Origin and Diversification of South American Polyploid Silene Sect. 
Physolychnis (Caryophyllaceae) in the Andes and Patagonia

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The Andes are an important biogeographic region in South America extending for about 8000 km from Venezuela to Argentina. They are – along with the Patagonian steppes – the main distribution area of ca. 18 polyploid species of Silene sect. Physolychnis. Using nuclear ITS and plastid psbE-petG and matK sequences, flow cytometric ploidy level estimations and chromosome counts, and including 13 South American species, we explored the origin and diversification of this group. Our data suggest a single, late Pliocene or early Pleistocene migration of the North American S. verecunda lineage to South America, which was followed by dispersal and diversification of this tetraploid lineage in the Andes, other Argentinian mountain ranges and the Patagonian steppes. Later in the Pleistocene South American populations hybridized with the S. uralensis lineage, which led to allopolyploidisation and origin of decaploid S. chilensis and S. echegarayi occurring at high elevations. Additionally, we show that the morphological differentiation in leaf shape correlated with divergent habitats (high elevation Andes vs. lower elevation Patagonian steppes) is also supported phylogenetically, especially in the ITS tree. Lastly, the species boundaries among the narrow-leaved Patagonian steppe species are poorly resolved and need more thorough taxonomic revision.

Keywords: Andes, chromosome counts, ITS, Patagonia, phylogeny, relative genome size

INTRODUCTION

Polyploidy has significantly contributed to the diversification and radiation of flowering plants (Soltis et al., 2009, 2016; Wood et al., 2009; Husband et al., 2013; Madlung, 2013) and at least two polyploidisation rounds have occurred in all angiosperms (jiao et al., 2011). Polyploid lineages often exhibit complex relationships among each other and with their lower-ploid ancestors (e.g., Popp et al., 2005; Marhold and Lihova, 2006; Brysting et al., 2011; Frajman et al., 2016). Recurrent formation of polyploids, (epi)genetic, transcriptomic and genomic changes as well as morphological, geographic and ecological divergence following polyploidisation are considered significant processes in the evolution of polyploids (Leitch and Bennett, 2004; Adams and Wendel, 2005; Johnston et al., 2005; Paridos et al., 2010; Sonnleitner et al., 2010; Balao et al., 2011; Weiss-Schneeweiss et al., 2013), and obviously increase taxonomic complexity. On the other hand,
polyploids have been suggested to have lower speciation and higher extinction rates than diploids (Mayrose et al., 2011). Consequently, Arrigo and Barker (2012) suggested that the high abundance of polyploids is a consequence of their high formation rate rather than of accelerated diversification of polyploids, but the inconsistent patterns of evolution following polyploidy make it challenging to develop a “unified theory” of polyploidy (Soltis et al., 2016).

The genus Silene L. (Caryophyllaceae) includes 800 to 900 mostly diploid (2n = 2x = 24) species distributed mainly across the Northern Hemisphere (Petri and Oxelman, 2011; Oxelman et al., 2013). Extensive polyploidisation has accompanied evolution of North American (Popp and Oxelman, 2007) and – to a lesser extent – arctic (Popp et al., 2005) and Asian (Petri and Oxelman, 2011) members of Silene sect. Physolychnis (Benth.) Bocquet, which includes diploids, tetraploids, hexaploids and octoploids (Petri and Oxelman, 2011; Rice et al., 2015). In North America most native species are tetraploids or octoploids (Popp and Oxelman, 2007), whereas little is known about the role of polyploidisation in the evolution of South American Silene sect. Physolychnis. The section includes species with upright stems, irregular thyroid inflorescences and mostly small, often dissected petal limbs (Petri and Oxelman, 2011). It was most thoroughly studied by Bocquet (1969), but based on molecular studies (e.g., Popp et al., 2005; Popp and Oxelman, 2007; Jenkins and Keller, 2010; Rautenberg et al., 2010; Petri and Oxelman, 2011) additional species were included in the section, thus establishing a morphologically rather diverse but phylogenetically well-supported monophyletic group (Petri and Oxelman, 2011).

For the Andes, the Andean Precordillera and other mountain areas (e.g., Sierra de Córdoba, Sierra de la Ventana) as well as for the Patagonian lowlands of South America, 17 species of S. sect. Physolychnis were recognized by Bocquet (1969) and two more by Pedersen (1984), namely S. melanotomatia Pedersen and S. antarctica (Kuntze) Pedersen; the latter was considered conspecific with S. patagonica (Speg.) Bocquet by Pedersen (1984), but this might be a result of eco-morphological differentiation and/or evolutionary divergence.

To disentangle the origin and diversification of South American Silene sect. Physolychnis, we use phylogenetic analyses of nuclear ITS and plastid psbE-petG and matK sequences, flow cytometric genome size estimations and chromosome counts, including 13 South American species. We address the following questions: (1) are the South American species of Silene sect. Physolychnis monophyletic, resulting from a single migration from North America?; (2) what role did polyploidisation play in the evolution of South American species and which lineages have been involved in their diversification?; and (3) is the morphological differentiation, especially the leaf shape, correlated to the evolutionary history or rather to divergent habitats?

**MATERIALS AND METHODS**

**Plant Material**

21 populations belonging to 11 species were collected in South America, mostly Argentina and Chile, between 2006 and 2008. In most cases silica-gel material and in all cases herbarium vouchers were collected, often with ripe seeds. Additionally, seven populations (2, 4, 5, 8, 13, 14, 22) belonging to five species were sampled from herbaria and 17 GenBank sequences from previous studies were added, thus resulting in a dataset of 29 populations belonging to 13 species (Figure 1 and Supplementary Table S1). The species were identified following Bocquet (1969) and Pedersen (1984); several accesses were collected at the localities listed by the two authors, thus facilitating the identification. Outgroup species, mostly from North America, were sampled from different herbaria and we also included 106 GenBank sequences from previous studies (Supplementary Table S2).
FIGURE 1 | Sampled populations of Silene in South America, detailed information is given in Supplementary Table S1. The frame in the left map shows the position of the area depicted in the right map. Leaf shapes are depicted next to the sampling points. The photos represent some of the analyzed species in their natural habitats.

Chromosome Numbers
Chromosome numbers were determined for six Silene species (Supplementary Table S1). Seeds were germinated on moist paper in Petri dishes and actively growing root tip meristems were collected and pre-treated with 0.002 M aqueous solution of 8-hydroxyquinoline for 2.5 h at room temperature and 2.5 h at 4°C, fixed in ethanol and glacial acetic acid (3 : 1), and stored at −20°C until use. Meristems were hydrolysed in 5N HCl for 20 min at room temperature, washed with tap water and stained with Schiff’s reagent for 1 h (Weiss-Schneeweiss et al., 2009). Chromosome spreads were prepared by squashing a stained meristem in a drop of acetic acid (60%) under the coverslip, and analyzed using an AxioImager M2 microscope (Carl Zeiss, Vienna, Austria). Images were acquired with a CCD camera and files processed using AxioVision ver. 4.8 (Carl Zeiss, Vienna, Austria) using only those functions that apply to the whole image equally.

Relative Genome Size (RGS) Estimation With Flow Cytometry
Relative genome size was typically measured for four to six individuals per population of 18 populations representing nine species (Supplementary Table S1). Relative fluorescence intensity of silica-dry samples was estimated by DAPI flow cytometry according to the simplified one-step methodology using Otto buffers as detailed in Doležel et al. (2007). Bellis perennis (2C = 3.38 pg; Schönswetter et al., 2007) served as an internal reference standard. A Partec PA II cytometer equipped with a UV HBO lamp was used to record fluorescence values of 3000–5000 particles, depending on the sample quality. Due to the relatively frequent left-hand shoulder of the sample peak, manual gating was applied before the standard/sample peak ratio was calculated with FloMax software (Partec, Germany). Coefficients of variation of the G0/G1 peaks of the internal standard and the sample varied from 1.98 to 3.23% (mean 2.46%) and from 2.52 to 7.82% (mean 4.78%), respectively.

DNA Extraction, PCR and Sequencing
Total genomic DNA was extracted from herbarium specimens or silica-gel dried material following the protocol described by Oxelman et al. (1997) and purified using the QIAquick purification kit protocol (QiaGen).

PCRs were performed as described by Frajman et al. (2009). PCR products were purified using Multiscreen PCR (Millipore) according to the manufacturer’s protocol and then sequenced with the PCR or nested primers (see Frajman et al., 2009) using the BigDye Terminator Cycle Sequencing Kit (Applied Biosystems), and visualized on ABI 3700 or ABI3730XL (Applied
Biosystems). Sequencing reactions were performed by Macrogen Inc.

Contigs were assembled and edited using Geneious Pro 5.3.6 (Drummond et al., 2011). Base polymorphisms were coded using NC-IUPAC ambiguity codes as described by Frajman and Oxelman (2007). GenBank numbers of ITS sequences are presented in Supplementary Tables S1, S2.

**Phylogenetic Analyses**

South American Silene accessions were first aligned to all Sileneae sequences in BoxTax database (Oxelman et al., 2013) available in 2016 using Geneious Pro 5.3.6 (Drummond et al., 2011) and the three datasets were analyzed using FastTree v.2.1.5 (Price et al., 2010) with the GTR model and Gamma20 likelihoods. Based on the inferred trees (not shown) as well as the study of Popp and Oxelman (2007) all South American accessions, their most closely related species and some additional taxa representing different groups of Silene sect. Physolychnis were sampled (see Supplementary Table S1) and their alignments analyzed as described below. Plastid matK and psbE-petG sequences were concatenated and analyzed together.

Maximum parsimony (MP) and MP bootstrap (MPB) analyses for ITS and plastid datasets were performed using PAUP 4.0b10 (Swofford, 2002). The most parsimonious trees were searched for heuristically with 100 replicates of random sequence addition, TBR swapping, and MulTrees off. The swapping was performed on a maximum of 1000 trees (nchuck = 1000). All characters were equally weighted and unordered. The data set was bootstrapped using full heuristics, 1000 replicates, TBR branch swapping, MulTrees option off, and random addition sequence with five replicates. Silene acaulis L. and S. menziesii Hooker were used for rooting in the plastid and ITS datasets, respectively, based on Popp and Oxelman (2007) and preliminary analyses described above.

Bayesian analyses were performed with MrBayes 3.2.1 (Ronquist et al., 2012) applying the SYM+Γ (ITS) and GTR+Γ (plastid dataset) substitution models proposed by the Akaike information criterion implemented in MrAIC.pl 1.4 (Nylander, 2004); matK and psbE-petG alignments were analysed separately with MrAIC but for both the same substitution model was proposed. Values for all parameters, such as the shape of the gamma distribution, were estimated during MrBayes analyses. The settings for the Metropolis-coupled Markov chain Monte Carlo process included four runs with four chains each (three heated ones using the default heating scheme), run simultaneously for 10,000,000 generations each, sampling trees every 1,000th generation using default priors. The posterior probability (PP) of the phylogeny and its branches was determined from the combined set of trees, discarding the first 1001 trees of each run as burn-in.

ITS and plastid datasets were pruned for divergence times estimation. Several distantly related outgroup taxa and some accessions of closely related species for which several accessions were available were removed. The resulting matrices thus included the same species. The analyses were performed using BEAST ver. 1.8.2 (Drummond et al., 2012). Prior to analyses, we excluded all longer (>5 bp) autapomorphic insertions from the plastid alignment. Birth-death speciation prior with incomplete sampling (Stanler, 2009) and GTR+Γ substitution model with estimated base frequencies were used for phylogeny inference. A lognormal relaxed clock with a weakly informative prior on the clock rate (exponential with mean 0.001) was applied. The prior age of the root was set to 3.37 million years with a normally distributed standard deviation of 0.9, which corresponds to the median age and 95% highest posterior densities (HPD) interval of the corresponding node obtained from the dating analysis by Sloan et al. (2009). In that study South American S. argentina (Pax) Bocquet was positioned in a polytomy with other species of S. sect. Physolychnis and the divergence of the clade was dated to 3.37 Ma (HPD, 1.69–5.21 Ma). The analyses were run for 10 million generations, logging parameters every 1000 generations. The performance of the analyses was checked in Tracer 1.6.0 (Rambaut et al., 2014); both the effective sample sizes and mixing were appropriate (i.e., ESS always exceeded 100 and the parameter traces showed the chain fluctuating around the equilibrium). The maximum clade credibility trees (MCC) showing the mean ages were produced and annotated with Tree Annotator (part of the BEAST package) after removing burnin and visualized with FigTree 1.4.2 (Rambaut, 2014).

**Leaf Shape Analyses**

To compare the leaf shapes among populations of different species we plotted the scanned leaf shapes of the basal leaves on Figure 1. Additionally, we measured (1) the leaf length, (2) the maximal leaf width as well as (3) the distance from the leaf basis to the point of maximal leaf width for all sampled populations of subsect. Chilenses. We also calculated the ratios between 1 and 2, and 3 and 1, to describe the leaf shape.

**RESULTS**

**Chromosome Numbers and Relative Genome Size**

Chromosome numbers were established for 12 populations representing six Silene species (Supplementary Table S1 and Figure 2). The basic chromosome number of all individuals was inferred to be x = 12. Two ploidy levels were recorded, i.e., tetraploid (2n = 4x = 48) and decaploid (2n = 10x = 120). Each species, regardless of the number of individuals analyzed, represented only one ploidy level. Four species were tetraploid [S. antarctica, S. chubutensis (Spegazz.) Bocquet, S. magellanica, S. patagonica] and two decaploid, although establishing unambiguously exact chromosome numbers for those two individuals was no trivial task [S. chilensis (Naudin) Bocquet, S. echegarayi (Hieron.) Bocquet]. The karyotypes cannot be directly compared without additional markers, as it is impossible to define homo- or homoeologous chromosome pairs based on chromosome morphology alone, especially given their very similar sizes and types (mostly metacentrics and submetacentrics with some acrocentrics). The average sizes of the chromosomes were similar within individual species, but differed among the species with S. magellanica having chromosomes larger than other analyzed taxa (Figure 2).
According to RGS the inference of ploidy-levels was not clear-cut (Figure 3), but available chromosome counts and RGS for some populations allowed the inference of DNA-tetraploidy in populations for which chromosome numbers were not estimated, at least for two populations of S. chubutensis, for which one count from a third population was available and for S. densifolia, which had the RGS very similar to that of S. chubutensis (ID 20), for which the chromosome number was established (Supplementary Table S1). The RGS of tetraploids ranged from 1.47 in S. cuspidata (ID 9) to 2.40 in S. antarctica (25), and in decaploids from 2.78 to 3.24, thus exhibiting 2.2-fold variation; both latter values are from S. chilensis (12, 7). Also among different populations of some species the RGS varied considerably (Supplementary Table S1 and Figure 3).

Plastid and ITS Sequences

Internal transcribed spacer sequences of South American accessions were between 746 (Silene sp. 22) and 748 (S. chilensis 7, 10, 12) bp long. Topologies inferred by parsimony and Bayesian analyses were congruent, as no strong incongruences (different strongly supported clades with PP > 0.95 and/or BS > 70%) were detected. Parsimony analysis inferred a lower number of clades, which mostly had lower support compared to Bayesian inference (Figure 4). South American Silene species were positioned in two monophyletic, but clearly divergent clades. One consisted of S. chilensis and S. echegarayi (posterior probability, PP 1, maximum parsimony bootstrap, BS 98%) and was positioned in a clade (PP 1, BS 58%) with S. suksdorfii B. L. Robinson, S. uralensis (Rupr.) Bocquet and S. violascens (Tolm.) V. V. Petrovsky & Elven. All other South American species were in a clade (PP 1, BS 85%) in trichotomy (PP 1, BS 70%) with North American S. occidentalis and S. verecunda.

All matK sequences of South American accessions were 869 bp long, those of psbE–petG varied in length between 1347 bp (S. densifolia (Dusen) Bocquet 21) and 1487 bp (S. chilensis 7). Topologies inferred by parsimony and Bayesian analyses of the concatenated plastid dataset were largely congruent, as no strong incongruences (different strongly supported clades with PP > 0.95 and/or BS > 70%) were detected. Parsimony analysis inferred a lower number of clades, which mostly had
lower support compared to Bayesian inference (Figure 5). South American Silene species formed a monophyletic group (P 0.99) sister to two accessions of North American S. verecunda (PP 0.97). All other outgroup species formed several clades with different support values. One of them, clearly separated from South American accessions, consisted of S. sachalinensis F. Schmidt, S. scouleri, S. sorensenis (B. Boivin) Bocquet, S. suksdorfii and S. uralensis (PP 1, BS 75%).

The relationships in phylogenetic trees inferred by Bayesian analyses using BEAST were largely unresolved, resulting in a basal polytomy (Figure 6), in accordance with the results described above (Figures 4, 5). Only some clades had a support exceeding PP 0.95. In the ITS tree (Figure 6A) one of these clades included most of the South American accessions (PP 1) as sister (PP 1) to one accession of S. verecunda. The other such clade included S. chilensis and S. echegarayi (PP 1) as sister (PP 1) to S. uralensis and S. suksdorfii. In the plastid tree (Figure 6B) all South American species were in a clade (PP 1) as sister (PP 1) to one accession of S. verecunda, whereas S. uralensis and S. suksdorfii were in a clade (pp 0.97) with S. nigrescens and some other species. The divergence of South American species from North American S. verecunda was dated to the late Pliocene to early Pleistocene, 2.45 Ma (HPD, 1.25–3.76 Ma) in the ITS tree and to 2.05 Ma (HPD, 0.85–3.35 Ma) in the plastid tree. Silene chilensis and S. echegarayi diverged from the S. uralensis alliance 1.89 Ma (HPD, 0.82–3.07 Ma).

Leaf Shape Variation
As shown in Figure 1 leaf shape differs among high- and low-elevation populations. Within S. subsect. Chilenses the high-elevation taxa have broader, spatulate to oblanceolate leaves, which are widest in the upper half, whereas the low-elevation species have narrower, usually linear leaves, which are normally widest at the base (Supplementary Table S1 and Figures 1, 7).

DISCUSSION
Single Migration From North to South America in Silene Sect. Physolychnis
Phylogenies inferred from nuclear ITS and plastid DNA sequences (Figures 4–6) suggest a single origin of South American species of S. sect. Physolychnis, followed by allopolyploidisation in the S. chilensis lineage. The datasets indicate that South America was colonized by a lineage sister to North American S. verecunda and S. occidentalis. Silene verecunda is a tetraploid, morphologically and ecologically exceptionally variable species occurring in the western
United States from 0 to 3400 m (Morton, 2005). Morphologically it is fairly similar to the South American species, even if many species of *Silene* sect. *Physolychnis* resemble in habit. In contrast, *S. occidentalis*, which is endemic to California, differs considerably in morphology from South American species and *S. verecunda*, especially in having a remarkably prolonged tubular calyx and deeply lobed pink petals (Morton, 2005). Many Northern American temperate plant lineages have colonized the South American Andes after overland connections were established between the two continents, but the timing of migrations differed among the groups (Bell and Donoghue, 2005; Morton, 2005).
in the plastid tree (Figure 6). The migration thus likely happened in the late Pliocene or early Pleistocene, i.e., after the final closure of the Isthmus of Panama and the final uplift of the northern Andes (Gregory-Wodzicki, 2000). This is in line with previous evidence that the upland Andean habitats have been available for plant colonization only since the late Pliocene and early Pleistocene 2 to 4 Ma and several plant groups, including Silene sect. Physolychnis, diversified in this area in the Pleistocene (Burnham and Graham, 1999; Gregory-Wodzicki, 2000; Hughes and Eastwood, 2006; Popp et al., 2011; Madriñán et al., 2013; Hughes and Atchison, 2015). In several genera like Astragalus (Fabaceae; Scherson et al., 2008), Calceolaria (Calceolariaeae; Cosacov et al., 2009) Gentianella (Gentianaceae; Von Hagen and Kaderieit, 2001), Hypericum (Hypericaceae; Nürk et al., 2014), Lupinus (Fabaceae; Hughes and Eastwood, 2006), and Valeriana (Valerianaceae; Bell and Donoghue, 2005) rapid diversifications leading to high numbers of species in the Andes have been evidenced, but in Silene sect. Physolychnis the number of species remained relatively low.

**Allopolyploidisation and Origin of Decaploid Species**

Most of the South American species analyzed here are tetraploid (Figures 2, 3 and Supplementary Table S1), similar to their North American sister taxon S. verecunda (Morton, 2005). A decaploid chromosome number (2n = 10x = 120) established for the high Andean species S. chilensis and S. echegarayi (Figure 2 and Supplementary Table S1) is the highest ploidy level ever reported in Silene sect. Physolychnis, since only octoploids were previously known (Popp and Oxelman, 2007; Petri and Oxelman, 2011; Rice et al., 2015); up to 240 chromosomes have been reported for European S. ciliata Pourr. (Blackburn, 1933). The diploid chromosome count for S. magellanica by Moore (1981) is probably erroneous, as no diploids were established by our chromosome counts neither for this nor any other South American species belonging to S. sect. Physolychnis.

A discordant position of decaploid South American species in ITS and plastid trees (Figures 4–6) suggests their allopolyploid origin. Decaploid species are positioned in the same clade with tetraploid South American species in the plastid tree (Figure 5), where most accessions (with exception of population 16 of S. chilensis) appear in the same clade with morphologically very similar and sympatric S. andicola; thus, the latter likely served as a maternal parent in the allopolyploid origin of the decaploids. On the other hand, in the ITS tree decaploids are positioned in a clade with the diploid arctic/subarctic S. uralensis, the North American octoploid S. suksdorffii and the North Asian S. violascens. Nowadays none of these or closely related species occur in South America and it remains unclear how the hybridisation has happened. In any case, our dating analyses indicate a time lag between the migration to South America and the putative hybridisation with the S. uralensis lineage resulting in the allopolyploidisation as the divergence of decaploids from the S. uralensis lineage was dated to 1.89 Ma (HPD, 0.82–3.07 Ma).

Previous research has shown that diploid S. uralensis was involved also in the allopolploid origin of tetraploid subarctic/arctic...
S. involucrata (Cham. & Schltdl.) Bocquet, but as a maternal lineage (Popp et al., 2005). In this case the distribution range of the paternal Siberian/northeastern Asian S. ajanensis (Regel.) Vorosch. lineage nowadays overlaps with that of S. involucrata (Popp et al., 2005). In combination with plastid DNA sequences, ITS has often been used to infer origins of polyploid plants (Sang et al., 1995; Ge et al., 1999; Popp et al., 2005; Popp and Oxelman, 2007; Kuzmanović et al., 2017). However, since the homoeologous ITS repeats in an allopolyploid often are homogenized toward one of the parental types by concerted evolution or chimeric sequences are formed (Wendel et al., 1995), the inference of allopolyploid origin is only possible if the homogenization is directed toward the paternal lineage (Smedmark and Eriksson, 2002; Popp et al., 2005), which was also the case in South American Silene sect. Physolychnis. The homogenisation seems to be complete, as we observed only two polymorphic sites in all ITS accessions of decaploid Silene.

**Morphological Differentiation in South American Silene Sect. Physolychnis**

The South American species of Silene sect. Physolychnis exhibit a great variability in habitats as well as in morphology. Most of the species occur at higher altitudes of the Andes, the Andean Precordillera and other mountain areas in Argentina (e.g., Sierra de Córdoba, Sierra de la Ventana), but some inhabit also Patagonian dry steppes and semi-deserts (Bocquet, 1969; Pedersen, 1984). Based on the calyx characters, Bocquet (1969) divided South American species into three subsections. The first, S. subsect. Genovevanae Bocquet includes five species from the northern Andes from Venezuela to northern Argentina. From this subsection we included S. mandonii (Rohrb.) Bocquet and S. thyssanodes Fenzl in our phylogenetic analyses; in the ITS tree S. thysanodes S. mandonii northern Andes from Venezuela to northern Argentina. From S. in our analyses.

**CONCLUSION**

Our study provides new insights into origin and diversification of Silene sect. Physolychnis from South America. The analyzed nuclear ITS and plastid sequences indicate a single migration of the S. verecunda lineage from North to South America and subsequent allopolyploidisation with the S. uralensis lineage. To corroborate this hypothesis further phylogenetic studies including DNA sequences from additional nuclear and plastid regions are needed. Moreover, sampling of South American species not included in our study is needed to corroborate a single origin of all South American members of this section. Finally, low phylogenetic resolution within different species groups and high intraspecific morphological variability calls for a taxonomic revision of South American Silene sect. Physolychnis.

**AUTHOR CONTRIBUTIONS**

BF designed the study, coordinated its implementation, conducted field work, performed karyological and phylogenetic analyses, and wrote major parts of the paper. PS conducted field work and improved a draft of the paper. HW-S coordinated and conducted karyological analyses, wrote corresponding parts of the paper and improved its draft. BO was involved in study design, performed the phylogenetic analysis of the complete Sileneae datasets, coordinated the sequencing of most of the sequences, and improved a draft of the paper.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2018.00639/full#supplementary-material
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