Cytotaxonomic study of the Chilean endemic complex *Alstroemeria magnifica* Herb. (Alstroemeriaceae)

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Abstract

*Alstroemeria* L. (Alstroemeriaceae) represents one of the most diverse genera of vascular plants in Chile. It contains approximately 54 taxa, 40 of which are endemic. The “complex” *Alstroemeria magnifica* is endemic to Chile, and it comprises four varieties: *A. magnifica* var. *magenta*, *A. magnifica* var. *magnifica*, *A. magnifica* var. *sierrae*, and *A. magnifica* var. *tofoensis*. It is distributed from Coquimbo to the Valparaíso Region. We analyzed karyotypes of 10 populations along its natural distribution. All the populations presented an asymmetric karyotype, with 2n = 16 chromosomes but with three different karyotypic formulae. *Alstroemeria magnifica* var. *magenta* and *A. magnifica* var. *sierrae* presented the same karyotypic formula, and *A. magnifica* var. *magnifica* and *A. magnifica* var. *tofoensis* each had a different formula. The scatter plot among CVCL vs. MCA shows different groupings between populations of the four varieties. Based on the results, it is possible to consider raising *Alstroemeria magnifica* var. *magenta* to species level (*A. magenta*) and *A. magnifica* var. *tofoensis* to subspecies level (*A. magnifica* subsp. *tofoensis*); *A. magnifica* var. *magnifica* and *A. magnifica* var. *sierrae* should each remain as varieties. Nevertheless, these taxonomic changes should be considered tentative, as additional sources of evidence become available.

Keywords: *Alstroemeria*, karyotype, species complex, asymmetry, Chile.

Received: May 30, 2017; Accepted: November 17, 2017.

Introduction

*Alstroemeria* is a South American genus, which comprises about 82 taxa distributed from Venezuela (3°N) to Tierra del Fuego (53°S) (Muñoz and Moreira, 2003). The centers of distribution of this genus are located in Central Chile and East of Brazil, representing a disjoint pattern of distribution produced by the isolating effect of the Cordillera de Los Andes and the South American Arid Diagonal (Muñoz and Moreira, 2003; Hofreiter, 2007; Chacón et al., 2012a).

In Chile, *Alstroemeria* represents one of the most diverse genera of vascular plants, comprising 49 taxa (33 species, 8 varieties, and 8 subspecies); 40 of which are endemic (Muñoz and Moreira, 2003). Recent studies suggest increasing to 54 the number of taxa recognized in *Alstroemeria*, with the validation of *Alstroemeria citrina* Phil. (Eyzaguirre, 2008) and *Alstroemeria parvula* Phil. (Muñoz et al., 2011). These modifications also include the discovery of *Alstroemeria hookeri* Lodd. subsp. *sansebastiania* C.M. Baeza & E. Ruiz (Baeza and Ruiz, 2011), *Alstroemeria marticorenae* Negritto & C. M. Baeza (Negritto et al., 2015) and *Alstroemeria traudliae* (Hoffmann et al., 2015).

Reports of chromosome studies in *Alstroemeria* are dated since 1882, recognizing a fundamental karyotype on about 30 taxa, 22 of them from Chilean species (Chacón et al., 2012b). A stable chromosome set of 2n = 16 was determined, with an asymmetric and bimodal karyotype of eight chromosomes: three or four are acrocentric and four or five are metacentric, submetacentric or subtelocentric (Baeza et al., 2008). Until today, no reports of polyploids have been observed in natural populations of *Alstroemeria* (Baeza et al., 2007a).

Cytogenetic studies have proven useful for the delimitation of entities in *Alstroemeria* since every studied taxon
presents a distinctive, unique, and largely stable karyotype. As such, these studies have contributed not only to the delimitation of species and varieties, but also to elicit underlying processes - at chromosomal levels - that determine the divergence of these taxa (Baeza et al., 2007b). In taxonomic complexes, a clear-cut discrimination of intraspecific taxa has resulted from the study of differences in the architecture and/or the asymmetry of chromosomes (Cajas et al., 2009; Baeza et al., 2010). For example, the study of karyotype was determinant for the delimitation of taxa within the *Alstroemeria hookeri* complex, providing evidence for the existence of *Alstroemeria hookeri* subsp. *sansebastiana* (Baeza and Ruiz, 2011) and supporting the proposal of Muñoz and Moreira (2003) to raise the status of *Alstroemeria hookeri* subsp. *cummingiana* to species level.

The difficult differentiation and delimitation of taxa within *Alstroemeria* have led to the definition of intraspecific complexes, which consist of two or four subspecies and/or varieties of the same species. Some of these complexes have become relevant taxa for the prospecting of development in different areas of national interest, given their economic potential as ornamental plants, and/or their importance as representative taxa of the Chilean biodiversity. Traditional taxonomic treatments in *Alstroemeria* have largely been based on patterns of variability of their conspicuous flowers, which present tepals with a wide display of morphological and coloring patterns (Bayer, 1987; Muñoz and Moreira, 2003). These structures, while useful for discrimination at interspecific levels, usually exhibit levels of variation beyond levels of stability required for a robust taxonomic discrimination in several groups of species (Baeza et al., 2010, 2015, 2016a,b). Therefore, since any potential development in these groups has been mostly restricted to inconclusive taxonomic interpretations the use of other possible sources of evidence, like cytological characters, could result useful to help taxonomic discrimination at intraspecific levels.

*Alstroemeria magnifica* Herb. is a species complex which comprises four varieties (Muñoz, 2003; Muñoz and Moreira, 2003). Along with *A. hookeri*, *A. magnifica* is one of the richest in terms of number of taxa in *Alstroemeria*. This complex is distributed from the locality of Chuncungo (29°26’S; Region of Coquimbo) to the north of Papudo (32°21’S; Region of Valparaiso). It is recurrent across coastal rocky bluffs and slopes, most likely in areas with permanent fog. The specific ranges of distribution and description of floral morphology are the following:

*Alstroemeria magnifica* Herb. var. *magenta* (Ehr.Bayer) Muñoz-Schick: This variety presents the largest range of distribution within the complex, ranging from 30°39’S to 32°21’S. It is distinguished by the presence of both small inflorescences and flowers. Their internal upper tepals show thick lines, which end in a respective spot at the apex. The internal lower tepal can present design or not (Figure 1B).

*Alstroemeria magnifica* Herb. var. *sierrae* (Muñoz) Muñoz-Schick: This variety occurs in a restricted distribution, from 29°36’S to 29°45’S. It presents flowers of large size, which are distinctive by the design present in the internal upper tepals, and having lines forming a large spot at the apex and basis of this structure. The internal lower tepal can present design (Figure 1C).

*Alstroemeria magnifica* Herb. var. *tofoensis* Muñoz-Schick: This variety has a very restricted distribution, from 29°26’S to 29°32’S. It is characterized by their internal upper tepals with a patch of a yellowish spot and a white background, which does not reach the borders. Scattered lines with no spotty end at the apex are also present. The internal inferior tepal is maculate at the basis (Figure 1D).

Until now, cytological work has not been extensive for completely clarifying the taxonomic status of the infraspecific taxa of the *A. magnifica* complex. Buitendijk and Ramanna (1996) analyzed the karyotype of *A. magnifica* subsp. *magnifica*, informing the morphology of chromosomes and interspecific variability in the C-bands patterns. Additional reports exist about the genomic size (Buitendijk et al., 1997, 1998), which in association with patterns of variation in C-bands, suggest discontinuous variation in the quantity of nuclear DNA of *A. magnifica*. Nevertheless, since these observations are circumscribed to cultivated specimens only (mostly cultivars and greenhouse varieties), no additional cytological information exists from local populations and/or range of variation within their natural environmental and geographic ranges.

Therefore, the present study aims to characterize and compare, at a cytotaxonomic level, the four varieties of the *A. magnifica* complex. Thus, using representative sampling from the total of the geographic range of distribution, we expect to offer a suggested clarification of the taxonomic status of each variety within the complex.

Materials and Methods

Plant material

A total of two to four individuals from 10 populations of *A. magnifica* were collected across the known range of distribution (Table 1). Voucher specimens from each population were deposited in the Herbarium of the University of Concepción (CONC). Figure 2 shows the distribution of the collected populations, which were used in the present study.

Methodology for the study of karyotypes

Rhizome roots (1-2 cm length) obtained from individuals in each population and held in a greenhouse, were cut and pre-treated with a solution of 8-hydroxyquinoline (2
**Figure 1** - Photographic representation of varieties present in the *A. magnifica* complex. (A) Photography of *A. magnifica* var. *magnifica*; (B) Photography of *A. magnifica* var. *magenta*; (C) Photography of *A. magnifica* var. *sierrae*; (D) Photography of *A. magnifica* var. *tofoensis*. Bar = 2 cm.

**Table 1** - Plant material for the analyzed populations.

| Species          | Population | Locality and Date of Collection. | Latitude S/ Longitudes W | Altitude (m) |
|------------------|------------|----------------------------------|--------------------------|--------------|
| *A. magnifica* var. *magnifica* | 4408       | Región de Coquimbo. Provincia de Elqui. Inicio Cuesta Buenos Aires. 5 de octubre de 2014 | 29°34'18''/71°14'35'' | 473          |
|                  | 4411       | Región de Coquimbo. Provincia de Elqui. Entre Cuesta Porotitos y Caleta Hornos. 7 de octubre de 2014 | 29°44'32''/71°19'20'' | 150          |
|                  | 4414       | Región de Coquimbo. Provincia de Limarí. Bosque Hidrófilo, parte alta. 8 de octubre de 2014 | 30°39'45''/71°40'57'' | 598          |
| *A. magnifica* var. *magenta* | 4379       | Región de Coquimbo. Provincia de Choapa. Bosque Santa Julia, fundo Agua Amarilla. 31 de octubre de 2013 | 31°49'48''/71°30'35'' | 110          |
|                  | 4380       | Región de Coquimbo. Provincia de Choapa. Entre quebrada El Negro y Los Vilos. 31 de octubre de 2013 | 31°57'20''/71°29'14'' | 138          |
|                  | 4381       | Región de Coquimbo. Provincia de Choapa. Fundo Palo Colorado. 5 km al norte de Puente Quilimarí, frente al Cerro Tentén. 1 de noviembre de 2013 | 32°05'58''/71°30'27'' | 80           |
|                  | 4383       | Región de Valparaíso. Provincia de Petorca. 2 km al sur de Los Molles. 1 de noviembre de 2013 | 32°14'35''/71°29'27'' | 37           |
| *A. magnifica* var. *sierrae* | 4406       | Región de Coquimbo. Provincia de Elqui. Juan Soldado. 5 de octubre de 2014 | 29°43'04''/71°18'25'' | 175          |
|                  | 4407       | Región de Coquimbo. Provincia de Elqui. Caleta Hornos. 5 de octubre de 2014 | 29°38'01''/71°17'08'' | 152          |
| *A. magnifica* var. *tofoensis* | 4409       | Región de Coquimbo. Provincia de Elqui. Mina El Tofo. 6 de octubre de 2014 | 29°26'56''/71°14'52'' | 676          |
mM) for 24 h at 4 ºC. These samples were subsequently fixed with a fresh solution of ethanol/acetic acid (3:1) for 24 h. Squash preparations from root tips were made using an acid hydrolysis pretreatment with HCL 0.5 N during 17 min at 42 ºC. After washing in distilled water, the material was stained with 1% orcein solution. Metaphase chromosome plates were photographed using a Zeiss Axioskop microscope, with an incorporated video camera. Chromosomes were measured with the assistance of the software MicroMeasure 3.3 (Reeves, 2001) and classified according to arm ratios (long arm/short arm; modified from Levan et al., 1964). From 10 metaphase plates in each analyzed population, randomly chosen from the total of individuals, an idiogram was constructed for each studied variety. Intra-chromosomal asymmetry (MCA) and interchromosomal asymmetry (CVCL) indices were calculated for each analyzed population, following the proposal of Peruzzi and Erôglu (2013). Both indices were placed in a scatter plot, accompanied by the total length of diploid chromosomes (TCL), for each analyzed population using the package plot3D v 1.1 (Soetaert, 2016) in R v 3.3.3 (R Core Team, 2017).

Results and Discussion

All analyzed populations of A. magnifica revealed a $2n = 2x = 16$. A. magnifica var. magnifica and A. magnifica var. sierrae presented the same haploid formula: two pairs of metacentric chromosomes, one submetacentric pair, two subtelocentric pairs, two subtelocentric pairs with satellite and one telocentric pair with satellite (2m + 1sm + 2st + 2st-sat + 1t-sat; Figure 3B). A. magnifica var. tofoensis presented a haploid formula of two metacentric chromosomes, one submetacentric pair, one subtelocentric pair, two telocentric pairs and two telocentric pairs with satellite (2m + 1sm + 1st + 2t + 2t-sat; Figure 3D). Figure 4 shows representative metaphase plates for each studied taxon. The values of CVCL, MCA and TCL per populations are summarized in Table 2. Figure 5 represents the scatter plot of CVCL and MCA indices.

Karyological studies have previously reported about the morphology of chromosomes, patterns and polymorphism of C-bands, nuclear content, and the genomic size of A. magnifica (Buitendijk and Ramanna, 1996; Buitendijk et al., 1997, 1998). The present study concurs with the findings made in those publications, specifically on the stability of the $2n = 16$ present in all varieties of A. magnifica. Additionally, our results support the typical asymmetric and bimodal karyotype present in Alstroemeria, with four to seven metacentric, submetacentric or subtelocentric chromosomes (Baeza et al., 2008). Within this cytological configuration, it is possible to distinguish, at least, two patterns that can discriminate the varieties of this complex.

First, while identical in structure, notorious differences are noticeable in the total length of chromosomes (TLC). This observation allows to distinguish A. magnifica var. magnifica from A. magnifica var. sierrae, as the former exhibits smaller chromosomes than the latter (Table 2), despite exhibiting identical karyotypes (Figures 3A and 3C). Populations from A. magnifica var. sierrae and A. magnifica var. magnifica are separated because of their differences in the CVCL index (Figure 5), which is directly related to the TLC values (Perruzzi and Erôglu, 2013). This cytological
pattern is intriguing, because of its recurrent presence in other complexes in *Alstroemeria*. For example, in the *Alstroemeria diluta* complex, both recognized subspecies, *A. diluta* subsp. *diluta* and *A. diluta* subsp. *chrysantha*, reveal similar karyotypes but a different TLC values (Baeza et al., 2016a). Such change in chromosome length could be the result of changes in the total genomic nuclear size of *A. magnifica* (Buitendijk et al., 1997, 1998), which could have implications as a mechanism of differentiation among taxa in *Alstroemeria*. Nonetheless, this circumstantial evidence should be further corroborated with additional studies based on nuclear DNA content (e.g., flow cytometry) and its variation across natural populations.

The second pattern is exhibited in *A. magnifica* var. *magenta* and *A. magnifica* var. *tofoensis*, which present different and unique karyotypes – compared to *A. magnifica* var. *magnifica* and *A. magnifica* var. *sierrae* (Figures 3 and 4). In this case, chromosome 3 of *A. magnifica* var. *magenta* is submetacentric, instead of subtelocentric or telocentric found in the other varieties of *A. magnifica*. In *A. magnifica* var. *tofoensis*, a polymorphism in the length of chromosome arms is detected between homologous chro-

**Figure 3** - Idiograms of *A. magnifica* varieties. (A) *A. magnifica* var. *magnifica*; (B) *A. magnifica* var. *magenta*; (C) *A. magnifica* var. *sierrae*; (D) *A. magnifica* var. *tofoensis*. 
mosomes of pair 5, which is also expressed in terms of greater levels of magnitude in standard variation related to TCL (Table 2). This pattern is in line with previous reports in A. philippii, where a population revealed length polymorphism between homologous in the chromosome pair 3 (Buitendijk et al., 1998). A similar situation has been found in species of Brachycome, Triticum, Tulipa, Secale, Allium (Houben et al., 2000), Scilla (Greilhuber and Speta, 1976),

![Metaphase plates in the varieties of A. magnifica complex.](image)

Table 2 - Karyotype features of the varieties of Alstroemeria magnifica. CV<sub>CL</sub> = Coefficient of variation of chromosome length; M<sub>CA</sub> = Mean centromeric asymmetry index according to Peruzzi and Erőglu (2013); SD = Standard deviation; TLC = Total length in diploid chromosomes.

| Varieties                      | CV<sub>CL</sub> ± SD | M<sub>CA</sub> ± SD | TLC ± SD   |
|--------------------------------|----------------------|---------------------|------------|
| A. magnifica var. magnifica (4408) | 62.0 ± 2.4           | 45.0 ± 1.6          | 128.2 ± 6.2|
| A. magnifica var. magnifica (4411) | 60.0 ± 4.3           | 42.0 ± 2.3          | 136.5 ± 8.3|
| A. magnifica var. magnifica (4414) | 62.0 ± 3.5           | 47.0 ± 2.0          | 130.4 ± 5.9|
| A. magnifica var. sierrae (4406) | 46.0 ± 3.6           | 51.0 ± 1.8          | 187.5 ± 7.2|
| A. magnifica var. sierrae (4407) | 47.0 ± 4.2           | 50.0 ± 2.1          | 193.5 ± 6.8|
| A. magnifica var. tofoensis (4409) | 55.0 ± 3.9           | 55.0 ± 1.5          | 173.9 ± 9.2|
| A. magnifica var. magenta (4379) | 54.0 ± 4.8           | 51.0 ± 2.4          | 104.6 ± 8.8|
| A. magnifica var. magenta (4380) | 55.0 ± 5.3           | 51.0 ± 1.3          | 100.1 ± 4.6|
| A. magnifica var. magenta (4381) | 55.0 ± 4.4           | 49.0 ± 1.8          | 109.4 ± 5.2|
| A. magnifica var. magenta (4383) | 53.0 ± 3.1           | 50.0 ± 2.2          | 110.4 ± 5.9|
Placea amoena (Baeza and Schrader, 2004), and Chaetanthera pentacaenoides (Baeza and Torres-Díaz, 2006).

In terms of the overall differentiation among the varieties of the *A. magnifica* complex, a better characterization is possible to achieve by using individual patterns of asymmetry in chromosomes. *A. magnifica* var. *magnifica* and *A. magnifica* var. *magenta* revealed substantial differences in their M<sub>CA</sub> and CV<sub>CL</sub> values, which also occurs with *A. magnifica* var. *tofoensis*; nonetheless, the latter with similar patterns than *A. magnifica* var. *magenta* (Figure 5). Despite this, the karyotype of *A. magnifica* var. *magnifica* and *A. magnifica* var. *magenta* exhibits notorious differences, especially in the unique presence of a submetacentric chromosome in pair 3 of *A. magnifica* var. *magenta* compared to the rest of the complex. Furthermore, *A. magnifica* var. *tofoensis* presents a polymorphism in the homologous metacentric chromosomes in pair 5 (see above), while this is subtelocentric in *A. magnifica* var. *magenta*. These unique features suggest that both asymmetry patterns and karyotype variation should be considered together if this evidence is to be used for the precise discrimination of the involved varieties.

The results of this study suggest that patterns of chromosome variation can be instrumental for discriminating among taxa and proposing taxonomic rearrangements in the species complexes of *Alstroemeria*, as they tend to exhibit higher levels of stability and resolution than traditional tepal morphological characters at intraspecific levels (Cajas et al., 2009; Baeza et al., 2010, 2015, 2016a,b). In this case, these changes would be further supported, as cytological data is integrated and contextualized with preliminary results observed from additional character sources (chloroplast DNA, colorimetric variation and morphometry in chromosomes.

Figure 5 - Scatter plot among populations of *Alstroemeria magnifica* varieties using values of M<sub>CA</sub> vs. CV<sub>CL</sub>.

try of tepals; Carrasco et al. in preparation). For example, given the concordance of cytological data with patterns of discrete morphological variation, sympatric distribution and differentiation based in chloroplast DNA, it is likely that *A. magnifica* var. *sierra* and *A. magnifica* var. *magenta* should retain their taxonomic status without modifications. Instead, *A. magnifica* var. *tofoensis* status should be changed to subspecies level, because, despite presenting clear differentiation in floral characters, isolated distribution, and a distinctive unique karyotype, it presents close genetic similarity with *A. magnifica* var. *magnifica* and *A. magnifica* var. *sierra*. Likewise, it would be recommendable to revalidate *A. magenta* Bayer, from *A. magnifica* var. *magenta*, as originally proposed by Bayer (1987), given its consistent differences in vegetative and floral characters (smaller plants and flowers), allopatric distribution, a distinctive karyotype and substantial genetic distance from the rest of the taxa of *A. magnifica* complex. Nonetheless, these proposals should be seen as tentative, contingent upon additional and more conclusive results that can be added from the suggested character sources.

Acknowledgments

This study was supported by FONDECYT N° 1130349 and FONDECYT N° 3160453. The authors thank the Corporación Nacional Forestal (CONAF) for providing the permits that allowed collecting specimens in the Fray Jorge National Park. Thanks are also to the Departamento de Botánica and the Herbario de la Universidad de Concepción (CONC) for their assistance provided in this study.

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Associate Editor: Marcelo Guerra

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