Statistical support for the branches was measured in Bayesian posterior probabilities (PP).

**Results**

**cpDNA diversity**

The combined cpDNA sequence data (*trnG-trnS* and *psbA-trnH*) for the 571 individuals yielded an 1127 bp alignment, with 65 variable and 50 parsimoniously informative sites resulting in 77 haplotypes. The median-joining network grouped species into four main groups (Fig. 2). We found low haplotype sharing among species, with only three haplotypes shared between two species each (Additional file 6: Table S3; Fig. 2): H34 and H41 were shared by *P. interior* and *P. inflata*, whereas H43 was observed in *P. inflata* and *P. integrifolia*. The largest number of haplotypes (27) was found in *P. depauperata*, the most frequent ones being H4 (found in 110 individuals) and H5 (found in 85 individuals). The smallest number of haplotypes (three) was observed in *P. bajeensis*, the species with the smallest number of both collection sites and individuals, and with the most restricted geographical range among the analysed taxa. For both *P. interior* and *P. depauperata*, haplotype differentiation clearly revealed subgroups, with different genetic components in these species.

**Fig. 2** Median-joining network of concatenated plastid haplotypes coloured by taxon. Transverse lines indicate mutations. The four haplogroups referred to in the text are presented in boxes.
Overall, the haplotypic diversity was $h = 0.93$, and the nucleotide diversity was $\pi = 0.32\%$. The highest values for haplotype and nucleotide diversity were observed for *P. interior* ($h = 0.91$ and $\pi = 0.34\%$), the lowest for *P. bajeensis* ($h = 0.52$ and $\pi = 0.09\%$). *Petunia depauperata* ($h = 0.77$ and $\pi = 0.13\%$), *P. integrifolia* ($h = 0.84$ and $\pi = 0.18\%$), and *P. inflata* ($h = 0.86$ and $\pi = 0.24\%$) presented the intermediate values of diversity. AMOVA results (Table 1) indicated greater interspecific than intraspecific variation (~64% vs ~36%).

The phylogenetic tree based on cpDNA haplotypes revealed two primary groups: one well supported (PP = 0.99) corresponding to the haplotypes of *P. interior* 45, 46, 53, 54, 55, and 56 (I) and another not supported clade corresponding to the rest of the haplotypes (II). Clade II has two subclades, one well supported corresponding mainly to haplotypes of *P. depauperata + P. integrifolia* and *P. bajeensis* (IIa), and another group mainly including haplotypes of *P. inflata* and *P. interior* (IIb), and a divergence time ca. 687 thousand years ago (Kya; Fig. 3a). These two groups of species inhabit different environments, the first one is from the Pampas grasslands, the second from the grasslands of the Highland inside the Atlantic rainforest, which forms a natural mosaic with the *Araucaria* forest. Within each group, the analyses supported several subclades. In group IIa, *P. depauperata* formed one clade (PP = 0.98; ~409 Kya) which further divided the haplotypes into Central, North, and South groups. *P. bajeensis* also formed a cluster with evidence for recent diversification (PP = 0.99; ~199 Kya) (Fig. 3a).

The Bayesian Skyline Plot (Additional file 7: Figure S2) showed different patterns for each group of species forming the two main clades of the Pampas and Highland grasslands, respectively, indicating periods of expansion and contraction. For *P. bajeensis*, the plot suggested a smooth yet constant population decline until ~25 Kya, from when population growth was observed until the present. On the other hand, *P. integrifolia* and *P. inflata* showed a recent population decline starting around 5 Kya until present. *Petunia interior* experienced a slight decrease over time followed by a recent expansion. However, these results should be taken with caution because of the large credibility intervals associated with population size estimates.

### Microsatellite diversity and structure

We analysed seven SSR loci. Six of these were polymorphic in *P. bajeensis*, *P. integrifolia*, and *P. inflata*; five were polymorphic in *P. interior*. All seven loci were polymorphic in *P. depauperata* (Additional file 8: Table S4). The average number of alleles found per locus per taxon was two for *P. bajeensis*, three for *P. integrifolia* and *P. depauperata*, four for *P. inflata*, and five for *P. interior*. The locus PID1G6 was monomorphic in *P. bajeensis*, as was PID3G5 in *P. inflata* and *P. interior*, and PID4C6 in *P. integrifolia* and *P. interior*. Genetic diversity evaluated across all loci was generally higher in

| Source of variation | Sum of squares | Variance component | Percentage of variation |
|---------------------|----------------|--------------------|------------------------|
| cpDNA               |                |                    |                        |
| Among groups        | 682.224        | 1.837              | 64.4                   |
| Within groups       | 574.663        | 1.015              | 35.6                   |
| SSR                 |                |                    |                        |
| Among groups        | 534.059        | 2.660              | 38.6                   |
| Within groups       | 779.188        | 4.234              | 61.4                   |
| AFLP                |                |                    |                        |
| Among groups        | 1444.143       | 24.713             | 17.2                   |
| Within groups       | 5348.117       | 118.847            | 82.8                   |

![Phylogenetic trees based on (a) haplotypes and (b) AFLP](image)
populations of *P. inflata* and *P. interior* than in the other species when estimated via allelic richness and gene diversity (Additional file 8: Table S4). In *P. bajeensis*, the inbreeding coefficient was negative when considering all loci, indicating an excess of heterozygotes. The other species showed negative values for one (*P. depauperata*) or two (*P. integrifolia*) loci and few positive values at the other loci. Private alleles were discovered at one locus for *P. integrifolia* and *P. inflata*, at two loci for *P. depauperata*, and three for *P. interior* (Additional file 8: Table S4). Only the locus PID1F1 in *P. interior* deviated from the Hardy–Weinberg equilibrium after a Bonferroni correction (*P* < 0.05; Additional file 9: Table S5). The frequency of null alleles was negative or close to zero for all loci calculated, suggesting an absence of null alleles (Additional file 9: Table S5). The AMOVA indicated that 38% of the genetic variation was a result of differences between taxa, while 62% was due to within-taxon variation (Table 1).

Based on the SSR data, we found the best value for K to be 2 in the *STRUCTURE* analysis (Fig. 4a), with the first cluster comprising *P. bajeensis* and various individuals of *P. interior* from a single collection site (T8), and the second one including all other species and the remaining populations of *P. interior*. For *K* = 5 (the number of morphological taxa included in our analyses; Fig. 4b), *P. bajeensis* showed a private genetic component. *Petunia interior* population T8 shared a genetic component with *P. inflata*; individuals of both *P. integrifolia* and *P. depauperata* had two different components, both of which were equally shared by both species. Finally, *P. inflata* and the remaining populations of *P. interior* showed three genetic components that did not distinguish them (one was exclusive to these taxa; the second was common to *P. integrifolia* and *P. depauperata*; the last was shared with the T8 population of *P. interior*).

**AFLP profiles and genetic structure**

Both gene diversity values (Hj) and percentage of polymorphic loci (#P_loc) found for the five taxa of the *P. integrifolia* complex fell into a narrow range (Hj = 0.00427–0.00512; #P_loc = 51.4–60%; Additional file 10: Table S6). AMOVA results (Table 1) indicated higher levels of intraspecific than interspecific variation (~83% vs ~17%).

The best number of groups in the AFLP-based clustering analysis using *STRUCTURE* was *K* = 3 (Fig. 4c), with all three genetic components present in all taxa, albeit in different proportions. Individuals belonging to *P. interior* were divided into two groups, of which one was very similar to *P. inflata*, whereas the others were close to *P. depauperata*. *Petunia bajeensis* displayed a pattern similar to that of *P. integrifolia* (Fig. 4c). For *K* = 5, each taxon presented a different genetic constitution, with most of the *P. bajeensis* and *P. integrifolia* individuals sharing the same genetic component, while *P. depauperata*, *P. inflata*, and *P. interior* were characterised by different predominant genetic components (Fig. 4d).

The phylogenetic tree based on all three AFLPs had well-supported branches, with the majority of the groups related to taxon morphology (Fig. 3b). The first clade was formed by most of the *P. bajeensis* individuals (PP = 1.0), while the second grouped individuals of all the other species of the *P. integrifolia* complex (PP = 1.0). This second clade was divided into two subgroups: the first included all individuals of *P. interior* and a single individual each of *P. inflata* and *P. bajeensis* (PP = 1.0). The second subgroup comprised all remaining individuals and species (PP = 0.85) and was further subdivided into two fully-supported clades, one formed only by *P. inflata* and the other by *P. integrifolia* and *P. depauperata* (Fig. 3b).

![Fig. 4](image)

*Fig. 4* Results of the *STRUCTURE* analyses based on microsatellites and AFLP markers. Each individual is represented as a vertical line partitioned into K-coloured components that represent an individual’s proportional assignment to one of the genetic clusters for SSRs and K = 2 (a); for SSRs and K = 5 (b); for AFLPs and K = 3 (c); and for AFLPs and K = 5 (d). *Petunia bajeensis* (baje); *P. integrifolia* Ssp. *integrifolia* (inte); *P. integrifolia* Ssp. *depauperata* (depa); *P. inflata* (inf); *P. interior* (teri)
Spatial distribution

The replicated models for the taxa resulted in AUC values >0.919 (Additional file 11: Table S7). MAXENT analyses suggested that the climate variables offering the best explanation for the geographic distribution varied between the different taxa of the *P. integrifolia* complex: the “mean temperature of the warmest quarter” made the greatest contribution to the model in *P. bajeensis* (61.2%) and *P. integrifolia* (28.3%), the “annual precipitation” for *P. depauperata* (82%), “temperature seasonality” for *P. inflata* (47.1%), and the “mean temperature of the coldest quarter” for *P. interior* (70.3%).

Ecological niche modelling over time (Fig. 5) showed that present and LIG climate conditions were even less suitable for *P. bajeensis* than MH and LMG conditions. The suitable areas for *P. integrifolia* were spread out over regions close to the Atlantic coast during the MH, and the modelling provided little evidence for the presence of this taxon in the coastal area under LGM and LIG conditions. The suitable areas for *P. depauperata* were larger under LGM conditions, when sea levels were lower, than under LIG, MH or present conditions. For *P. inflata*, the results of ENM were, for the most part, stable over LGM, MH, and present periods, but the suitable area was smallest in the LIG. The suitable area for *P. interior* was greater in size than in the present versus past-tested intervals except for the LIG. The stable areas for all taxa were very similar under LGM, MH, and present conditions, with the exception of *P. depauperata*, for which the suitable area has diminished in size since the LGM (Fig. 5). In general, suitable areas were smaller during the LIG for all taxa than during the other periods.

The analysis of biogeographical regions found *P. integrifolia*, *P. depauperata*, and *P. inflata* to be present in Bioregion 1; individuals of *P. integrifolia*, *P. inflata*, and *P. interior* were found in Bioregion 2; while in Bioregion 3, only *P. integrifolia* and *P. bajeensis* were found (Fig. 1a; Additional file 12: Table S8). The bioregions identified for all taxa of the *P. integrifolia* complex revealed that, while restricted to the Pampas region, *P. integrifolia* is present in all areas, whereas *P. inflata* and *P. interior* are found in the grasslands at higher elevations, in a region within the Araucaria forest domain. The reconstruction of the ancestral distribution (Fig. 1b) fitted the distribution of *P. integrifolia* and included the entire complex. All identified bioregions encompassed the suitable areas for each taxon that remained stable over time according to ENM results.

![Fig. 5 Ecological Niche Modelling for the Petunia integrifolia complex. Black dots represent the localities where species were found. Models were calibrated for the present, Mid-Holocene, Last Glacial Maximum (LGM) and Last Interglacial (LIG). Darker areas represent higher probabilities of occurrence. Country boundaries obtained from http://www.diva-gis.org](image-url)
Discussion
The Petunia integrifolia complex inhabits a transitional area [16]. Transitional areas are usually characterised by the presence of heterogeneous habitats; in the present case, three bioregions are found within a small geographical area (Fig. 1). Environmental heterogeneity is considered a trigger for speciation processes; here, speciation may have occurred without promoting deep morphological changes (Additional file 3: Box S2). On the other hand, phenotypic plasticity and ecological or geographical adaptations to a heterogeneous environment (luminosity, open or forested areas, and different types of soil) might explain some of the morphological variation among the populations of these closely related taxa of recent evolutionary origin.

Here, we compared the genetic diversity of five taxa of the Petunia integrifolia complex based on different molecular markers (sequences of plastid gene spacers, nuclear microsatellites, and AFLP profiles). We focused on phylogenetic context, group cohesiveness, and niche conservatism, considering multiple individuals and populations per taxon. We evaluated the ability of the different markers to correctly assign individuals to their corresponding morphological-based taxonomic group and modelled niche distribution in an effort to identify the drivers of diversification and taxon delimitation. Nevertheless, the different molecular markers analysed in this study demonstrated differences in their potential to correctly assign individuals to their respective morphological taxa.

A recurrent pattern among Petunia species and subspecies is the absence of reciprocal monophyly and a large degree of interspecific plastid haplotype sharing, even between species that form morphologically well-delimited taxa [26, 29, 30]. Despite the minimal morphological differentiation within the Petunia integrifolia complex (Additional file 3: Box S2), plastid marker analyses confirmed the separation of Petunia integrifolia, Petunia depauperata, and Petunia bajeensis (Fig. 2) with a minimum of ancestral sharing.

Moreover, the Petunia integrifolia subspecies (Petunia integrifolia and Petunia depauperata) are equally distant from each other as Petunia integrifolia and Petunia bajeensis (Figs. 2 and 3). Based on this, we suggest considering Petunia integrifolia and Petunia depauperata as two independent species, which can be identified based on their habit (procumbent in Petunia depauperata and decumbent in Petunia integrifolia) as well as the shape of their leaves and their geographical distribution in nature (Additional file 3: Box S2). The flowers of Petunia depauperata are significantly smaller than those of Petunia integrifolia [35], and the corolla is darker in Petunia depauperata. Changes in nomenclature are necessary and shall be discussed in an appropriate forum.

For Petunia interior and Petunia inflata, cpDNA analysis revealed a more complex pattern. Four Petunia interior populations from the northernmost region of the species’ distribution, where Petunia inflata does not occur, and two populations from its southernmost area, which are near populations of Petunia inflata (see Fig. 1), form a separate group. However, the majority of populations of both species are grouped together (Figs. 2 and 3). In accordance with the results of the Bayesian Skyline Plot (Additional file 7: Figure S2) and niche modelling (Fig. 5), this pattern could be the result of a recent population expansion of Petunia interior to the south. However, because plastid haplotype sharing among the populations is rare, recent hybridization does not offer the best explanation for this pattern. Haplotype sharing has been described as a consequence of recent interspecific hybridization of Petunia axillaris and Petunia exserta [15, 26]; however, in that case, it was paired with various morphological signals. Other authors have found plastid haplotype sharing in the absence of hybridization, a finding that likely reflects differing patterns of genetic drift between recently diverged species [21]. Such incomplete lineage sorting supports the idea that insufficient time has elapsed to allow complete divergence of Petunia plastids [27]. The most recent common ancestor (MRCA) of the Petunia genus is thought to have lived a minimum of 1.3 Ma ago (Mya) [21] and a maximum of 2.8 Mya [83]. The divergence of Petunia integrifolia and Petunia depauperata has been estimated at 400 Kya [27], and in the present work, group II is estimated to have diverged around 687 Kya, while Petunia bajeensis diverged from the other taxa ~199 Kya, making incomplete lineage sorting a realistic possibility [84–86]. Alternatively, these results can be interpreted as an indication that genetically, these individuals belong to the same species, and that morphological differences may be attributed to phenotypic plasticity. We thus suggest a taxonomic review.

Microsatellites are highly variable and can be highly effective in discriminating closely related species [87]. On the other hand, even though AFLPs generally have lower mutation rates than SSRs, studies in ferns [88], mangroves [89], and Quercus [90] have shown that AFLPs are more efficient at distinguishing populations than microsatellites. To date, there have been few studies comparing the effectiveness of microsatellites and AFLPs in discriminating plant species. It has been suggested that, in animals, both types of markers have an approximately 70% chance to correctly discriminate between species and assign individuals to their respective groups [91]. In the present study, we found SSRs and AFLPs (Figs. 3 and 4) less effective than cpDNA haplotypes (Fig. 2) in discriminating between the species of the Petunia integrifolia complex, according to the currently accepted morphological classification. The genetic structure revealed by the AMOVA highlights this difference among markers (Table 2): whereas 64.4% of the total cpDNA variation was between groups, the corresponding variation was only 38.6% for SSRs and 17.2% for AFLPs. In addition, we believe that the high variability observed within taxa could explain the discrepancies found
between the AFLP-based tree and STRUCTURE analyses and the results obtained from these analyses using the other markers (Figs. 3 and 4).

The results of the STRUCTURE analyses using AFLPs and SSRs agree with the complicated morphological classification of the *P. integrifolia* complex and are in line with the phenotypic continuum found for a range of morphological traits (Additional file 1: Figure S1 and Additional file 3: Box S2). Plastid markers usually show more local structure, mostly because their maternal inheritance reflects the importance of the habitat in lineage isolation [29, 92].

It is important to highlight that the geographical distribution of *P. bajeensis* falls completely within the range of *P. integrifolia*. However, there was no suggestion of population reduction (Additional file 7: Figure S2) or hybridization (based on cpDNA and SSRs analyses) between *P. bajeensis* and *P. integrifolia*, and the populations of *P. bajeensis* occupy a different bioregion, where some populations of *P. integrifolia* also occur. The ecological and/or environmental factors responsible for this isolation remain to be determined. Despite morphological similarities, phylogenetic analyses [24, 25] found that *P. bajeensis* is not closely related to the remaining taxa of the complex; it is the sister species of the highland subclade in the short corolla tube clade.

The *P. integrifolia* complex is comprised of four geographically isolated and stable lineages (Fig. 5), which are associated to different bioregions (Fig. 1). The fact that these closely related taxa occupy very different habitats, such as the coastal plains, the Pampas, and the Atlantic forest, suggests that eco-geographical isolation has acted as one of the most significant reproductive barriers within the genus *Petunia*. The dynamic of new adaptations spreading through a population might be influenced by the levels of gene flow and demographic structure [93, 94]. The phylogeographic patterns found in the *P. integrifolia* complex could therefore be the result of low levels of gene flow through seed dispersion, either by free fall or barochory [16, 95], combined with differential selection in a heterogeneous environment. This is consistent with the fact that these species share a pollinator, and with the absence of reproductive barriers, at least in controlled situations [16]. Up to date, there are no studies on the pollination biology of these species, and the available data come from sporadic field observations.

The geographical area inhabited by the taxa of the *P. integrifolia* complex is of a variable topography and is characterised by a humid subtropical climate (Cfa) [96]. It is a region of transition between two important Neotropical phytogeographical formations: the Pampas grassland [97] and the Brazilian Atlantic Forest (BAF) [98]. In the central portion of Rio Grande do Sul, for example, populations of *P. integrifolia* and *P. inflata* are located only ca. 22 km from one another, but while *P. integrifolia* inhabits the Pampas grasslands up to a maximum elevation of 167 m, *P. inflata* and *P. interior* are found in mosaics of grasslands and BAF above 450 m, in the *Araucaria* forest domain. Both the Pampas grassland and the BAF have undergone a series of changes in range and floristic composition during the Quaternary [99]. Even during warm periods, open fields naturally covered the Pampas grasslands with small patches or long strips of gallery forests along rivers and on the escarpments of low-elevation sandstone hills [100]. In the southern BAF, typified by mosaics of high-elevation grasslands and forests, the dynamics of the Quaternary are also well documented through pollen records [100, 101]. During glacial periods, forests were restricted to more humid areas along watercourses and slopes, enabling the expansion of the grassland areas. In contrast, the conditions of the interglacial periods allowed forests to gain more territory over the grasslands [99]. The plant migrations triggered by the Quaternary climate oscillations, combined with the non-uniform landscape composition, may have contributed to the diversification of the evolutionary lineages of the *P. integrifolia* complex. This dynamic also underlies the diversification and range expansion of other *Petunia* species [21] and of the genus as a whole [24], and mirrors that of other plant species from the same region [31, 102, 103], in line with the theory of Andean components in the southern Brazilian flora [104]. The differentiation between habitats and the stability found in niche modelling analysis reinforce the idea that there are four genetic lineages: *P. integrifolia*, *P. depauperata*, *P. bajeensis*, and *P. inflata + P. interior*.

**Conclusions**

The species is one of the fundamental units of biology. In nature, plant species identification is often complicated by the presence of a continuum of morphological traits across taxa. Overall, our results suggest that the *Petunia integrifolia* species complex represents four historically isolated lineages. We suggest *P. depauperata* be elevated to species level and *P. inflata* and *P. interior* status should be investigated in a broader approach including morphological data. We also point out *P. bajeensis* is not part of the complex because its phylogenetic position, which could be attributed to incomplete lineage sorting. The cpDNA markers were more efficient to discriminate among *Petunia* taxa classified based on morphological traits than SSRs and AFLPs. The Niche modelling and bioregions analyses revealed that the habitat played an important role in preserving species boundaries over time. The sharing of morphological, ecological, and genetic characters do not necessarily accommodate a definition according to the biological species concept,
but are completely in agreement with the definition of a species as a metapopulation. Beyond the ornamental \textit{P. hybrida}, a popular model of molecular and developmental genetics, wild \textit{Petunia} species thus serve as a remarkable case study of evolutionary biology.

### Additional files

Additional file 1: Figure S1. Morphologies of the species of the \textit{Petunia integrifolia} complex and evolutionary relationships among \textit{Petunia} species. (A) \textit{P. bajeensis}; (B) \textit{P. integrifolia} spp. depauperata; (C) \textit{P. infleta}; (D) \textit{P. integrifolia} spp. integrifolia; (E) \textit{P. interior}; (F) Phylogenetic tree adapted from Reck-Kortmann et al. (2014): the long corolla tube clade is shown in pink, the short corolla tube clade is shown in black; species of the \textit{P. integrifolia} complex are in bold type. (PNG 4747 kb)

Additional file 2: Box S1. Simplified taxonomic changes of the \textit{Petunia integrifolia} complex over time. (DOCX 19 kb)

Additional file 3: Box S2. Morphological characteristic of the \textit{Petunia integrifolia} complex. (DOCX 14 kb)

Additional file 4: Table S1. Sample Collection. (DOCX 111 kb)

Additional file 5: Table S2. Collection sites used in the Environmental Niche Modelling (ENM) analysis to the taxa of \textit{Petunia integrifolia} complex. (DOCX 47 kb)

Additional file 6: Table S3. Plastid haplotypes observed for each species of the \textit{Petunia integrifolia} complex. (DOCX 13 kb)

Additional file 7: Figure S2. Changes in the effective population size over time (years ago) for each species of the \textit{Petunia integrifolia} complex. Bayesian skyline plot showing the effective population size fluctuation throughout time for \textit{P. bajeensis}, \textit{P. integrifolia} spp., \textit{integrifolia}, \textit{P. infleta} and \textit{P. interior} (solid line, median estimators of Ne; blue lines, confidence interval). (TIFF 70 kb)

Additional file 8: Table S4. Diversity indices for the \textit{Petunia integrifolia} complex based on microsatellite variation. (DOCX 17 kb)

Additional file 9: Table S5. Microsatellite-based diversity indices for species of \textit{Petunia integrifolia} complex. (DOCX 14 kb)

Additional file 10: Table S6. AFLP-based genetic diversity of the \textit{Petunia integrifolia} complex. (DOCX 12 kb)

Additional file 11: Table S7. AUC values and standard deviations of the Ecological Niche Modeling. (DOCX 14 kb)

Additional file 12: Table S8. Selected bioregions for each species of the \textit{Petunia integrifolia} complex. (DOCX 12 kb)

### Abbreviations

- \textit{PI}: loci; \textit{AFLP}: polymorphic loci; \textit{A}: Allele number; \textit{AFLP}: Amplified fragment length polymorphism; \textit{MIC}: Akaike information criterion; \textit{AMOVA}: Analysis of molecular variance; \textit{AR}: Allele richness; \textit{AUC}: Area under the curve; \textit{BAF}: Brazilian Atlantic Forest; BHCB: Universidade Federal de Minas Gerais Herbarium; \textit{Bl}: Bayesian inference; \textit{bp}: Base pair; \textit{BSP}: Bayesian Skyline Plot; \textit{Cfa}: Humid subtropical climate; \textit{cpDNA}: Plastid DNA; \textit{CTAB}: Cetyltrimethylammonium bromide; \textit{ENM}: Ecological niche modelling; \textit{ESS}: Effective sample size; \textit{Ex}: Exclusive alleles; \textit{Fis}: Inbreeding coefficient; \textit{GD}: Gene diversity; \textit{GPS}: Global positioning system; \textit{GTR}: General Time reversible; \textit{h}: Haplotypic diversity; \textit{He}: Expected heterozygosity; \textit{Hj}: AFLP Nei’s gene diversity; \textit{HKY}: Hasegawa, Kishino, and Yano nucleotide substitution model; \textit{HO}: Observed heterozygosity; \textit{ICN}: Universidade Federal do Rio Grande do Sul Herbarium; \textit{indels}: Insertions/deletions; \textit{K}: Groups tested in model; \textit{LIG}: Last Interglacial; \textit{MCMC}: Markov chain Monte Carlo; \textit{MH}: Mid-Holocene; \textit{MRCA}: Most recent common ancestor; \textit{Mya}: Million years ago; \textit{NUL}: Proportion of null alleles; \textit{PCR}: Polymerase Chain Reaction; \textit{PEG}: Polyethylene glycol; \textit{PIC}: Polymorphic information content; \textit{PP}: Posterior probability; \textit{Rst}: F-statistic analogues; \textit{SSR}: Simple sequence repeats; \textit{r}: Nucleotide diversity

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### Availability of data and materials

Microsatellite and AFLP datasets used and analysed during the current study are available from the corresponding author on reasonable request. All cpDNA sequences were deposited at GenBank, and herbarium vouchers were deposited for each population and species (see main text for more information).

### Authors’ contributions

LBF conceived the study; ALAS, MR-K, and CT generated the data and ran the statistical analyses; LBF provided reagents and equipment for developing the experiments. All authors have commented on and approved the final manuscript. Collectively, the group is interested in investigating evolutionary processes of plants.

### Ethics approval and consent to participate

The vouchers were deposited at the BHCB (Universidade Federal de Minas Gerais, Belo Horizonte, Minas Gerais, Brazil) or ICN (Universidade Federal do Rio Grande do Sul, Porto Alegre, RS, Brazil) herbaria. This work was conducted under Brazilian Federal Government permit MP.2.186/16 for the access of plant genetic information to develop evolutionary or taxonomic studies.

### Consent for publication

Not applicable

### Competing interests

The authors declare that they have no competing interests.

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