A taxonomic and evolutionary review of the South American *Hierochloë* section *Monoecia* (Poaceae: Anthoxanthinae)

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An integrative approach combining morphological, molecular and cytological information was used to assess the taxonomy and biogeography of *Hierochloë* section *Monoecia*. More specifically, we aimed to evaluate (1) if the morphological and molecular data are in concert with the current taxonomy of the group and (2) if speciation in this exclusively South American group could be linked to the formation of the Andes. Our analysis of 31 macro- and micromorphological characters, four plastid and nuclear DNA regions, and nuclear DNA content suggests that the taxonomic status of several of the species in the section is not justified based on either the phylogenetic (apomorphic) or the phenetic species concepts. We propose that only four out of the eight species in the section (*H. pusilla*, *H. juncifolia*, *H. quebrada* and *H. redolens*) should be recognized and the remaining taxa (*H. altissima*, *H. gunckelii*, *H. spicata* and *H. utriculata*) should be reduced to varieties of the widespread *H. redolens*. In addition, we recover a biogeographical scenario for section *Monoecia* including genetic exchange between the southern and the central Andes, recent and incomplete diversification in the southern Andes, and longer isolation history for those species in the section with restricted ecological and/or geographical ranges.

 $ADDITIONAL\ KEYWORDS:\ biogeography-molecular\ phylogenetic\ analysis-morphometric\ analysis-nuclear\ DNA\ content-South\ America-taxonomy.$

INTRODUCTION

All South American *Hierochloë* spp. have monoecious spikelets, which are unique in the genus. This trait has led different authors to recognize their close relationships (Parodi, 1941; De Paula, 1975; Connor & Renvoize, 2009) and to propose their inclusion in section *Monoecia* Connor (Connor, 2012). Here, we use micro- and macromorphological, molecular and karyological data to clarify the taxonomy in the section and to discuss the biogeography and evolution of *Hierochloë* in the southern New World.

be maintained ... because of their distinctive floral

Floral structure is key to the taxonomy of the closely related genera *Hierochloë* and *Anthoxanthum*

L. (e.g. Connor, 2012; Pimentel et al., 2013), the only two components of subtribe Anthoxanthinae (Poaceae). They have laterally compressed spikelets bearing three florets, the two lower being male or neuter in Anthoxanthum and invariably tristaminate in Hierochloë, whereas the apical floret is usually bisexual, bistaminate and protogynous in both genera. Schouten & Veldkamp (1985) proposed merging the two genera based on the existence of intermediate forms, the putatively hybrid Anthoxanthum section Ataxia R.Br (Pimentel et al., 2013). However, Connor (2012) indicated that 'both genera should

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biology and of their separate pathways to individual floral expression'. Andromonoecism (i.e. perfect apical floret and male lower florets) is dominant in boreal New World, Old World and Australasian Hierochloë (Weimarck, 1971; Connor, 2008), with only the South American taxa (section *Monoecia*) showing monoecism (i.e. the apical floret is functionally female, whereas the lower florets are male; Parodi, 1941). The number of species comprising section Monoecia is difficult to establish because no recent taxonomic review is available (Parodi, 1941, De Paula, 1975). In his study on floral biology, Connor (2012) included seven species in section Monoecia: H. juncifolia (Hack.) Parodi, H. quebrada Connor & Renvoize, H. utriculata (Ruiz & Pav.) Kunth, H. altissima Steud., H. redolens (Vahl) Roem. & Schult., H. gunckelii Parodi and H. pusilla Hack. All these species apart from the recently described H. quebrada from the central Andes of Peru (Connor & Renvoize, 2009) were also recognized by De Paula (1975) and Parodi (1941) in their taxonomic reviews of the genus in Chile and Argentina and South America, respectively. Parodi (1941) included one additional taxon in his study, H. spicata Parodi, a species also accepted by De Paula (1975) who described it as endemic to southern Chile. In her review, De Paula (1975) added two more taxa, H. sorianoi De Paula and H. moorei De Paula, currently considered extreme forms of the widespread H. redolens (e.g. Anton & Zuloaga, 2012; but see Villalobos & Finot, 2016).

Taxa in the exclusively South American Hierochloë section Monoecia grow from Venezuela to Tierra del Fuego (De Paula, 1975; Connor, 2008; Anton & Zuloaga, 2012) and are restricted to temperate to cold environments in the Andes, defined in this paper following Nagy & Grabherr (2009) and Körner, Paulsen & Spehn (2011). An inverse relationship between elevation and latitude common to most C3 grass groups (Still et al., 2003) can be observed in the distribution of the section. The central Andean H. quebrada from Peru is restricted to tropical alpine areas > 4000 m a.s.l. (Connor & Renvoize, 2009), whereas H. redolens presents a disjunct distribution, growing in tropical alpine areas from Venezuela to Peru (up to 3850 m a.s.l.) and in moist areas in southern South America (central to southern Chile and Argentina) at lower elevations. Some doubt has been cast on the identity of the plants from both regions (e.g. Parodi, 1941; Connor, 2012). This taxon has also been recorded in New Zealand, Australia and New Guinea, but populations from these areas show fundamental differences in floral constitution and development and should be considered as representing a different species (Zotov, 1973; De Paula, 1975; Connor, 2012). Hierochloë utriculata, H. altissima, H. spicata and H. gunckelii show overlapping ranges at different latitudes in

central to southern South America, on one or both sides of the Andes (De Paula, 1975), whereas *H. juncifolia* grows in volcanic or sandy soils in mountains up to 1750 m a.s.l. in central Chile and central-western Argentina, showing a discontinuous distribution. Finally, *H. pusilla* is restricted to moist areas between 100 and 1000 m a.s.l. at high latitudes in Argentina (Santa Cruz and Tierra del Fuego provinces) and Chile (Magallanes Region; De Paula, 1975).

Morphological differentiation in section *Monoecia* is complicated, casting doubts on the taxonomic status of some of its members (Villalobos & Finot, 2016). Hierochloë juncifolia and H. pusilla are the only species showing a clear-cut differentiation, based on awn insertion in the male floret and leaf shape and on plant size and lack of awned male florets, respectively (De Paula, 1975). For the remaining taxa, artificial limits in inflorescence shape, an otherwise continuous trait, are used to differentiate H. altissima, H. quebrada, H. spicata and H. utriculata (spiciform panicles) and H. gunckelii and H. redolens (lax to somewhat contracted panicles). The use of micromorphological traits for species differentiation is promising (Connor, 2008; Villalobos & Finot, 2016); however, more complete analyses including more individuals and populations are needed to assess their usefulness.

Hierochloë (basic chromosome number, x = 7) also displays a wide variation in ploidy, with diploids, tetraploids, hexaploids and duodecaploids having been identified (e.g. Weimarck, 1971). Still, to the best of our knowledge, chromosome numbers or genome size estimations are unknown in section *Monoecia*. These cytogenetic characters have proven to be essential in the clarification of taxonomic problems in reticulated groups with high variation in chromosome numbers, a feature common to many sections in Anthoxanthinae (Chumová *et al.*, 2015).

Overlapping morphologies and distribution ranges associated with climatic conditions highlight the need for a reappraisal of the taxonomy of *Hierochloë* section *Monoecia*, a possible model for diversification of C3 taxa in the Andes due to its distribution encompassing the whole mountain range. Combined analyses using data from different sources have been deemed especially useful in solving taxonomic problems (Ruhfel, Stevens & Davis, 2013; Besse, 2014; Szlachetko et al., 2017). Here, we follow this approach to clarify the taxonomy of the section and to discuss the evolution and biogeography of this group of South American grasses. First, nuclear DNA content values were assessed for the different species in the section using flow cytometry (FCM) (e.g. Doležel, Greillhuber & Suda, 2007). Second, we conducted multivariate analyses of macro- and micromorphological data to assess the boundaries among taxa and the reliability

of the traits traditionally used in the taxonomy of section *Monoecia*. This methodology has been successfully applied to other complex groups of plants (e.g. Repka, 2003; Ospina, Sylvester & Sylvester, 2016), including those in Anthoxanthinae (Pimentel, Estévez & Sahuquillo, 2007; Pimentel, Catalán & Sahuquillo, 2010). Thirdly, we sequenced several commonly used plastid and ribosomal nuclear DNA markers (e.g. Pimentel et al., 2013) to determine whether the taxonomy of the section is in concert with the phylogenetic tree. Given the multi-copy nature of the nuclear regions used, a cloning strategy was devised to detect possible instances of reticulate evolution (Díaz-Pérez et al., 2014; Wan et al., 2014). Molecular data were also analysed in the light of what is known about the evolution and biogeography of Andean plants (reviewed by Luebert & Weigend, 2014).

MATERIAL AND METHODS

PLANT MATERIAL

Ninety-six field-collected or herbarium specimens were used in this study (Appendix 1); 94 plants were included in the morphometric analyses and subsets of specimens were considered in the anatomical (66 samples), molecular (39 samples, 31 used for cloning) and FCM (83 samples) analyses. In *Hierochloë* section

Monoecia, four to 22 specimens were sampled per species (Appendix 1), with the exception of H. quebrada and H. spicata, for which only one plant was included due to the difficulty in differentiating them in the field and their poor representation in the analysed herbaria. Sixty-seven of the analysed specimens were directly collected in the field (15 populations, five specimens per population on average; for population details see Appendix 1). In each population we collected specimens in anthesis that were at least 3 m apart to avoid sampling clones. Collected plants were pressed, leaves were taken and preserved in silica gel for DNA extraction and, when available, seeds were stored for cytometry studies. The selection of localities was based on De Paula (1975) and vouchers were deposited in SANT and CONC (Holmgren, Holmgren & Barnett, 1990). For the geographical location of populations see Figure 1. Twenty-nine herbarium specimens were added to the database in order to (1) complete the representation of section Monoecia and (2) include samples of two of the three other exclusively American Anthoxanthum or Hierochloë spp. outside Monoecia, i.e. Anthoxanthum mexicanum Mez and Hierochloë occidentalis Buckley, and populations of H. redolens from South-East Asia. All plants used in the study were tentatively identified (or, for herbarium samples, their identity was confirmed) following De Paula (1975). Herbarium specimens were obtained from

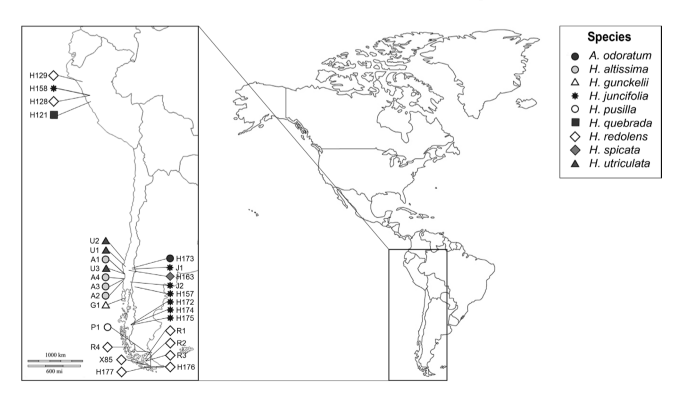


Figure 1. Sampled populations of $Hierochlo\ddot{e}$ section Monoecia and Anthoxanthum odoratum in South America (population names follow Appendix 1).

CONC, K, M, MA, PH, SANT, UPS and US (Holmgren et al., 1990).

GENOME SIZE ESTIMATION BY FLOW CYTOMETRY

Nuclear DNA content was estimated through FCM following Galbraith *et al.* (1983) and Loureiro *et al.* (2007). The obtained values were expressed in picograms (pg) and in mega base pairs (Mbp) using the formula of Doležel *et al.* (2003) (1 pg = 978 Mbp). At least five individuals were assessed per species. A univariate analysis of variance (ANOVA) and a Tukey's test were used to characterize the groups with similar measures of nuclear DNA content (2C values; *sensu* Greilhuber *et al.*, 2005). For a full description of the procedure see Supporting Information Appendix S1.

MACRO- AND MICROMORPHOLOGICAL ANALYSES

Fifteen macromorphological (eight quantitative and seven qualitative) and 16 micromorphological (three quantitative and 13 qualitative; 11 from the leaf epidermis and five from the leaf transverse section) characters were measured. The characters were selected because (1) they are commonly used in the taxonomy of Anthoxanthinae and (2) we observed a high variability among the samples included in the analyses. To standardize data collection, leaf and spikelet data were gathered in the second leaf of the plant from the base and the basal spikelet, respectively. Micromorphological data were obtained following Devesa (1992) and Pimentel & Sahuquillo (2003), with minor modifications. All quantitative macromorphological traits and one quantitative micromorphological character (long cells length; LlcL) were logarithmically transformed to limit the influence of allometry on the results (Dufrene, Gathoye & Tyteca, 1991; Almeida-Pinheiro de Carvalho et al., 2004) and quantitative micromorphological data were obtained by calculating the mean value of 12 measurements. Qualitative characters were scored as binary variables (presence/absence). Those qualitative characters that presented more than two states (three macro- and eight micromorphological) were transformed into binary traits. All macro- and micromorphological characters used are listed in Appendix 2.

Each specimen measured was treated as an independent operational taxonomic unit (OTU) for all statistical tests. Macro- and micromorphological databases were built and analysed separately due to the different number of samples, but the same statistical techniques were used. Qualitative and quantitative data were analysed jointly and separately. All statistical analyses were performed using IBM SPSS Statistics for Mac v. 20.0 (IBM Corp., 2011) and

R v. 3.3.2 (R Core Team, 2016a) as implemented in RStudio v. 1.0.136 (RStudio Team, 2016) and Rcmdr v. 2.3-1 (Fox, 2005, 2017; Fox & Bouchet-Valat, 2016). The packages 'RcmdrMisc' (Fox, 2016) and 'foreign' (R Core Team, 2016b) were used to translate our Exceland SPSS-built databases into an R-readable format. Plots were built using the packages 'lattice' (Sarkar, 2008) and 'rgl' (Adler & Murdoch, 2016). A detailed description of the methods applied can be found in Appendix S2.

DNA ISOLATION, AMPLIFICATION, CLONING AND SEQUENCING

DNA from silica-gel dried, field-collected leaves and herbarium samples was extracted using the NORGEN Plant/Fungi DNA isolation kit (Norgen Biotek Corporation, Thorold, Ontario, Canada) and the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany), respectively. We followed the manufacturer's protocols, with modifications in the latter case following Bendiksby (2011).

The selected plastid and nuclear regions were chosen based on results from Pimentel et al. (2013) and Tusiime et al. (2017). The plastid trnT-L and trnL-F (including the trnL intron plus the trnL-F intergenic spacer; Taberlet et al., 1991) regions were amplified and sequenced using primers a and b (trnT-L intergenic spacer; Taberlet et al., 1991) and primers c and f. Amplification of the *trnT-L* region followed Galley & Linder (2007), whereas for the trnL-F PCR conditions followed Torrecilla et al. (2003). Amplification of the ribosomal nuclear ITS (ITS1-5.8S-ITS2) and ETS regions followed Hsiao et al. (1995) and Gillespie, Soreng & Jacobs (2009), respectively. Although these regions have produced useful results when comparing closely related taxa (e.g. Consaul. Gillespie & Waterway, 2010; Pimentel et al., 2013), homology issues connected to concerted evolution have been raised (Alvarez & Wendel, 2003). Considering the above, and in view of multiple bands being registered for the ETS of several specimens, this region was cloned in 31 plants belonging to all species in section Monoecia plus A. mexicanum and South-East Asian H. redolens populations (cloning failed for the northern American *H. occidentalis*). Two plants were cloned per population, except for *H. altissima* population A2, H. pusilla population P1 and all herbarium samples (Appendix 1), for which only one sample was available. When cloning did not produce enough colonies, one additional plant was added (*H. altissima* population A1 and *H. juncifolia* population J2). Six to 19 clones were sequenced for each plant (Appendix 1). Clones were analysed separately and were only added to the general DNA matrix when direct sequencing of non-cloned ETS amplicons failed. Cloning and plasmid extractions were performed using the StrataClone PCR cloning kit (Agilent Technologies, Santa Clara, CA, USA) and the QIAprep kit (Qiagen), respectively, following the manufacturers' protocols. An ETS sequence from *Anthoxanthum odoratum* L. was used as an outgroup (Appendix 1).

Non-cloned products were purified using ExoSap-IT PCR cleanup reagent (Affymetrix, Santa Clara, CA, USA), and all products were sequenced using the BigDye Terminator Cycle Sequencing Ready Reaction v3.1 kit (Applied Biosystems, Paisley, UK) on an Applied Biosystems 3710 automated sequencer at the Universidade da Coruña Sequencing Service. Amplification primers were used for all sequencing except for the clones, for which the M13/T7 primer pair was used. Conservative approaches are recommended to reduce the impact of PCR artefacts on the phylogenetic reconstruction based on clones (Popp & Oxelman, 2007). Here, we followed a mixed approach based on Popp & Oxelman (2007) and Díaz-Pérez et al. (2014). A maximum parsimony (MP) analysis based only on potentially parsimony-informative characters (see Sequence alignment and phylogenetic analyses below) was run to detect putative chimeric sequences (long terminal branches due to homoplasy; Popp & Oxelman, 2007). These sequences were excluded from all subsequent analyses. Monophyletic groups of clones differing only by autopomorphic substitutions were considered as a single sequence (Popp & Oxelman, 2007). For the remaining clones, the position of which in the parsimony tree was in a polytomy, we followed Díaz-Pérez et al. (2014). A p-distance matrix was built using MEGA v. 6.0 (Tamura et al., 2013). All sequences with a *p*-distance < 0.01 base substitutions *per site* were collapsed into a single type. Consensus sequences were built using SeaView v. 4 (Gouy, Guindon & Gascuel, 2010).

SEQUENCE ALIGNMENT AND PHYLOGENETIC ANALYSIS

The forward and reverse sequences were edited and assembled using CodonCode Aligner v. 4.0 (CodonCode Corp., Dedham, MA, USA). DNA regions were independently aligned using the MUSCLE algorithm (Edgar, 2004) as implemented in SeaView v. 4 (Gouy et al., 2010) and manually adjusted upon the detection of errors. For this study, 539 new sequences were generated: 409 ETS (363 clones), 41 ITS, 48 trnL-F and 41 trnT-L (Appendix 1). Independent databases were built for each region (plus a fifth database including all clones). Plastid and nuclear sequences from four Eurasian Hierochloë spp. were also obtained and added to the different databases to clarify the phylogenetic position of section Monoecia

in the genus. Two additional sequences belonging to the closely related but separate *Anthoxanthum* section *Anthoxanthum* were also added and used as outgroups (Appendix 1).

Separate plastid, nuclear and nuclear-clone matrices were built and analysed independently. Sequences missing due to PCR and/or sequencing problems were coded as missing data in the concatenated data sets. Plastid and nuclear datasets were also concatenated and jointly analysed once conflicting samples were removed (see Results). MP or Bayesian inference (BI) analyses were run depending on the data set; MP was used to detect putative chimaeric sequences in the clones database, whereas BI was applied to the inference of phylogenetic relationships in all matrices (plastid, nuclear, combined and clones). MP and Bayesian analyses followed Torrecilla et al. (2003) and Pimentel et al. (2013), and are described in Supporting Information Appendix S3. Gaps were treated as missing data in all analyses, except in the combined plastid + nuclear tests, for which all gapped positions were excluded.

PHYLOGENETIC NETWORKS

Phylogenetic networks were computed to represent groupings in the data and evolutionary distances among taxa simultaneously (López-Pujol et al., 2012). Statistical parsimony analyses as implemented in the software TCS v. 1.21 (Clement et al., 2002) and PopArt v. 1.7 (http://popart.otago.ac.nz) were applied to the plastid, nuclear and ETS clone databases. Gaps were ignored and the remaining parameters were set by default. This method was chosen due to its simplicity of representation, because it allows for the detection of haplotypes that are candidates for being products of recombination (Templeton, Crandall & Sing, 1992; Posada & Crandall, 2001) and because it is useful in the definition of species under the phylogenetic species concept (Hart & Sunday, 2007).

RESULTS

GENOME SIZE ESTIMATIONS

Holoploid genome sizes (2C) ranged from $2C = 12.82 \,\mathrm{pg}$ in H. pusilla to $2C = 27.43 \,\mathrm{pg}$ in H. altissima and are listed in Table 1 and Supporting Information Figure S1. The coefficient of variation (CV) of genome size values was < 3% for all populations represented by more than one specimen except for one H. utriculata population (U1; CV = 4.77%; for population codes see Appendix 1). Hierochloë altissima is the only species showing significant genome size variation across its populations.

Table 1. Nuclear DNA content in the South American species of Hierochloë section Monoecia analysed in this study

Species	Pop.	Genome s	ize (2C, pg)				G^*	n. spec.
		Mean	SD	Min.	Max.	CV (%)		
H. altissima	A1	18.58	0.36	18.19	19.08	1.94	G3	5
H. altissima	A2	27.43	0.37	27.02	27.91	1.34	G5	5
H. altissima	A3	27.04	0.40	26.65	27.58	1.48	G5	5
H. altissima	A4	27.30	0.25	27.04	27.55	0.90	G5	5
H. gunkelii	G1	19.43	0.32	19.06	19.84	1.64	G4	5
H. juncifolia	J1	14.75	0.24	14.36	14.98	1.66	G2	5
H. juncifolia	J1	14.09	0.21	13.86	14.38	1.47	G2	5
A. mexicanum	M1	18.89	_	_	_	_		1
A. mexicanum	M2	17.62	_	_	_	_		1
$H.\ occidentalis$	O1	21.84	_	_	_	_		1
H. pusilla	P1	12.82	0.21	12.63	13.12	1.61	G1	5
H. quebrada	Q1	18.87	_	_	_	_		1
H. redolens	R1	19.65	0.25	19.37	19.94	1.26	G4	5
H. redolens	R2	19.46	0.42	18.86	19.99	2.17	G4	5
H. redolens	R3	19.84	0.58	18.76	20.42	2.92	G4	6
H. redolens	R4	19.02	0.51	18.34	19.60	2.66	G3	5
H. spicata	S1	13.28	_	_	_	_		1
H. utriculata	U1	18.54	0.88	16.89	19.34	4.77	G3	6
H. utriculata	U2	18.27	0.26	17.90	18.47	1.45	G3	5
H. utriculata	U3	18.46	0.27	18.10	18.84	1.47	G3	5

^{*}See Discussion in this paper. Pop., population; SD, standard deviation; Min., minimum value; Max., maximum value; CV, coefficient of variation; G, DNA content group; n. spec., number of specimens measured. For population codes see Appendix 1.

According to the ANOVA and Tukey's test conducted, there are five groups based on 2C nuclear DNA content (Table 1; Fig. S1). Groups G1 (H. pusilla, 2C = 12.63–13.12 pg), G2 (H. juncifolia, 2C = 13.86–14.98 pg) and G5 (H. altissima populations A2, A3 and A4; 2C = 27.04–27.43 pg) are clearly differentiated, whereas groups G3 (H. altissima population A1, H. redolens population R4 and H. utriculata; 2C = 16.89–19.60 pg) and G4 (H. gunckelii and H. redolens populations R1, R2 and R3; 2C = 18.76–20.42 pg) are partially overlapping. Anthoxanthum mexicanum, H. occidentalis, H. quebrada and H. spicata were represented by only one sample and were excluded from the statistical tests.

MACROMORPHOLOGICAL ANALYSES: QUANTITATIVE CHARACTERS

Descriptive statistics (not shown) indicate that all characters overlap among most species with plant height (PH) and spikelet length (SL) showing the highest differentiation. Despite this, the ANOVA and Tukey's test revealed significant differentiation among species for all characters. The Kaiser-Meyer-Olkin test analysis performed indicated that our data were adequate for multivariate analyses (Fig. 2A, B). In the principal components analysis (PCA), the three first principal

components (C1, C2 and C3) accounted for 94.50% of the variability and the characters were grouped as in Table 2. Our accessions clustered into three groups: (1) *H. pusilla*, (2) *H. juncifolia* and (3) the other six species (Fig. 2B). Vegetative characters differentiated H. pusilla from the rest, whereas floral characters were responsible for the differentiation of H. juncifolia. A linear discriminant test (LDA) was performed based on the PCA results. Discriminant function 1 (F1) corresponded with the first component (C1; vegetative characters); F2 was composed of the three components of the PCA; and F3 corresponded to C2 and C3 (spikelet length, SL; flower length, FL; and lower glume length, LGL; all floral characters). The three functions are significant according to Wilks's lambda. The plot representing F1 and F3 (Fig. 2A) reveals a clear separation of *H. pusilla* and *H. juncifolia*, whereas the rest of the species are intermingled. Discriminant functions correctly classified 74.7 % of the specimens, although this percentage was different across species (not shown). The results of hierarchical cluster analysis (not shown) were consistent with the LDA.

MACROMORPHOLOGICAL ANALYSES: QUALITATIVE CHARACTERS AND COMBINED ANALYSES

The chi-square test of independence revealed that qualitative characters differed among species. The

Table 2. Matrix of rotated components (Varimax rotation) of the PCA performed using macromorphological quantitative data

	Component lo	adings	
Characters	Component 1	Component 2	Component 3
PH_ln	0.874	0.285	0.357
LL_ln	0.861	0.242	0.399
LW_ln	0.899	0.331	-0.017
IL_ln	0.863	0.322	0.349
$SLSB_{ln}$	0.826	0.378	0.300
SL_{ln}	0.283	0.387	0.860
FL_{ln}	0.418	0.838	0.250
LGL_{ln}	0.319	0.754	0.495
% variability	78.65	11.08	4.75

Morphological characters showing highest factor loadings on the first three principal components are in bold type. For character codes, see Appendix 2.

permutational multivariate analysis of variance using the distance matrix test (ADONIS; Supporting Information, Appendix S2) showed that the most similar species are H. gunkelii and H. utriculata (94.2% similarity; p-value = 0.942); all other speciesshowed a higher differentiation (< 30% similarity). The principal coordinate analysis (PCoA; not shown) and the non-metric multidimensional scaling test (NMDS; Fig. S2A) using the Jaccard index for three axes (stress = 0.07) separate the species into two groups: (1) H. altissima, H. gunckelii, H. redolens and H. utriculata; and (2) A. mexicanum, H. juncifolia, H. occidentalis and H. pusilla. The hierarchical cluster analysis (not shown) revealed the existence of two clear groupings in the data that did not follow the taxonomy of the section. Increasing the number of groups did not improve species discrimination, highlighting data dispersion. The taxa H. pusilla, H. redolens and, to a lesser extent, H. juncifolia had a less ambiguous adscription into one of the groups.

The NMDS (three axes, stress = 0.09; Fig. 3A) and the PCoA (not shown) performed with combined data using Gower's similarity coefficient clustered the species into three groups: (1) H. altissima, H. gunckelii, H. redolens and H. utriculata (highly overlapped); (2) A. mexicanum, H. juncifolia and H. occidentalis (slightly overlapped); and (3) H. pusilla (clearly differentiated). The multiple correspondence analysis (MCA; not shown) yielded the same results as the NMDS.

MICROMORPHOLOGICAL ANALYSES: QUANTITATIVE CHARACTERS

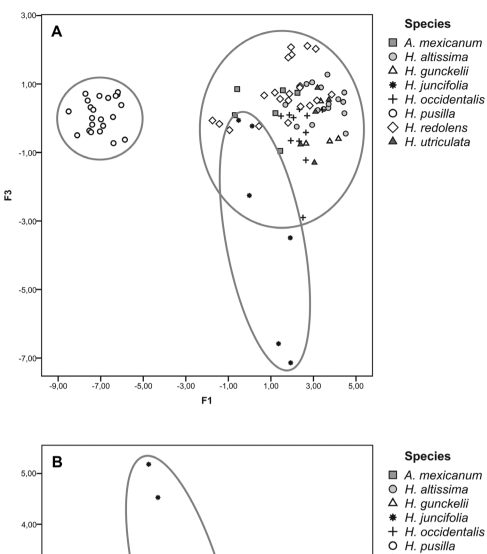
Descriptive statistics showed that all variables overlap among species (not shown; for character

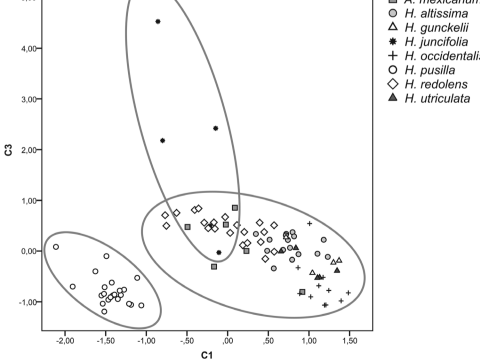
names see Appendix 2). The ANOVA indicates that stomata size (StS) does not differentiate among species (p-value = 0.176), whereas long cells length (LlcL) and number of ribs (NRi) showed significant differences across taxa. Only these variables (LlcL and NRi) were included in the discriminant functions obtained in the LDA using the stepwise method. The LDA plot (not shown) indicated that only H. juncifolia and H. pusilla are clearly different from the rest of the taxa. Hierochloë juncifolia shows low values for these two characters, whereas H. pusilla has low values for number of ribs (NRi) but intermediate values for long cells length (LlcL). The other five species are grouped together, especially H. altissima, H. gunckelii and H. occidentalis, showing higher values for number of ribs (NRi); H. redolens and H. utriculata presented intermediate values.

MICROMORPHOLOGICAL ANALYSES: QUALITATIVE CHARACTERS AND COMBINED ANALYSIS

The chi-square test of independence showed that only two traits, furrow depth (FD) and vascular bundles with sclerenchyma associated only in the margin of the leaf (PBEx) (for character names see Appendix 2), were not significantly different among taxa. The ADONIS analysis indicated that A. mexicanum, H. occidentalis and H. pusilla showed the highest consistency (100% internal similarity; p-value = 1.000) followed by H. redolens and H. utriculata (30.4% similarity; p-value = 0.304). The NMDS based on the Jaccard index for three axes (stress = 0.1, Supplementary Information Fig. S2B) separated the species into four groups: (1) H. altissima, H. gunckelii, H. pusilla, H. redolens and H. utriculata; (2) H. juncifolia; (3) A. mexicanum; and (4) H. occidentalis. The hierarchical cluster analysis (not shown) revealed low differentiating power for micromorphological qualitative data.

Qualitative and quantitative characteristics that did not have discriminating power according to the exploratory tests (StS, FD and PBEx; for character names see Appendix 2) were excluded from the combined analyses. The PCoA, MCA (not shown) and the NMDS plots performed using Gower's similarity coefficient for three axes (stress = 0.1; Fig. 3B) gave similar results and showed differentiation among the taxa. Consistent results were obtained in the hierarchical cluster analysis when the number of groups chosen equals the number of taxa. These analyses revealed that hook density (low density, HoF; intermediate density, HoN; high density, HoM), presence and shape of ribs (presence, RS; round-flattened, RSD; round, RSR; quadrangular, RSQ; polygonal, RSP) and leaf section shape (planar, LSP; open, LSO; closed, LSC) were the most differentiating traits.





PHYLOGENETIC RECONSTRUCTION

The independent analyses of the nuclear ribosomal DNA markers produced congruent topologies and the ITS and ETS regions were combined. Tree comparison indicated that the combination of the complete plastid and nuclear data was untenable due to topological conflict (Pirie *et al.*, 2009). A joint plastid + nuclear analysis was conducted after removing the only conflicting sample (*H. quebrada*).

The aligned plastid and nuclear matrices included 48 and 47 terminals, respectively (37 and 36 in section Monoecia). They represented 16 species (14 Hierochloë + one South American Anthoxanthum + one Eurasian Anthoxanthum, outgroup) and 1788 (plastid: trnL-F, 1-1084; trnT-L, 1085-1788) and 1288 (nuclear; ETS, 1–626; ITS, 627–1288) characters. The number of potentially parsimony-informative characters (as estimated using the program PAUP v. 410b; Sinauer Associates, Sunderland, MA, USA) was low, in particular 105 (5.9%) and 128 (9.9%) in the plastid and nuclear data, respectively. The Bayesian consensus plastid and nuclear trees are represented in Figures 4 and 5. The topologies were largely congruent (better general resolution in the plastid tree) and only the different position of H. quebrada was supported. The results of the concatenated analysis (H. quebrada removed from the matrices) are presented in Supporting Information Figure S3. In all trees, A. mexicanum is clearly differentiated from Hierochloë spp. (regardless of their geographical origin), constituting a highly supported clade. The plastid and concatenated topologies showed a well-supported and monophyletic Hierochloë section Monoecia as sister to the Eurasian and North American species. The nuclear tree failed to recover this structure, and section Monoecia was paraphyletic with H. quebrada as sister to all the other *Hierochloë*. A large polytomy including seven independent clades can be observed in the nuclear topology (H. redolens from South-East Asia; H. alpina; H. odorata + H. hirta + H. glabra; H. laxal; H. pusilla; H. occidentalis; Hierochloë section Monoecia except H. quebrada and H. pusilla; Fig. 5). The plastid tree recovered a weakly supported position for the South-East Asian H. redolens in the Eurasian + North American clade as the earliest diverging group, whereas the concatenated tree strongly supports H. occidentalis as sister to all other Hierochloë not in Monoecia. With regard to Hierochloë section Monoecia, H. pusilla is well supported as the first diverging group (Figs 4, S3) followed by H. quebrada (not represented in the concatenated tree). The position of *H. juncifolia* is not well resolved in the plastid tree (Fig. 4), but the nuclear and concatenated topologies (Figs 5, S3) clearly place it as sister to all other Monoecia, except H. pusilla and H. quebrada. The phylogenetic resolution of the H. redolens species complex, including H. redolens (central and southern Andean), H. altissima, H. gunckelii, H. utriculata and H. spicata (also H. juncifolia in the plastid tree; Fig. 4), is poor (Figs 4, 5, S3). The nuclear and concatenated topologies (Figs 5, S3) recover a well-supported position for an Argentinian H. redolens sample (H177, Appendix 1), positioning it as sister to all other specimens in the clade. In the plastid topology (Fig. 4) this position is occupied by a central Andean H. redolens (H129, Appendix 1), with low support. No structure could be obtained for H. altissima, H. gunckelii, H. spicata and H. utriculata.

The final aligned ETS clone matrix included 33 consensus sequences (see Material and Methods) and 851 characters representing eight South American Hierochloë spp. + A. mexicanum and A. odoratum (outgroup). Consensus sequences were generated for 353 ETS clones. Only H. quebrada, H. pusilla and H. redolens (Indonesia) were represented by one monospecific consensus sequence, including nine, ten and eight clones, respectively. Hierochloë juncifolia (78 clones) was represented by 12 consensus sequences, 11 of which were monospecific. Hierochloë altissima (80 clones) and H. utriculata (57 clones) were represented by six and eight consensus sequences, three and five of which included only one taxon, respectively. South American H. redolens (79 clones) and H. spicata (12 clones) participated in seven (four monospecific) and two consensus sequences, respectively, and all H. gunckelii (20 clones) were included in one non-monospecific consensus sequence (Appendix 1, Supporting Information Fig. S4). The number of clones represented for each consensus sequence was highly variable. Fourteen of them were based on one clone only (six of them belonging to *H. juncifolia*), whereas one consensus sequence accounted for 264 (72.7%) of the clones. This latter sequence represented a high percentage of the clones of all species of Hierochloë section Monoecia, except H. juncifolia and H. pusilla. Clones obtained from an individual tended to be represented by the same consensus sequence; monoclonal consensus sequences represent the exceptions to this trend.

The Bayesian consensus tree based on the ETS clone matrix of consensus sequences is represented in Figure

Figure 2. Scatterplots of macromorphological quantitative data. (A) Two-dimensional scatterplot of the linear discriminant analysis (LDA), classification functions 1 and 3. (B) Two-dimensional scatterplot of the principal components analysis (PCA), components 1 and 3. The percentage of variability explained by each component and the contributions of single characters are indicated in Table 2.

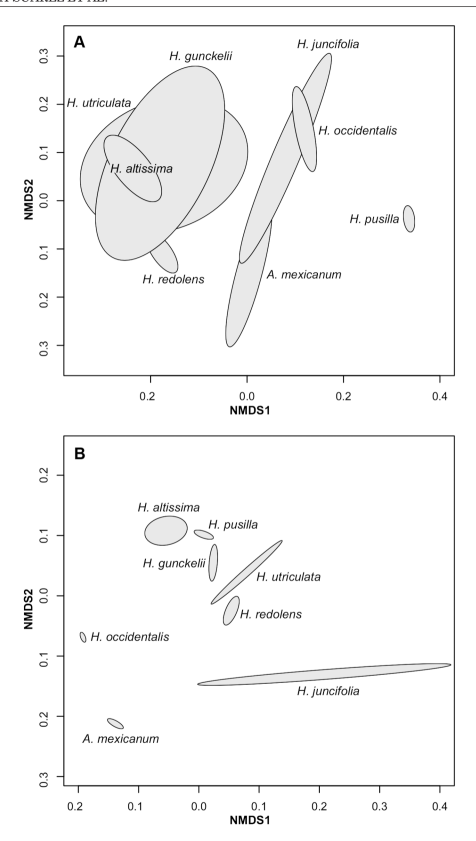


Figure 3. Plots of the non-metric multidimensional scaling (NMDS) analysis. (A) Combined macromorphological quantitative and qualitative data. (B) Combined micromorphological quantitative and qualitative data. Data combination was conducted using Gower's similarity coefficient.

S4, and is largely consistent with the phylogenetic tree based on nuclear data (Fig. 5). Resolution in the tree is generally low and only few relationships are strongly supported. Hierochloë section Monoecia is recovered as monophyletic with respect to A. mexicanum, whereas H. quebrada is sister to the remaining taxa in the section with strong support. All consensus sequences including clones from specimens of H. altissima, H. utriculata, H. juncifolia, H. pusilla, H. gunckelii, H. spicata and H. redolens (all populations) are grouped (low support) in a large polytomy with only three moderately to strongly supported clades: a strongly supported group of two consensus sequences representing two clones of H. utriculata and H. altissima; a moderately supported clade grouping all but two *H. juncifolia* consensus sequences and one H. redolens from the Central Andes; and a third group including three consensus sequences of *H. redolens*, H. juncifolia and H utriculata (six clones).

Phylogenetic networks were built to represent ambiguities in the phylogenetic reconstructions. Different explanations lie behind these ambiguities, including hybridization, homoplasy and, in the case of nuclear ribosomal sequences, recombination (Posada & Crandall, 2001). The lack of resolution in the phylogenetic trees for Hierochloë section Monoecia (Figs 4, 5) renders this approach useful even if it has been traditionally used at an intraspecific level (Clement, Posada & Crandall, 2000). The statistical parsimony networks produced with plastid (Supporting Information Fig. S5) and nuclear sequences (not shown) were largely consistent with the plastid and nuclear Bayesian topologies (Figs 4, 5). Samples of Hierochloë section Monoecia except H. juncifolia, H. pusilla and H. quebrada are grouped in one haplotype, with these three latter species closely related to it. Eurasian and North American species (+ South-East Asian H. redolens) occupy more distant positions in the network. Two missing intermediaries were created (Clement et al., 2000) and one cycle was observed in the plastid network affecting Eurasian, North American, South American and South-East Asian samples (Fig. S5). No cycle is defined in the nuclear network and eight missing intermediaries were inferred. The statistical parsimony network based on clone data (Fig. 6) groups 264 out of 352 clones (75%) in one ribotype. Twentyfour additional ribotypes were defined in the network: H. juncifolia (nine), H. juncifolia + H. altissima (one), H. altissima (two), H. altissima + H. redolens (one), H. redolens (six including the South-East Asian populations), H. utriculata (four) and H. quebrada (one). Two cycles were defined in this network and ten missing intermediaries were inferred.

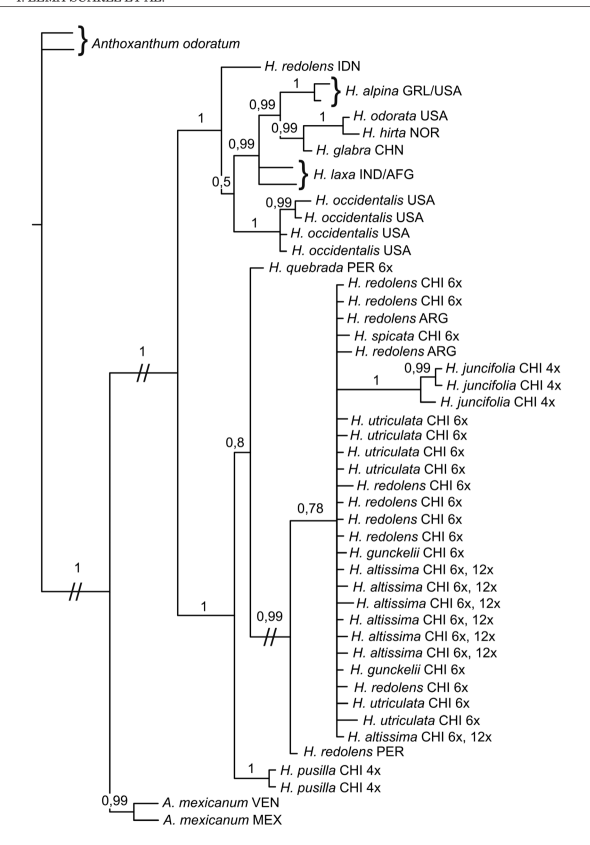
DISCUSSION

Integrative approaches offer the best results in exploring biodiversity (e.g. Hörandl, 2007; Schlick-Steiner et al., 2010). Here, we combine macromorphological, micromorphological, molecular and cytological information to assess the taxonomy of Hierochloë section Monoecia. We conclude that the taxonomic status of several of the species in the section is not justified based on either the phylogenetic (apomorphic) or the phenetic species concepts (for a discussion on the application of different species concepts in plants, see Soltis et al., 2007). In addition, we recovered a scenario of recent and incomplete diversification in the southern Andes for section Monoecia and a longer isolation history for those species with restricted ecological and/or geographical ranges.

Inference of DNA ploidy in $\emph{HIEROCHLOË}$ SECTION $\emph{MONOECIA}$

Hierochloë displays a wide variation in ploidy, with diploids, tetraploids, hexaploids and duodecaploids being identified (e.g. Weimarck, 1971). As a reference for the tentative translation of our nuclear DNA content values into DNA ploidies we used the measurements performed by Murray, de Lange & Ferguson (2005) in New Zealand Hierochloë. These authors found duodecaploids: H. redolens, H. fusca Zotov and *H. brunonis* Hook (12x; 2n = 84; 2C = 27.55-29.97 pg);hexaploids: *H. equiseta* Zotov (6x; 2n = 42; 2C = 18.10 pg);and tetraploids: *H. novae-zelandiae* Gand (4x; 2n = 28;2C = 12.54 pg). Considering this, our groups G1 and G2 (2C = 12.82–14.45 pg; Table 1, Fig. S1), including H. pusilla and H. juncifolia, correspond to DNA tetraploids; groups G3 (*H. altissima* population A1, H. redolens population R4 and H. utriculata) and G4 (H. gunckelii and H. redolens populations R1, R2 and R3) are putative DNA hexaploids (2C = 18.27–19.84 pg) and group G5 (*H. altissima* populations A2, A3, A4; 2C = 27.04 - 27.43 pg) most probably corresponds to DNA duodecaploids (for population codes see Appendix 1 or Fig. 1). Hierochloë quebrada and H. spicata are probably a DNA hexaploid (2C = 18.87 pg) and a DNA tetraploid (2C = 13.28 pg), respectively, but only one measurement could be attained for these taxa (Table 1).

Genome size variation and the number of chromosomes are not always positively correlated (e.g. Chumová, Mandáková & Trávníček, 2016), and the likelihood of chromosome rearrangements increases with evolutionary distance (Bhutkar et al., 2008). New Zealand taxa of Hierochloë do not belong to section Monoecia based on morphology; however, recent phylogenetic trees of Anthoxanthinae based on nuclear data (I. Lema-Suárez et al., unpubl. data) indicate that H. fusca is nested in the Monoecia clade and close to the



South American *H. redolens* complex. This evolutionary proximity gives consistency to our results; however, the inferred ploidies must be interpreted with caution. B chromosomes have been found in Anthoxanthinae (Chumová *et al.*, 2016), including in some *Hierochloë* Weimarck (1978). However, according to Chumová *et al.* (2015), the contribution of B chromosomes to the total nuclear DNA content is small in different groups of Anthoxanthinae. It is therefore unlikely that they are responsible for the differences in nuclear DNA content observed among and (in some cases) within taxa (Chumová *et al.*, 2015).

EVOLUTION AND BIOGEOGRAPHY OF $HIEROCHLO\ddot{E}$ SECTION MONOECIA

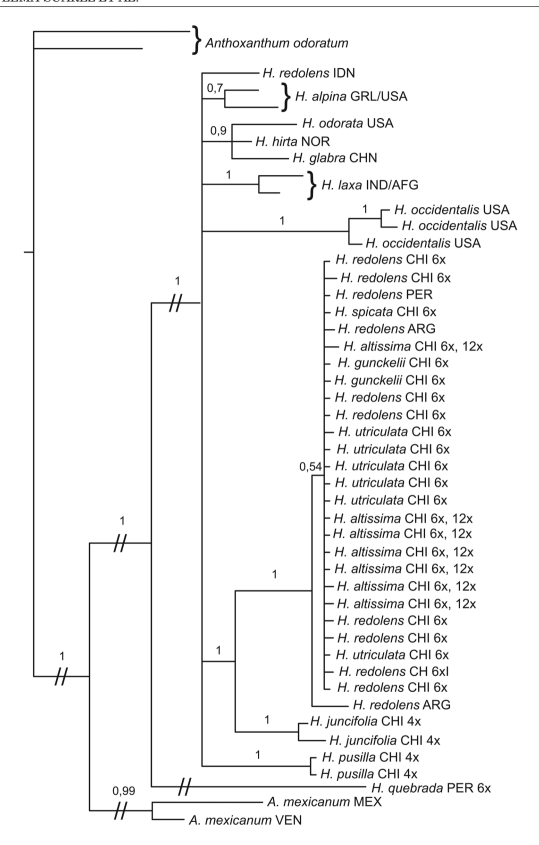
Phylogenetic analyses (e.g. Figs 4, 5, S3) recover a well-defined and monophyletic section *Monoecia* that is clearly separated from other South American groups of Anthoxanthinae. This result is consistent with Connor & Renvoize (2009) who established that spikelet monoecism is a good indicator of evolutionary isolation in the genus. Andromonoecious spikelets are dominant in *Hierochloë* (Connor, 2012) and only one transition to monoecism has been detected. Reversals in this transition have not been reported, although recent phylogenetic results (I. Lema-Suárez *et al.*, unpubl. data) indicate that some andromonoecious taxa are nested in the *Monoecia* clade (e.g. *H. fusca*, a New Zealand endemic).

All analyses recover the split of A. mexicanum [A. davidsei (Pohl) Veldkamp is not represented in the tree] prior to the divergence of section Monoecia and the development of monoecism in Hierochloë. Connor & Renvoize (2009) suggested that monoecious Hierochloë section Monoecia species evolved from non-South American andromonoecious ancestors through the development and fixation of male sterility genes in the apical floret. According to these authors, 'the Ataxia system could not give rise to monoecism' and therefore the central to northern South American Ataxia spp. A. mexicanum and A. davidsei can be excluded as ancestors of section Monoecia (Connor & Renvoize, 2009). Our molecular phylogenetic trees and networks support this view. Our limited sampling and phylogenies do not allow us to indicate with certainty where these ancestors grew or when the section diverged from the remaining *Hierochloë*, but Nearctic

(and Central American) colonizers are among the main components of the Andean flora since the mid-Miocene (e.g. Simpson, 1983). A Nearctic origin has been detected for other groups of high-elevation Andean grasses such as Festuca L. (Inda et al., 2008), and longdistance colonization of high mountain environments by cold-adapted lineages has been observed in different tropical alpine areas of the world (e.g. Gehrke & Linder, 2014; Luebert & Weigend, 2014; Merckx et al., 2015), including in Anthoxanthinae (Tusiime et al., 2017). Our well-supported plastid and concatenated phylogenetic trees (Figs 4, S3) do not seem to support this scenario, as they recover an old divergence for section Monoecia and do not show a close association between South American and Nearctic (or Holarctic) Hierochloë.

Taxa in section Monoecia represent a mixture of well-defined species and other species not yet well differentiated based on morphology or molecular data (e.g. Figs 2, 3A, 4, 5). A clear correlation can be observed between morphological and phylogenetic differentiation and ecological, geographical and/or cytological isolation. The central Andean hexaploid H. quebrada and the southern Andean tetraploid H. pusilla are the first groups to diverge in section Monoecia (Figs 4, 5, S3). Only the plastid phylogenetic tree did not recover a strongly supported position for *H. quebrada* (Fig. 4), and all the analyses based on ETS clones (Figs 6, S4) failed to produce a well-differentiated *H. pusilla*. This latter species, well characterized regarding macromorphological vegetative characters, especially plant size (Fig. 2; e.g. Parodi, 1941), is ecologically restricted to moist meadows in the Patagonian and Tierra del Fuego floristic provinces (De Paula, 1975). Only the hexaploid *H. redolens* grows in similar areas, but both taxa seem to be completely isolated (Figs 4, 5). Hierochloë pusilla shows low levels of macro- and micromorphological variation (Fig. 3). This might reflect its small distribution area and the general pattern of low morphological diversity for cold-adapted Antarctic/Arctic species (Grundt et al., 2006), although the number of analysed specimens was too low to draw any final conclusions. The ETS clones obtained from H. pusilla were reduced to one, monospecific consensus sequence indicating isolation, but the ETS clones-based network (Fig. 6) and phylogenetic tree (Fig. S4) failed to recover a well-supported position for this taxon. The high level of differentiation observed

Figure 4. Majority rule consensus tree inferred from Bayesian analysis (MrBayes) of plastid DNA sequences (*trnT-L* and *trnL-F*). Forty-eight samples representing 17 taxa of *Hierochloë* and *Anthoxanthum* section *Ataxia* and one outgroup (*A. odoratum*) are represented in the tree. The symbol // denotes branches that were shortened to simplify presentation. Posterior probability support values are represented above the branches. For each terminal, the species name is followed by country of origin (IDN, Indonesia; GRL, Greenland; USA, United States of America; NOR, Norway; CHN, China; IND, India; AFG, Afghanistan; PER, Peru; CHI, Chile; ARG, Argentina; VEN, Venezuela; MEX, Mexico).



is consistent with an old relative age for *H. pusilla*, a species restricted to the southernmost tip of the Andes (*sensu* Nagy & Grabherr, 2009; Körner *et al.*, 2011). Mountain southern species are generally older than those from the central or northern Andes (Doan, 2003; Luebert & Weigend, 2014) due to earlier uplift (Blisniuk *et al.*, 2005).

The central Andean putative hexaploid H. quebrada occupies different positions in the plastid and nuclear trees (Figs 4, 5). Discordance in topologies is common in plants and can be due to different causes, most commonly hybridization and incomplete lineage sorting(Renoultetal., 2009). The frequency of reticulate evolution in Hierochloë and in Anthoxanthinae (Pimentel et al., 2010, 2013; Chumová et al., 2017; Tusiime et al., 2017) and the variation of ploidy found in the section suggest hybridization as a likely explanation. Nevertheless, specific tests using more samples should be made to confirm this hypothesis for H. quebrada. With regard to the possible parents in a putative reticulation event, a plant from section Monoecia is the most likely plastid donor (Fig. 4), whereas the paternal parent must be outside the section. Hierochloë quebrada is restricted to high-elevation (4000–4600 m a.s.l.), arid areas on the western slopes of the central Andes (Connor & Renvoize, 2009). Two other species of section *Monoecia* currently grow at lower elevations in the central Andes: the hexaploid H. redolens and the tetraploid H. juncifolia (Tovar, 1993). The presence of the latter species has been questioned due to the description of H. quebrada based on materials identified as H. juncifolia (Connor & Renvoize, 2009). An analysis of a specimen of H. juncifolia collected in Ancash, Peru (US 2882401, Appendix 1), seems to indicate that H. juncifolia does grow in the central Andes. Our network based on ETS clones (Fig. 6) shows a close proximity between H. juncifolia and H. quebrada, pointing to H. juncifolia as the most likely plastid donor. Only two species of Anthoxanthinae outside Monoecia grow in the northern to central Andes, namely A. mexicanum and A. davidsei. The ploidy of these species and their phylogenetic position in Anthoxanthinae remain poorly known, but their lineages are clear candidates to be involved in the putative hybridization. With regard to the age of this reticulation event, the fact that all *H. quebrada* clones cluster in one consensus

sequence points to an old origin and the effect of concerted evolution (Okuyama *et al.*, 2005).

Nuclear and concatenated phylogenetic trees recover the tetraploid *H. juncifolia* as sister to the H. redolens complex (Figs 5, S3), whereas its position is not well supported in the plastid topology (Fig. 4). All analyses reveal H. juncifolia as a consistent group in section Monoecia, and highly distinct based on morphology or molecules (e.g. Figs 3, 4, S3). High variation is also detected in the species. Hierochloë juncifolia samples occupy a large morphological space in the NMDS analyses based on the combined macro- or micromorphological data (Fig. 3A, B). A similar pattern is observed in the ETS clone analyses. The phylogenetic tree (Fig. S5) recovered some (poorly supported) structure for the species, which was consistent with the clone-based network (Fig. 6). In addition, the 77 H. juncifolia clones were collapsed into 12 mostly monospecific consensus sequences, a number much higher than that obtained for the other species of Monoecia with a similar number of specimens sampled (Appendix 1). This scenario of high consistency combined with high interpopulational diversity may be due to the current ecology and distribution of the species. Hierochloë juncifolia grows in sandy or volcanic soils up to 1750 m a.s.l. in mountains of the Andean areas of the Argentine provinces of Río Negro, Neuquén and Chubut and in the adjacent regions of Chile (also in the central Andes of Peru, although no Peruvian sample could be used in the molecular analyses). Hierochloë juncifolia populations show complete ecological and (to a lesser extent) cytological and geographical isolation from the H. redolens complex (De Paula, 1975). They are also isolated from each other, since H. juncifolia has a highly discontinuous distribution due to its ecological requirements (De Paula, 1975; Anton & Zuloaga, 2012). This isolation is probably related to the emerging genetic structure in the species.

The *H. redolens* complex, also including *H. altissima*, *H. gunckelii*, *H. spicata* and *H. utriculata*, is the only group in the section that grows from the northern to the southern Andes. The phylogenetic analyses conducted recovered an almost complete lack of resolution for this group (e.g. Figs 4, 5, S3, S5), which might reflect its recent evolutionary origin and lack of reproductive isolation. The main exception to this

Figure 5. Majority rule consensus tree inferred from Bayesian analysis (MrBayes) of nuclear DNA sequences (ETS and ITS). Forty-seven samples representing 16 taxa of *Hierochloë* and *Anthoxanthum* section *Ataxia* and one outgroup (*A. odoratum*) are represented in the tree. The symbol // denotes branches that were shortened to simplify presentation. Posterior probability support values are represented above the branches. For each terminal, the species name is followed by country of origin (IDN, Indonesia; GRL, Greenland; USA, United States of America; NOR, Norway; CHN, China; IND, India; AFG, Afghanistan; PER, Peru; CHI, Chile; ARG, Argentina; VEN, Venezuela; MEX, Mexico).

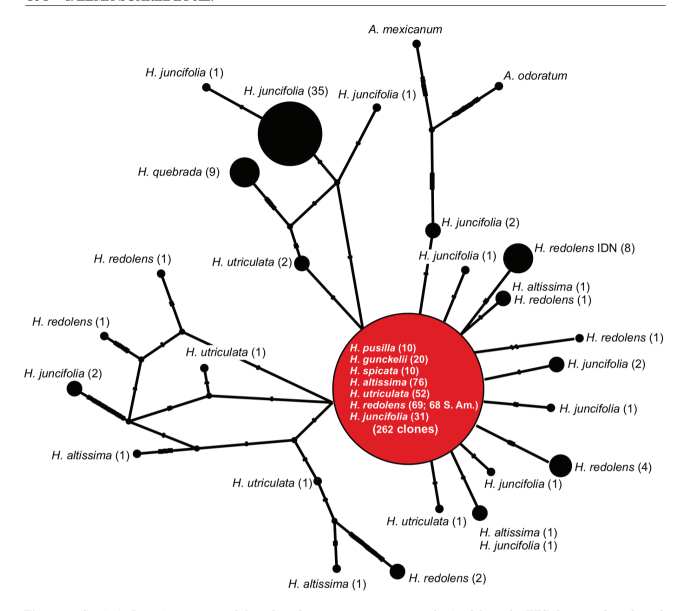


Figure 6. Statistical parsimony network based on the consensus sequences obtained from the ETS clones and conducted using the software TCS v. 1.21 and PopArt v. 1.7. S. Am., South America.

pattern is the position of the *H. redolens* accessions from Indonesia, which are clearly differentiated from the South American samples in all molecular analyses. Other authors have highlighted the need for a taxonomic reappraisal of *H. redolens* and have called for the separation of the Asian-Australian and South American andromonoecious populations of this taxon (Zotov, 1973; De Paula, 1975; Connor, 2012). It is important to note that although most of the ETS clones obtained from the Indonesian *H. redolens* sample clustered together in one exclusive consensus sequence, one Indonesian ETS clone clustered with the consensus sequence including most South American *H. redolens* complex clones. More research is needed to assess a

possible relationship between the South-East Asian and the South American populations.

The distribution range of the *H. redolens* complex in South America is clearly disjunct. It grows at low altitudes in temperate to subantarctic areas of Chile and Argentina and at mid- to high altitudes (up to 3850 m a.s.l.; Connor & Renvoize, 2009) in the tropical central and northern Andes. The plastid phylogenetic tree (Fig. 4) shows a well-supported sister relationship between the central and the southern Andean populations that is not recovered in the nuclear or concatenated trees (Figs 5, S3). Analyses based on clones recover a scenario of low differentiation between central and southern Andean populations. All clones

obtained from the Peruvian specimen collapsed in a consensus sequence that also comprised most clones in the southern South American H. redolens complex (Figs 6, S4). The biogeographical limit between the central and the southern Andes is the southern limit of the dry puna (Luebert & Pliscoff, 2006), an area of high aridity that acts as a filter to Andean elements, to the north and south (Arroyo et al., 1998). This barrier formed in the Miocene-Pliocene (e.g. Sepulchre et al., 2009; Heibl & Renner, 2012) as a result of Andean uplift together with the combined effect of the Humboldt Current and the Pacific anticyclone (Luebert & Weigend, 2014). It currently precludes the possibility of gene flow between the southern temperate and the northern tropicalalpine H. redolens, although whether gene flow was interrupted at the time of the formation of the barrier or afterwards remains to be determined.

The taxa of the *H. redolens* complex in southern South America have distribution and ecological ranges that are partially overlapping on a north-south gradient (De Paula, 1975; Anton & Zuloaga, 2012), and the obtained results must be interpreted in light of this. These groups, except *H. spicata* (tetraploid) and some *H. altissima* populations (duodecaploid), are putative hexaploids (Table 1), but differences in ploidy are not reflected in our phylogenetic analyses (e.g. Figs 4, 5). Differentiation among the species is minimal regardless of the analysis conducted, morphological or otherwise (e.g. Figs 2A, 4, 5, S2A), and only micromorphological data can separate all taxa (Figs 3B, S2B), except H. spicata. This species was not included in the analysis, but Villalobos & Finot (2016) indicated that it is not well defined based on leaf anatomy. The use of foliar micromorphological characters in grass taxonomy has been hindered by the effect of environmental parameters on leaf anatomy (Aiken, Darbyshire & Lefkovitch, 1984; Dubé & Moriset, 1996), including in Anthoxanthinae (Pimentel & Sahuquillo, 2008). Given the north-south distribution ranges of the species, new analyses based on additional populations are necessary to assess this point.

Analyses based on ETS clones revealed that (1) almost no structure exists in the data as regards the *H. redolens* complex (Figs 6, S4) and (2) most clones obtained from specimens of the complex collapse into one consensus sequence (80–100% of clones for *H. altissima*, *H. gunckelii*, *H. redolens*, *H. spicata* and *H. utriculata*; Fig. S4). Lack of phylogenetic resolution (Fig. S4) prevents us from making inferences about reticulation and introgression in this group; however, the general lack of differentiation observed, together with the abundance of multispecific consensus sequences (c. 50%) and the similar nuclear DNA estimations (and ploidy) lead us to believe that gene flow does exist in this group.

TAXONOMIC ASSESSMENT OF $HIEROCHLO\ddot{E}$ SECTION MONOECIA

Different species concepts have been applied in plants, especially in groups in which polyploidization is common (e.g. Soltis *et al.*, 2007). Here, we combined the morphologically based phenetic species concept (Sokal & Crovello, 1970) with the DNA-focused apomorphic species concept (Judd *et al.*, 2002). The first approach is practical for taxonomic purposes (e.g. Soltis *et al.*, 2007) and assumes that species are separated by a gap in morphological variation (Sokal & Crovello, 1970), whereas the second reduces subjectivity by recognizing only monophyletic groups that share several apomorphies, morphological or otherwise (Mishler & Theirot, 2000).

Ouranalyseshaveshownthatthemacromorphological traits used by Parodi (1941) and De Paula (1975) to underpin their classification of section Monoecia (plant size, inflorescence structure, awn insertion) are not useful for differentiating among the species unambiguously. Based on the two above-mentioned concepts and considering our results, only four species should be recognized in *Hierochloë* section *Monoecia*: H. juncifolia, H. pusilla, H. quebrada and H. redolens pro parte. This last name should be restricted to South American specimens following De Paula (1975, but see Zotov, 1973), and it would also include *H. altissima*, H. gunckelii, H. spicata and H. utriculata, as varieties. Given their micromorphological differences, we consider that they should be recognized taxonomically despite their lack of isolation. Although the limits between varieties and subspecies are far from clear-cut (e.g. McNeill et al., 2012), in our view the overlapping ecological and geographical ranges of these taxa would render the varietal rank more appropriate. Hierochloë sorianoi and H. moorei, not considered in our study, have already been transferred to *H. redolens* by other authors (e.g. Anton & Zuloaga, 2012), but recent results on H. moorei (Villalobos & Finot, 2016) call for a reassessment of this species.

CONCLUSIONS

Only four out of the eight *Monoecia* species (*H. pusilla*, *H. juncifolia*, *H. quebrada* and *H. redolens*) should be maintained based on morphological and molecular data. The remaining taxa (*H. altissima*, *H. gunckelii*, *H. spicata* and *H. utriculata*) should be reduced to varieties in *H. redolens*. The inclusion of South-East Asian and South American *H. redolens* populations in the same species is untenable according to our data.

Our results support the hypothesis proposed by Connor & Renvoize (2009), who suggested that species of *Hierochloë* section *Monoecia* evolved from non-South

American andromonoecious ancestors, and not from other South American Anthoxanthinae.

Our phylogenetic trees reveal a probable hybrid origin for the central Andean H. quebrada, although further studies are needed to confirm this hypothesis. The highly ecologically differentiated H. pusilla and H. juncifolia were shown to be the oldest species of the group. Extensive gene flow due to ecological, chorological and cytological similarities probably exists in the H. redolens complex.

New studies using a greater number of specimens and hypervariable nuclear markers are needed to clarify the structure in the *H. redolens* complex. Conducting chromosome counts in the different species is also necessary to confirm ploidies across the section.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Figure S1. Nuclear DNA content (pg/2C) in 82 plants of *Hierochloë* section *Monoecia* estimated using flow cytometric analysis. Specimens sorted according to increasing 2C-values. Putative DNA ploidies (see Discussion) were also included. Group G1 corresponds to *H. pusilla* (4x); G2 to *H. juncifolia* (4x); G3 and G4 to *H. altissima* population A1, *H. redolens*, *H. utriculata* and *H. gunckelii* (all 6x); G5 to *H. altissima* populations A2, A3 and A4 (12x). **Figure S2.** Plots of the non-metric multidimensional scaling (NMDS) analysis. (A) Qualitative macromorphological data. (B) Qualitative micromorphological data. Data combination was conducted using Gower's similarity coefficient.

Figure S3. Majority rule consensus tree inferred from Bayesian analysis (MrBayes) of concatenated nuclear (ETS and ITS) and plastid (trnT-L and trnL-F) DNA sequences. Forty-eight samples representing 17 taxa of $Hierochlo\ddot{e}$ and Anthoxanthum section Ataxia and one outgroup (Anthoxanthum odoratum) are represented in the tree. The symbol // denotes branches that were shortened to simplify presentation. Posterior probability support values are represented above the branches. For each terminal, the species name is followed by country of origin (IDN, Indonesia; GRL, Greenland; USA, United States of America; NOR, Norway; CHN, China; IND, India; AFG, Afghanistan; PER, Peru; CHI, Chile; ARG, Argentina; VEN, Venezuela; MEX, Mexico).

Figure S4. Majority rule consensus tree inferred from Bayesian analysis (MrBayes) of the consensus sequences obtained from the ETS clone DNA matrix. In total, 353 samples representing 17 taxa of *Hierochloë* and *Anthoxanthum* section *Ataxia* and one outgroup (*Anthoxanthum odoratum*) are represented in the tree. The symbol // denotes branches that were shortened to simplify presentation. Posterior probability support values are represented above the branches. S. Am.; South America. Numbers in parentheses represent the number of clones included in each ribotype (for more information see the Material and Methods).

Figure S5. Statistical parsimony network based on plastid data and conducted using the software TCS v. 1.21 and PopArt v. 1.7.

Appendix S1: Detailed description of the methodology followed in genome size estimation using flow cytometry.

Appendix S2: Detailed description of the statistical applyance applied to merphological data.

Appendix S2: Detailed description of the statistical analyses applied to morphological data.

Appendix S3: Maximum parsimony (MP) and Bayesian analyses conducted on molecular data.

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Appendix 1. Specimens and populations used in the analyses. Macrom, macromorphological characters; Microm, micromorphological characters; Phylog., phylogenetic tests; FCM, flow cytometry; Prov., province; Co, county; Mt., mountain. IDN, Indonesia; GRL, Greenland; USA, United States of America; NOR, Norway; CHN, China; IND, India; AFG, Afghanistan; PER, Peru; CHI, Chile; ARG, Argentina; VEN, Venezuela; MEX, Mexico. Numbers represent how many specimens were used per population in each analysis.

A. odoratum CHI IX Región de La Araucanía: Conguillio N.P. 2006 H173 CONC 173419 - H. alpina GRL Kangerlussuaq 2014 X149 - - H. alpina USA AK. Selavis Waring Mountains 2005 H135 US 3403659 - H. hirta NOR Hechnark: Hamar 2007 H43 OS 3103678 - H. hirta NOR Hechnark: Hamar 2007 H43 OS 3103678 - H. hirta NOR Hechnark: Hamar 2007 H43 OS 313689 - H. laxa AFG Kumark Prov.: Chapa Dara District 1977 H180 MASB005986 - H. dava AKC Valdex-Cordova C.A.: mile 173 Glenn 1959 H117 US 3495920 - A. mexicanum USA AK. Valdex-Cordova C.A.: mile 173 Glenn 1956 H10 US 3391684 1 A. mexicanum MEX Ohiapas State: Sierra de los Cuchumatanes 1966 H109 US 3391698 1 A. mexicanum <th>8</th> <th>1 111111 1 666 1 6</th> <th>ETS MF374535 MF374536 MF374545 MF374551 MF374552 MF374552 — MF374556 — MF374556</th> <th>TTS MF374580 MF374581 MF374582 MF374594 MF374596 — — — — — — — — — — — — — — — — — —</th> <th>MF678684 MF678685 MF678686 MF678697 MF678697 MF678701 MF678702 MF678702 — — — — — — — — — — — — — — — — — —</th> <th>MF678732 MF678733 MF678734 MF678740 MF678741 MF678741 MF678741</th> <th>CLONES MF688256- MF688264</th> <th></th>	8	1 111111 1 666 1 6	ETS MF374535 MF374536 MF374545 MF374551 MF374552 MF374552 — MF374556 — MF374556	TTS MF374580 MF374581 MF374582 MF374594 MF374596 — — — — — — — — — — — — — — — — — —	MF678684 MF678685 MF678686 MF678697 MF678697 MF678701 MF678702 MF678702 — — — — — — — — — — — — — — — — — —	MF678732 MF678733 MF678734 MF678740 MF678741 MF678741 MF678741	CLONES MF688256- MF688264	
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							MF688316	
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del Campo: Capitán Prat: Cochrane							MF688370	
H. juncifolia CHI XI Región Aysén del General Carlos Ibáñez 2006 H174 CONC 166409 –		I	ı	1	I	ı	MF688371-	1
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Appendix 1. Continued

District	Species	Country	Location	Date	Code	Herbarium	Macrom	Microm	Phylog					$_{ m FCM}$
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PRR													MF688443	
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195A ORe Courty Co. 195B Hot of PR682604 1 -	H. occidentalis	USA	CA: Marin Co.	1960	H89	PH 625586		ı	ı	1	ı	ı	1	ı
USA OR Coxe Co. 1929 HPF 568460 1 - <td>$H.\ occidentalis$</td> <td>$_{ m USA}$</td> <td>OR: Curry Co.</td> <td>1928</td> <td>H90</td> <td>$PH\ 662901$</td> <td>1</td> <td>ı</td> <td>MF374554</td> <td>MF374597</td> <td>MF678704</td> <td>MF678750</td> <td>1</td> <td>1</td>	$H.\ occidentalis$	$_{ m USA}$	OR: Curry Co.	1928	H90	$PH\ 662901$	1	ı	MF374554	MF374597	MF678704	MF678750	1	1
USA CA-Marin Co. 1936 1947 776544 1 - <td>$H.\ occidentalis$</td> <td>$_{ m USA}$</td> <td>OR: Coos Co.</td> <td>1929</td> <td>H91</td> <td>$PH\ 684550$</td> <td>1</td> <td>ı</td> <td>ı</td> <td>ı</td> <td>I</td> <td>1</td> <td>ı</td> <td>ı</td>	$H.\ occidentalis$	$_{ m USA}$	OR: Coos Co.	1929	H91	$PH\ 684550$	1	ı	ı	ı	I	1	ı	ı
USA CAN Marin Co. I 1985 H93 PH 994966 1 - <th< td=""><td>$H.\ occidentalis$</td><td>$_{ m USA}$</td><td>CA: Del Norte Co.</td><td>1936</td><td>H92</td><td>PH 756844</td><td>1</td><td>ı</td><td>I</td><td>I</td><td>ı</td><td>1</td><td>ı</td><td>ı</td></th<>	$H.\ occidentalis$	$_{ m USA}$	CA: Del Norte Co.	1936	H92	PH 756844	1	ı	I	I	ı	1	ı	ı
USA CAR Humboldt Bay 1918 H107 US 1061985 1 -	$H.\ occidentalis$	$_{ m OSA}$	CA: Marin Co.	1935	H93	$PH\ 994366$	1	1	I	ı	MF678705	MF678751	ı	ī
USA CA Monterey Co. Bigs Wikver 1949 1124 US 1649421 1 1 MF374555 MF374556 MF374550 MF678706 MF678752 - CO. Monterey Co. Bigs Wikver 1949 1124 US 2076057 1 1	$H.\ occidentalis$	$_{ m USA}$	CA: Humboldt Co: around Humboldt Bay	1918	H107	$US\ 1061985$	1	1	I	1	ı	I	1	ı
USA CA' Marin Co: Lagunitas Creek 1949 H124 US 2076155 1 -<	$H.\ occidentalis$	$_{ m NSA}$	CA: Monterey Co: Big Sur River	1937	H123	$US\ 1649421$	1	1	MF374555	MF374598	MF678706	MF678752	1	ı
Second Control of the Control of t	$H.\ occidentalis$	USA	CA: Marin Co: Lagunitas Creek	1949	H124	US 2079007	2	2	ı	I	I	ı	I	1
State Stat	$H.\ occidentalis$	Ω SA	CA: Santa Cruz Co: along Lockhart Gulch	1950	H125	$US\ 2976155$	1		1	1	1	1	1	ı
CHI XII Region de Magallanes y de la Antártida 2014 P1 SANT 73539 22			Road											
CHI XII Region de Magallanes y de la Antártida 2014 P1 SANT 73539 22	$H.\ occidentalis$	$_{ m NSA}$	CA: MarinCo: Mt. Tamalpais	1925	H126	$US\ 3182072$	1	1	1	ı	ı	1	1	ı
Chilena: near Pali Aike N.P.	H. pusilla	CHI	XII Región de Magallanes y de la Antártida	2014	P1	SANT 73539	22	4	MF374557	MF374600	MF678708	MF678754	MF688444-	70
Blanca Arcash Region: Recuay Prov.: Cordillera 1997 H121 US 3423081 1 MF374569 MF374602 MF678710 MF678756 MF			Chilena: near Pali Aike N.P.						MF374558	MF374601	MF678709	MF678755	MF688453	
Blanca	H. $quebrada$	PER	Ancash Region: Recuay Prov.: Cordillera	1997	H121	US~3423081	1	1	MF374559	MF374602	MF678710	MF678756	MF688454-	1
Herberton: Moat Herberton: Moat	$H.\ redolens$	ARG	Blanca Región de la Patagonia: Tierra del Fuego:	2008	H176		ı	ı	MF374560	MF374603	MF678711	MF678757	MF688462 -	I
Lake Roca	H. redolens	ARG	Herberton: Moat Región de la Patagonia: Tierra del Fuego:	2009	H177		I	1	MF374561	MF374604	MF678712	MF678758	I	1
CHI XII Region de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF374567 MF374505 MF374507 MF678719 MF678764 MF678719 MF678	H modelene	III	Lake Roca	9017	20 >				MESTARES	ME974608	017070710	MDG707EO		
CHI XII Region de Magallanes y de la Antártida 2014 R1 SANT 73533 6 4 MF374564 – MF678716 MF678718 MF678718 MF678718 MF678718 MF678718 MF678718 MF678718 MF678718 MF678718 MF678719 MF6			Chilono, I also Dogs						ME974569	ME974606	MF679714	MEG78760		
Chilena: Porvenir: Cape Boquerón MF678716 MF678716 CHI XII Región de Magallanes y de la Antártida 2014 R2 SANT 73534 4 4 MF374565 MF678717 MF678712 MI Chilena: Cameron: Cape Cameron: Inútil Bay MF678718 MF678718 MF678718 MF678763 MF678764 MI CHI XII Región de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF974567 MF678719 MF678764 MI	H. redolens	CHI	Ciliferal. Lane 100ca XII Región de Magallanes y de la Antártida	2014	R1	SANT 73533	9	4	MF374564	- WIT 01 #000	MF678715	MF678761	MF688463-	ro
CHI XII Región de Magallanes y de la Antártida 2014 R2 SANT 73534 4 4 MF374565 MF374607 MF678717 MF678762 MI Chilena: Cameron: Cape Cameron: Inútil Bay MF678718 MF678718 MF678718 MF678763 MF678763 MF678764 MI CHI XII Región de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF974567 MF678719 MF678764 MI			Chilena: Porvenir: Cape Boquerón								MF678716		MF688481	
Chilena: Cameron: Cape Cameron: Inútil MF678718 MF678718 MF678763 Bay Bay CHI XII Región de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF974567 MF678719 MF678764 MI	H. redolens	CHI	XII Región de Magallanes y de la Antártida	2014	R2	SANT~73534	4	4	MF374565	MF374607	MF678717	MF678762	MF688482-	20
Bay CHI XII Región de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF974567 MF678719 MF678764 MI			Chilena: Cameron: Cape Cameron: Inútil						MF374566		MF678718	MF678763	MF688492,	
CHI XII Región de Magallanes y de la Antártida 2014 R3 SANT 73532 4 1 MF374567 MF374608 MF678719 MF678764 MI			Bay										MF688494-	
Children Converse Con	H redolens	CHI	XII Reción de Macallanes v de la Antártida	2014	83	SANT 73539	4	-	MF374567	MF374608	MF678719	MF678764	MF688505 MF688506-	9
MF014303 MF016120 MF016103			Chilena: Cameron: Cape Cameron		}		,	,	MF374568		MF678720	MF678765	MF688525	,

Appendix 1. Continued

Species	Country	Country Location	Date	Code	Herbarium	Macrom	Microm	Phylog					FCM
								ETS	ITS	trnL- F	trnT- L	CLONES	
H. redolens	СНІ	XII Región de Magallanes y de la Antártida Chilena: Torres del Paine N.P.: above Lake	2014	R4	SANT 73535	Ø	63	MF374569	MF374609	1	MF678766	MF688526- MF688534	ro
H. redolens	IDN	Grey New Guinea: Papua Prov.: Maoke Mountains:	1938	H130	US 1761684	1	1	1	1	I	1	ı	ı
$H.\ redolens$	IDN	Sudirman Range: Lake Habbema New Guinea: Papua Prov.: Maoke Mountains:	1938	H131	$US\ 1761695$	1	1	MF374570	I	MF678721	MF678767	MF688535-	I
$H.\ redolens$	PER	Sudirman Range: 11 km NE of PuncakTrikora Ancash Region: Yungay Prov.: Huascarán	1984	H128	US 3097904	-	1	I	I	1	ı	MF688543 -	I
		N. P., between Lake Llanganuco and											
H. redolens	PER	rortacnuelo Cajamarca Region: Cajamarca Province: road	1988	H129	$US\ 3480791$	1	1	MF374571	MF374610	MF678722	MF678768	MF688544-	1
H. redolens	PNG	from Cajamarca to Bambamarca Central Prov.: Wharton Range: Mt. Albert	1974	H119	US 2741741	1	1	I	I	I	I	MF688550 -	1
H. redolens	PNG	Edward East Sepik Prov.: Mt. Burgers	1977	H120	US 2978618	73	73	I	I	I	ı	I	I
$H.\ spicata$	CHI	IX Región de La Araucanía: Cautín: Temuco	1923	H163	$US\ 1189418$	1	1	MF374572	MF374611	MF678723		MF688551-	1
$H.\ utriculata$	СНІ	VIII Región del Biobío: Arauco: Lake	2014	U1	SANT 73536	က	61	MF374573	MF374612	MF678724	MF678769	MF688560 MF688561-	9
$H.\ utriculata$	CHI	Lanalhue VIII Región del Biobío: Concepción: Hualpén:	2014	U2	SANT 73537	21	61	MF374574 MF374575	MF374613 MF374614	MF678725 MF678726	MF678770 -	MF688579 MF688580-	ro
$H.\ utriculata$	CHI	Cala Lenga IX Región de La Araucanía: Lake Budi:	2014	U3	,	I	ı	MF374576 MF374577	MF374615 MF374616	MF678727 MF678728	I	MF688596 MF688597-	70
		Puancho				96	99	MF374578 46	41	MF678729 48	41	MF688612 363	83

APPENDIX 2

Morphological characters used in this study. Micromorphological characters were selected and described according to Devesa (1992).

Macromorphological characters. Quantitative: 1. Plant height (mm) (PH). 2. Leaf length (mm) (LL) measured in the second leaf from the base of the plant. 3. Leaf width (mm) (LW) measured in the second leaf from the base of the plant. 4. Inflorescence length (mm) (IL). 5. Spike lower branch length (mm) (SLBL). The next three characters were measured in the second spikelet from the base of the inflorescence. 6. Spikelet length (mm) (SL). 7. Flower length (mm) (FL). 8. Lower glume length (mm) (LGL). Qualitative. Binary: 9. Convolute leaves (CL): planar (0), convolute (1). 10. Awn in male florets (AW): absence (0), presence (1). 11. Lower floret (LF): empty (0), male (1). **12. Intermediate floret** (MF): empty (0), male (1). Multi-estate: 12-I. Lower male floret awn inserted in the middle part (LGAM): yes (1), no (0). 12-II. Lower male floret awn inserted in the upper part (LGAS): yes (1), no (0). 13-I. Upper male floret awn inserted in the middle part (UGAM): yes (1), no (0). 13-II. Upper male floret awn inserted in the upper part (UGAS): yes (1), no (0). **14-I. Apical floret empty** (FE): yes (1), no (0). 14-II. Apical floret hermaphrodite (FH): yes (1), no (0). **14-III. Apical floret female** (FF): yes (1), no (0). 14-IV. Apical floret with staminodes (FFE): yes (1), no (0).

Micromorphological characters. Quantitative: 1. Long cells length (μm) (LlcL). 2. Stomata size (μm) (StS). 3. Number of ribs (NRi). Qualitative. Binary: 4. Companion cells type (CoC): round S5 (0), ax S2 (1). 5. Cilia (C): absence (0), presence (1). 6. Median vascular bundle sheath type (MVBT):

single (0), complete (1). **7. Median nerve** (MN): similar to the other nerves (0), bigger (1). 8. Furrows depth (FD): <3/4 of the total width (0), >3/4 of the total width (1). Multi-estate: 9-I. Long cell walls smooth (I3) (I3N): no (0), yes (1). 9-II. Long cell walls moderately undulating (I3P): no (0), yes (1). 9-III. Long cell deeply undulating (I3R): yes (1), no (0). **10-I. Bulliform cells** (CB): absence (0), presence (1). 10-II. Bulliform cells very abundant (CBM): no (0), yes (1). 11-I. Low density of hooks (HoF): absence (0), presence (1). 11-II. Intermediate density of hooks (HoN): absence (0), presence (1). 11-III. High **density of hooks** (HoM): absence (0), presence (1). 12-I. Stomata in adaxial surface (StAd): absence (0), presence (1). 12-II. Stomata in abaxial surface (StAb): absence (0), presence (1). 12-III. Stomata in **both surfaces** (StBo): absence (0), presence (1). **13-**I. Subepidermal sclerenchyma discontinuous (SScD): no (0), yes (1). 13-II. Subepidermal sclerenchyma continuous (SScC): no (0), yes (1). 13-III. Subepidermal sclerenchyma continuous and abundant (SScCA): no (0), yes (1). 14-I. Vascular bundles without sclerenchyma associated (PBS): no (0), yes (1). 14-II. Vascular bundles with sclerenchyma associated in the abaxial surface (PBAb): no (0), yes (1). 14-III. Vascular bundles with sclerenchyma associated only in the margin of the leaves (PBEx): no (0), yes (1). 15-I. Ribs shape (RS): presence of ribs (0), absence of ribs (1). 15-II. Ribs round-flattened (RSD): no (0), yes (1). 15-**III. Ribs round** (RSR): no (0), yes (1). **15-IV. Ribs** quadrangular (RSQ): no (0), yes (1). 15-V. Ribs polygonal (RSP): no (0), yes (1). 16-I. Leaf section planar (LSP): no (0), yes (1). 16-II. Leaf section open (LSO): no (0), yes (1). 16-III. Leaf section closed (LSC): no (0), yes (1).