

See discussions, stats, and author profiles for this publication at: <http://www.researchgate.net/publication/24283520>

# Karyotypes, heterochromatin, and physical mapping of 18S–26S rDNA in Cactaceae

ARTICLE *in* CYTOGENETIC AND GENOME RESEARCH · FEBRUARY 2009

Impact Factor: 1.91 · DOI: 10.1159/000200090 · Source: PubMed

---

CITATIONS

8

DOWNLOADS

58

VIEWS

181

## 4 AUTHORS, INCLUDING:



[Maria Laura Las Peñas](#)

National University of Cordoba, Argentina

19 PUBLICATIONS 37 CITATIONS

[SEE PROFILE](#)



[Juan Urdampilleta](#)

National University of Cordoba, Argentina

22 PUBLICATIONS 55 CITATIONS

[SEE PROFILE](#)



[Gabriel Bernardello](#)

National University of Cordoba, Argentina

84 PUBLICATIONS 1,041 CITATIONS

[SEE PROFILE](#)

# Karyotypes, heterochromatin, and physical mapping of 18S-26S rDNA in Cactaceae

M.L. Las Peñas<sup>a</sup> J.D. Urdampilleta<sup>b</sup> G. Bernardello<sup>a</sup> E.R. Forni-Martins<sup>b</sup><sup>a</sup>Instituto Multidisciplinario de Biología Vegetal (UNC-CONICET), Córdoba (Argentina)<sup>b</sup>Laboratório de Biosistemática, Departamento de Botânica, Instituto de Biologia Universidade Estadual de Campinas, Campinas (Brasil)

## Abstract

Karyotype analyses in members of the four Cactaceae subfamilies were performed. Numbers and karyotype formula obtained were: Pereskioideae = *Pereskia aculeata* ( $2n = 22$ ;  $10m + 1sm$ ), Maihuenioideae = *Maihuenia patagonica* ( $2n = 22$ ,  $9m + 2sm$ ;  $2n = 44$ ,  $18m + 4sm$ ), Opuntioideae = *Cumulo-puntia recurvata* ( $2n = 44$ ;  $20m + 2sm$ ), Cactoideae = *Acanthocalycium spiniflorum* ( $2n = 22$ ;  $10m + 1sm$ ), *Echinopsis tubiflora* ( $2n = 22$ ;  $10m + 1sm$ ), *Trichocereus candicans* ( $2n = 22$ ,  $22m$ ). Chromosomes were small, the average chromosome length was  $2.3 \mu\text{m}$ . Diploid species and the tetraploid *C. recurvata* had one terminal satellite, whereas the remaining tetraploid species showed four satellited chromosomes. Karyotypes were symmetrical. No  $\text{CMA}^-/\text{DAPI}^+$  bands were detected, but  $\text{CMA}^+/\text{DAPI}^-$  bands associated with NOR were always found. Pericentromeric heterochromatin was found in *C. recurvata*, *A. spiniflorum*, and the tetraploid cytotype of *M. patagonica*. The locations of the 18S-26S rDNA sites in all species coincided with  $\text{CMA}^+/\text{DAPI}^-$  bands; the same occurred with the sizes and numbers of signals for each species. This technique was applied for the first time in metaphase chromosomes in cacti. NOR-bearing pair no.1 may be homeologous in all species examined. In Cactaceae, the 18S-26S loci seem to be highly conserved.

Copyright © 2009 S. Karger AG, Basel

Fluorochrome banding (CMA/DAPI) and molecular cytogenetic methods (genomic and fluorescence in situ hybridization, GISH and FISH, respectively) have been applied to cytotaxonomical and evolutionary studies in different plant groups (e.g., Zhang and Sang, 1998; Adams et al., 2000; Schwarzacher, 2003).

The fluorochromes CMA and DAPI exhibit preferential staining for CG- and AT-rich DNA sequences, respectively, allowing the identification of different types of heterochromatin. This has been applied to chromosome analysis and has provided additional chromosome markers and informative characters for several Angiosperms (e.g., Moscone et al., 1996; Guerra, 2000; Souza and Benko-Iseppon, 2004; Gitaí et al., 2005; Urdampilleta et al., 2006).

More informative markers are often provided by FISH, a method that allows hybridization of known labeled marker sequences (probes) to homologous chromosomal targets (e.g., Adams et al., 2000; Schwarzacher and Heslop-Harrison, 2000; Schwarzacher, 2003). FISH enables the physical mapping of sequences to their location with-

Grants from 'Consejo Nacional de Investigaciones Científicas y Técnicas' (CONICET, Argentina), FONCYT, and 'Universidad Nacional de Córdoba' (SECyT, Argentina) are acknowledged. The 'Coordenação de Aperfeiçoamento de Pessoal de Nível Superior' (CAPES, Brazil) provided invaluable support.

in the genome, in particular repetitive sequences that cannot be mapped easily by any other method (Schwarzacher, 2003). Repetitive sequences change rapidly during evolution, providing excellent markers for the identification of chromosomes and chromosome segments, and for following evolutionary chromosome rearrangements. The 5S and 18S-26S rDNA genes, in particular, have been used extensively to establish possible chromosomal homeologies (e.g., Moscone et al., 1999; Taketa et al., 1999; Adams et al., 2000; Cai et al., 2006; Hasterok et al., 2006). Effectively, they were commonly used as molecular-cytogenetic markers because they are abundant and highly conserved in higher plants (Schwarzacher, 2003), e.g., *Hypochoeris* (Cerbah et al., 1998), *Rhynchospora* (Vanzela et al., 2003), *Plantago* (Dhar et al., 2006), *Citrus* (Moraes et al., 2007), and *Cephalanthera* (Moscone et al., 2007).

The 18S-26S rDNA associated with nucleolar organization regions (NOR) consists of tandem repeat units, comprising coding and internal transcribed spacers (ITS) (Appels and Honeycutt, 1986). The high copy number, rapid concerted evolution, and small size of the ITS regions make them informative in determining evolutionary relationships between closely related groups (Baldwin et al., 1995; Susanna et al., 1999; Ananthawat-Jonsson and Bodvardsdottir, 2001).

The family Cactaceae has been scarcely studied cytogenetically. Cactaceae is a relatively small family with about 130 genera and 1,500–1,800 species distributed in America, from southern Patagonia to Canada (Barthlott and Hunt, 1993; Anderson, 2001; Hunt et al., 2006). Morphological and molecular evidence suggest that the family is monophyletic (Wallace, 1994) and is composed of four subfamilies (Anderson, 2001; Hunt et al., 2006): Pereskioideae (monotypic with the genus *Pereskia*), Maihuenioideae (also monotypic with the genus *Maihuenia*), Opuntioideae (with 15 genera), and Cactoideae (the larger subfamily with around 110 genera).

The majority of cytological studies in cacti only provide chromosome counts, making clear that their basic chromosome number is  $x = 11$  (e.g., Ross, 1981; Pinkava et al., 1985; Parfitt, 1987; Cota and Philbrick, 1994; Cota and Wallace, 1995; Bandyopadhyay and Sharma, 2000; Arakaki et al., 2007). On the other hand, there are few detailed karyotypic studies available (Johnson, 1980; Palomino et al., 1988; Cota and Wallace, 1995), being more rare for South American members of the family (Das and Mohanty, 2006; Las Peñas et al., 2008). Regardless of the importance of the application of chromosome banding techniques, there is only one article on the spe-

cies of the exclusively Argentinean genus *Pyrrhocactus* (Las Peñas et al., 2008). There are no published reports on the use of FISH on somatic metaphase chromosomes and only one reported use of FISH on pollen mother cells to detect the ploidy status of the taxa studied (Tel-Zur et al., 2004).

Thus in this work we examined for the first time the karyotypes using CMA/DAPI chromosome banding, and FISH in six Argentinean species representing the four subfamilies of Cactaceae to correlate the banding pattern with the physical mapping of 18S-26S rDNA among the studied groups. In addition, we discuss these data in light of existing information on systematic and evolutionary relationships in Cactaceae.

## Material and methods

### *Plant material*

Collection data of the species studied are included in Table 1. Vouchers were deposited in the herbarium of the Museo Botánico de Córdoba (CORD). Specimens were planted in earthenware pots in an equal part mixture of sand and soil.

### *Karyotype analysis*

The preparation of metaphase chromosomes was done from adventitious roots pretreated with 2 mM 8-hydroxyquinolin for 8 h at 4°C and fixed in 3:1 ethanol:acetic acid. For slide preparation, root tips were hydrolyzed with 5 N HCl for 30 min at room temperature and then washed, stained with Feulgen for 2 h, and squashed in a drop of 2% acetic carmine (Jong, 1997). Permanent mounts were made following Bowen's method (Bowen, 1956). Ten metaphases per species were photographed with a phase contrast optic Zeiss Axiophot microscope (Jena, Germany) and a Leica DFC300FX camera (Wetzlar, Germany). Photographs were used to take measurements of the following features for each chromosome pair: s (short arm), l (long arm), and c (total chromosome length); the length of the satellite was added to the respective chromosome arm. The arm ratio ( $r = l/s$ ) was then calculated and used to classify the chromosomes as recognized by Levan et al. (1964). In addition, mean chromosome length (C), mean total haploid chromosome length of karyotype based on the mean chromosome length (tl), and mean arm ratio (R) were calculated. Idiograms were based on the mean values for each species. The chromosomes were arranged first into groups according to their increasing arm ratio and then according to decreasing length within each group. Karyotype asymmetry was estimated using the intrachromosomal ( $A_1$ ) and interchromosomal ( $A_2$ ) indices of Romero Zarco (1986).

### *Fluorochrome banding and FISH*

Root tips were washed twice in distilled water (10 min each), digested with a 2% cellulase (Sigma-Aldrich, Vienna, Austria), 20% pectinase (from *Aspergillus niger*; Sigma-Aldrich, Vienna, Austria) solution (45 min at 37°C), and squashed in a drop of 45% acetic acid (Schwarzacher et al., 1980). Only one root tip was used in each slide. After coverslip removal in liquid nitrogen, the slides were stored at -20°C.

**Table 1.** Cactaceae species studied (all from Argentina). Collection data: collector and number, province, locality, date, and in brackets number of individuals, number of cells studied, respectively.

Taxa	Collection data
Pereskioideae <i>Pereskia aculeata</i> Mill.	G. Barboza et al. 1036, Misiones, Guaraní, May-15-2004 (15, 15)
Maihuenioideae <i>Maihuenia patagonica</i> (Phil.) Britton and Rose	P. S. Steibel and H. Troiani, 16213, Rio Negro, Valcheta, December-08-2004 (20, 40), G. Barboza et al. 1247, Neuquen, Barrancas, February-05-2005 (10, 30)
Opuntioideae <i>Cumulopuntia recurvata</i> Guilmes and Thomas	M. L. Las Peñas 182, San Juan, Los Medanos, April-16-2005 (14, 20)
Cactoideae <i>Acanthocalycium spiniflorum</i> (K. Schum.) Backeb.	M. L. Las Peñas and D. Uñates 118, San Luis, Quines, November-30-2004 (5, 15)
<i>Echinopsis tubiflora</i> (Pfeiff.) Zucc.	M. L. Las Peñas and D. Uñates 278, Salta, Guachipas, January-20-2006 (7, 15)
<i>Trichocereus candicans</i> (Gillies ex Salm-Dyck) Britton and Rose	M. L. Las Peñas and D. Uñates 152, San Juan, Caucete, December-05-2005 (10, 25)

**CMA/DAPI banding:** Slides were stained with a drop of 0.5 mg/ml chromomycin A<sub>3</sub> (CMA) in McIlvaine buffer, pH 7.0 and distilled water (1:1) containing 2.5 mM MgCl<sub>2</sub> for 90 min and subsequently stained with 2 µg/ml 4'-6-diamidino-2-phenylindole (DAPI) (both Sigma-Aldrich, Vienna, Austria) for 30 min, and finally mounted in McIlvaine's buffer-glycerol v/v 1:1 (Schweizer, 1976; Schweizer and Ambros, 1994). The amount of heterochromatin was expressed as a percentage of the total length of the haploid karyotype.

**FISH:** This technique was applied for all species, except for the cytotype diploid of *Maihuenia patagonica* for which very few individuals and cells were available. The location and number of rDNA sites were determined by FISH using as probe the pTa71 containing the 18S-5.8S-26S rDNA (Gerlach and Bedbrook, 1979) labeled with biotin-14-dATP (BioNick, Invitrogen Carlsbad, USA). The FISH protocol was according to Schwarzacher and Heslop-Harrison (2000), with minor modifications. The preparations were incubated in 100 µg/ml RNase, post-fixed in 4% (w/v) paraformaldehyde, dehydrated in a 70–100% graded ethanol series, and air-dried. On each slide 30 µl of hybridization mixture was added (4–6 ng/µl of probe, 50% formamide, 10% dextran sulfate, 3.3 ng/µl of salmon DNA, 2× SSC and 0.3% SDS), previously denatured at 70°C for 10 min. Chromosome denaturation/hybridization was done at 90°C for 10 min, 48°C for 10 min, and 38°C for 5 min using a thermal cycler (Mastercycler, Eppendorf, Hamburg, Germany), and slides were placed in a humid chamber at 37°C overnight. The probe was detected with avidin-FITC conjugate and counterstained and mounted with 25 µl antifade (Vectashield Vector Lab., Burlingame, USA), containing 1 µl propidium iodide. Photomicrography was done with a BX51 Olympus photosystem (Tokyo, Japan) coupled with Evolution MIT CCD using Image ProPlus v4.5 software (Maryland, USA) for image capture.

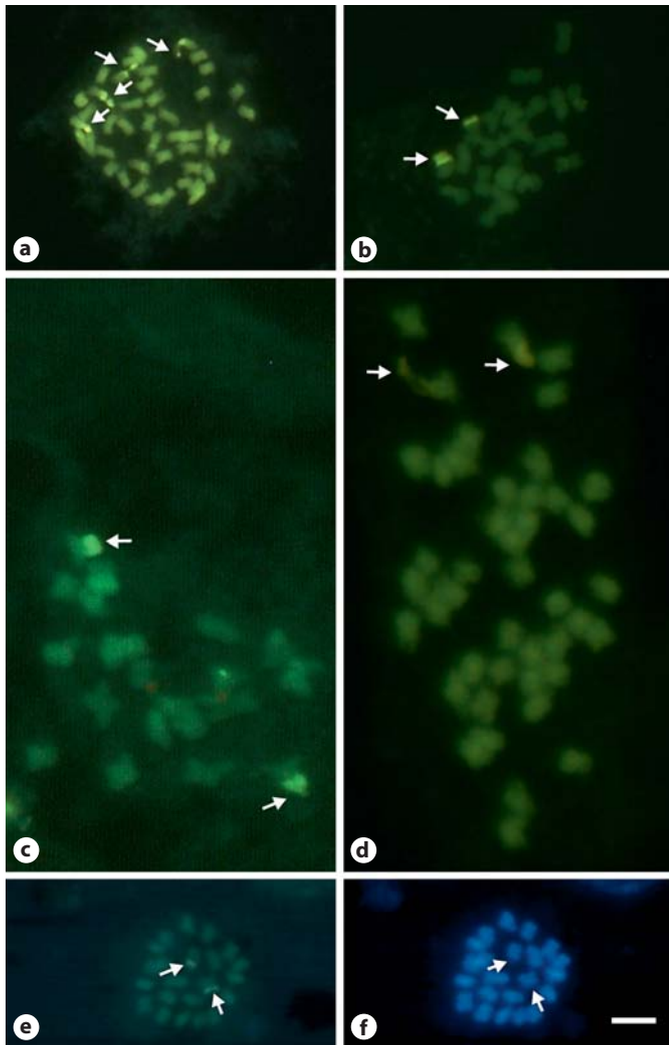
## Results

### Karyotypes

*Pereskia aculeata*, *Echinopsis tubiflora*, and *Acanthocalycium spiniflorum* were diploid (2n = 22), whereas *Cumulopuntia recurvata* and *Trichocereus candicans* were tetraploid (2n = 44), in all cells examined (Figs. 1, 2). On the other hand, the two populations of *Maihuenia patagonica* studied showed different ploidy levels: diploid from Neuquen province (2n = 22) and tetraploid from Rio Negro province (2n = 44) (Fig. 1a, e).

In general, the chromosomes were small, 2.3 µm being the average chromosome length for all taxa (Table 2). The karyotypes of the diploid species had one terminal satellite on the short arms of pair no.1, as also occurs in the tetraploid *Cumulopuntia recurvata* (Figs. 1, 3). The remaining tetraploid species showed four satellited chromosomes on the short arms of the longest *m* chromosomes (Figs. 1, 3). The frequency of appearance of the satellites in both homologues reached 70% of the examined cells in all taxa.

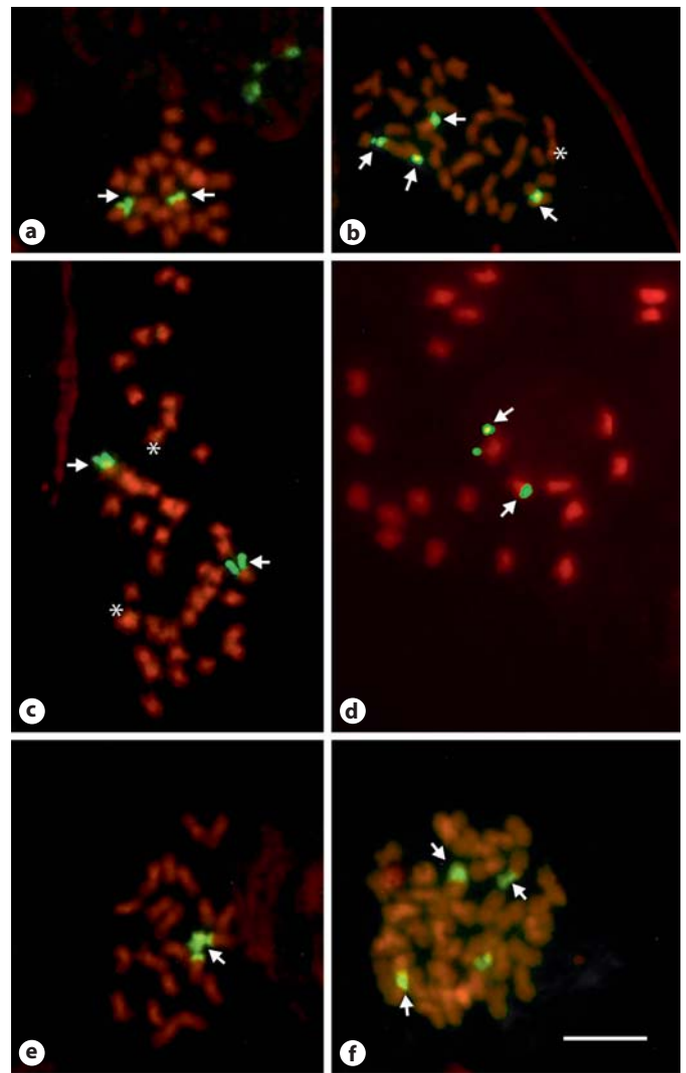
Karyotypes were symmetrical, considering both centromere position (most are *m* with 1 or 2, exceptionally 4 *sm* pairs) and chromosome size variation (there are slight differences among the chromosomes of the complement) (Table 2, Fig. 3). This is expressed by the asymmetry indices: A<sub>1</sub> ranges from 0.16 to 0.31 and A<sub>2</sub> from 0.14 to 0.26 (Table 2).



**Fig. 1.** Somatic metaphases with fluorochrome banding. (a) *Maihuenia patagonica* ( $2n = 44$ ). (b) *Acanthocalycium spiniflorum* ( $2n = 22$ ). (c) *Pereskia aculeata* ( $2n = 22$ ). (d) *Cumulopuntia recurvata* ( $2n = 44$ ). (e-f) *Maihuenia patagonica* ( $2n = 22$ ). (a-e) CMA fluorescence. (f) DAPI fluorescence. Arrows indicate CMA<sup>+</sup>/DAPI<sup>-</sup> NOR-associated heterochromatin. Bar = 5  $\mu\text{m}$ .

#### Chromosome banding

The banding patterns always showed CMA<sup>+</sup>/DAPI<sup>-</sup> constitutive heterochromatin associated with NOR (CG-rich). On the other hand, no CMA<sup>-</sup>/DAPI<sup>+</sup> bands were detected in any species. The diploids *Echinopsis tubiflora*, *Pereskia aculeata*, *Acanthocalycium spiniflorum*, and *Maihuenia patagonica* (Neuquen population), as well as the tetraploid *Cumulopuntia recurvata*, showed a NOR-associated heterochromatin region in the satellited chro-



**Fig. 2.** Somatic chromosomes detected by FISH using 18S-26S rDNA probe. (a) *Pereskia aculeata*. (b) *Maihuenia patagonica* ( $2n = 44$ ). (c) *Cumulopuntia recurvata*. (d) *Acanthocalycium spiniflorum*. (e) *Echinopsis tubiflora*. (f) *Trichocereus candicans*. Arrows show 18S-26S rDNA sites and asterisks indicate hybridization in pericentromeric region. Bar = 5  $\mu\text{m}$ .

mosome pair no.1. *Maihuenia patagonica* (Rio Negro population) and *Trichocereus candicans*, both tetraploid, had four NOR-associated heterochromatin regions (Fig. 1).

Additional CMA<sup>+</sup>/DAPI<sup>-</sup> pericentromeric bands were found in different numbers: four bands in two *m* pairs in *Cumulopuntia recurvata* and two bands in one *m* pair in *Acanthocalycium spiniflorum* and the tetraploid cytotype of *Maihuenia patagonica* (Figs. 1, 3). The highest and low-

**Table 2.** Karyotype data of Cactaceae. Abbreviations = tl: mean total haploid karyotype length; C: mean chromosome length; A<sub>1</sub>: mean intrachromosomal asymmetry index; A<sub>2</sub>: mean interchromosomal asymmetry index, %: amount of CMA<sup>+</sup>/DAPI<sup>-</sup> heterochromatin expressed as percentage of the haploid karyotype length; Lengths are in  $\mu\text{m}$ . Chromosome nomenclature after Levan et al. (1964). An asterisk indicates that the first chromosome pair has satellites and two asterisks two satellited pairs (# 1 and 2).

Taxa	2n	Haploid karyotype formula	tl	C	A <sub>1</sub>	A <sub>2</sub>	%
Pereskioideae							
<i>Pereskia aculeata</i>	22	10 m* + 1 sm	28.80	2.60	0.16	0.24	4.96
Maihuenioideae							
<i>Maihuenia patagonica</i>	22	9 m* + 2 sm	29.71	2.47	0.27	0.26	3.30
	44	18 m** + 4 sm	56.00	2.45	0.30	0.26	6.20
Opuntioideae							
<i>Cumulopuntia recurvata</i>	44	20 m* + 2 sm	39.50	1.80	0.23	0.20	3.10
Cactoideae							
<i>Acantocalycium spiniflorum</i>	22	10 m* + 1 sm	28.00	2.55	0.17	0.14	5.10
<i>Echinopsis tubiflora</i>	22	10 m* + 1 sm	26.90	2.40	0.20	0.25	4.62
<i>Trichocereus candicans</i>	44	22 m**	45.90	2.10	0.25	0.14	4.82

est total amounts of CMA<sup>+</sup>/DAPI<sup>-</sup> heterochromatin were found in two tetraploid taxa: *Cumulopuntia recurvata* with 3.1% and Rio Negro population of *Maihuenia patagonica* with 6.2% (Table 2).

#### Chromosomal mapping of the 18S-26S rDNA by FISH

The locations of the 18S-26S rDNA sites in all species studied coincided with CMA<sup>+</sup>/DAPI<sup>-</sup> bands described above (Figs. 2, 3), i.e., they are located on the secondary constrictions and the adjacent satellites at telomeric positions. The same occurred with the sizes and numbers of signals for each species.

#### Discussion

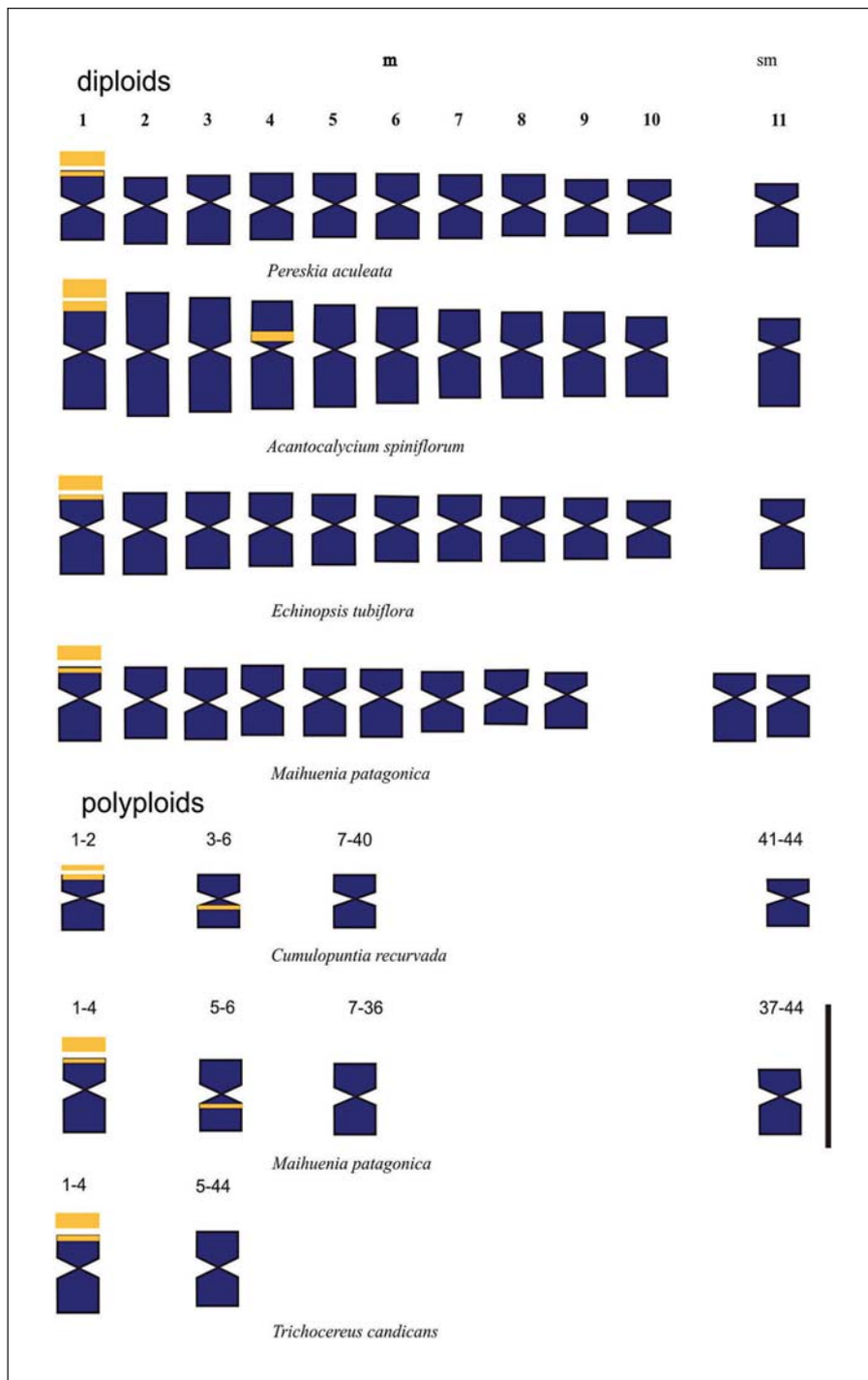
The basic chromosome number for the family is  $x = 11$  (e.g., Pinkava et al., 1977, 1985, 1992; Parfitt, 1987; Cota and Wallace, 1995; Bandyopadhyay and Sharma, 2000; Das and Mohanty, 2006). It was found in all taxa here, examined at the diploid or tetraploid levels.

Most numbers are here reported for the first time. The exceptions are: *P. aculeata* for which we confirmed previous reports (Leuenberger, 1986; Lombello and Forni-Martins, 1998) and *M. patagonica* for which we found a diploid population, as registered by Leuenberger (1997), as well as the novelty of a tetraploid population, a fact known for several species (Johnson, 1980; Pinkava, 2002). This tetraploid cytotype of *M. patagonica* is the first report of polyploidy in a member of subfamily Maihuenioi-

deae. Polyploidy has been reported for ca. 25% of the cacti investigated so far (cf. Palomino et al., 1988; Cota and Wallace, 1995; Pinkava, 2002; Arakaki et al., 2007); particularly, it seems to have played an important role in the evolution of Opuntioideae where it reaches a percentage of 64%.

Although variation in karyotype composition regarding chromosome size and morphology was found, the taxa here studied and previous data show that Cactaceae has symmetrical karyotypes mostly composed of *m* and *sm* chromosomes (e.g., Pinkava et al., 1985, 1992; Parfitt, 1987; Cota and Wallace, 1995; Bandyopadhyay and Sharma, 2000; Das and Mohanty, 2006; Las Peñas et al., 2008). *St* chromosomes are rare and only reported for two species (Johnson, 1980; Las Peñas et al., 2008), whereas *t* chromosomes have never been registered.

CMA/DAPI banding has been widely used in flowering plants to characterize heterochromatin bands with respect to their highly repeated DNA composition, known also as satellite DNA. The numbers of bands and the heterochromatin amount varied, but the general patterns were relatively conserved at the generic level (Galasso et al., 1996; Moscone et al., 1996; Miranda et al., 1997; Ran et al., 1999; Marcon and Guerra, 2005; Fregonezi et al., 2006). There is a positive association between the ploidy level and the amount of heterochromatin in the cytotypes of *Maihuenia patagonica*. The scarce data available in Cactaceae are reduced to seven *Pyrrhocactus* species (Las Peñas et al., 2008) and our data on members of the four subfamilies. Both data sets showed that CMA<sup>+</sup>/DAPI<sup>-</sup>



**Fig. 3.** Idiograms of Cactaceae studied. Blue blocks = euchromatin, yellow blocks = CMA<sup>+</sup>/18S-26S rDNA sites. Bar = 5 μm.

heterochromatin blocks were always present on a terminal position associated with secondary constrictions; on the other hand, there is variability in the presence and the number of pericentromeric bands, which may be useful for evolutionary and systematic purposes.

Physical mapping by FISH of 18S-26S rDNA were here performed for the first time in metaphase chromosomes in cacti. FISH signals with pTa71 do not exhibit variation in size among the different species studied. The presence of these sites in terminal regions of both chromosome

arms is frequent in Angiosperms (e.g., Zhang and Sang, 1998; Benko-Iseppon and Morawetz, 2000; Vaio et al., 2005; Urdampilleta et al., 2006).

The number of 18S-26S rDNA sites in most species studied varied from two in the diploids to four in the tetraploids; thus, they may be indicative of the ploidy level. The same numbers of sites were found on pollen mother cells to detect the ploidy level in species and hybrids of *Hylocereus* and *Selenicereus* (Tel-Zur et al., 2004). Nevertheless, the tetraploid *C. recurvata* here analyzed showed signals in only one pair. This reduction in the locus number of 18S-26S rDNA units in several polyploids has been reported for other plant species (Appels et al., 1980; Hanson et al., 1996; Adachi et al., 1997; Vanzela et al., 2003). Reduction in 18S-26S rRNA gene locus number has previously been attributed to diploidizing (Melo and Guerra, 2003; Leitch and Bennett, 2004).

In many Angiosperm species, the centromere is associated with blocks of heterochromatin that contain a core of tandem satellite repeats, many of which are species specific and show chromosome-specific variants (Heslop-Harrison et al., 2003; Schwarzacher, 2003). Diverse mechanisms have been postulated to account for the variation in size, number, and position of rDNA sites and their apparent mobility, i.e., chromosome rearrangements, homologous and nonhomologous unequal crossing-over, gene conversion, transpositional events (Schubert and Wobus, 1985; Leitch and Heslop-Harrison, 1992; Hanson et al., 1996). These processes could act alone or in combination and do not necessarily imply changes in overall chromosome morphology (Hall and Parker, 1995). In this study, pericentromeric fluorescence signals were less intense than terminal signals. A possible hypothesis explaining the presence of these pericentric sites might be that they originated as active, subtelomeric 18S-26S rRNA genes that were subsequently shifted to the centromeric region through an inversion mechanism; nevertheless, information based on additional techniques, as AgNOR staining or FISH using IGS as probe, are needed to confirm it. Frequently, intergenic spacers could organize other chromosome regions (Hemleben et al., 2007), e.g., the satellite DNA sequences of some species of *Vigna* (Fabaceae) apparently originated from subrepeats of the intergenic spacer of rDNA (Unfried et al., 1991; Macas et al., 2003). In *Nicotiana* and *Solanum* (Solanaceae), independent satellites highly homologous to rDNA intergenic spacers were observed in terminal and pericentromeric regions, respectively (Stupard et al., 2002; Lim et al., 2004), and the interspecific variation indicated a dynamic nature of the repetitive DNA.

Both fluorochrome banding and rDNA gene mapping have been valuable in the identification of homeologous chromosome pairs among plant species (e.g., Moscone et al., 1995, 1996, 2007). In Cactaceae, NOR-bearing pair no.1 may be homeologous in all species examined so far (Las Peñas et al., 2008, this study). Although more data are needed, in Cactaceae the 18S-26S loci seem to be highly conserved as detected in members of its four subfamilies. This NOR-associated heterochromatin (18S-26S rDNA locus) is the rule in plants as a whole (Sinclair and Brown, 1971; Morawetz, 1986; Benko-Iseppon and Morawetz, 2000; Urdampilleta et al., 2006).

In general, karyotype features of cacti allowed individual species to be distinguished. Chromosome variation, although not always large, accompanied evolutionary divergence of the species. In addition, our comparative study of chromosome banding patterns and data on *Pyrrhocactus* (Las Peñas et al., 2008) suggested that repeated DNA segments also contributed to karyotypic differentiation. On the other hand, our data on the 18S-26S rDNA locus implied that its physical position seems to be conserved in the family. In summary, more data are needed using these techniques in cacti, as they provided useful information in the cytogenetics of these peculiar plants.

### Acknowledgements

Roberto Kiesling (CRICYT, Mendoza) kindly identified the samples.

### References

- Adachi J, Watanabe K, Fukui K, Ohmido N, Kosuge K: Chromosomal location and reorganization of the 45S and 5S rDNA in the *Brachyscome lineariloba* complex (Asteraceae). *J Plant Res* 110:371–377 (1997).
- Adams SP, Leitch IJ, Bennett MD, Chase MW, Leitch AR: Ribosomal DNA evolution and phylogeny in *Aloe* (Asphodelaceae). *Am J Bot* 87:1578–1583 (2000).
- Anamthawat-Jonsson K, Bodvarsdottir SK: Genomic and genetic relationships among species of *Leymus* (Poaceae: Triticeae) inferred from 18S-26S ribosomal genes. *Am J Bot* 88:553–559 (2001).
- Anderson EF: *The Cactus Family* (Timber Press, Portland 2001).
- Appels R, Honeycutt RL: rDNA: evolution over a billion years, in Dutta SK (ed): *DNA Systematics*, pp 81–125 (CRC Press, Boca Raton 1986).
- Appels R, Gerlach WL, Dennis ES, Swift T, Peacock WJ: Molecular and chromosomal organization of DNA sequences coding for the ribosomal RNAs in cereals. *Chromosoma* 78:293–311 (1980).

- Arakaki M, Soltis DE, Speranza P: New chromosome counts and evidence of polyploidy in *Haageocereus* and related genera in tribe Trichocereae. *Brittonia* 59:290–297 (2007).
- Baldwin BG, Sanderson MJ, Porter JM, Wojciechowski MF, Campbell CS, Donoghue MJ: The ITS region of nuclear ribosomal DNA: a valuable source of evidence on angiosperm phylogeny. *Ann Missouri Bot Gard* 82:247–277 (1995).
- Bandyopadhyay B, Sharma A: The use of multivariate analysis of karyotypes to determine relationships between species of *Opuntia*. *Caryologia* 53:121–126 (2000).
- Barthlott W, Hunt D: Cactaceae, in Kubitzki K, Rohwer JG, Bittrich V (eds): *The Families and Genera of Vascular Plants. Vol. II. Flowering Plants Dicotyledons*, pp 161–197 (Springer, Berlin 1993).
- Benko-Iseppon AM, Morawetz W: Cytological comparison of Calyceraceae and Dipsacaceae with special reference to their taxonomic relationships. *Cytologia* 65:123–128 (2000).
- Bowen C: Freezing by liquid carbon dioxide in making slides permanent. *Stain Technol* 31:90 (1956).
- Cai Q, Zhang D, Liu ZL, Wang XR: Chromosomal localization of 5S and 18S rDNA in five species of subgenus *Strobos* and their implications for genome evolution of *Pinus*. *Ann Bot* 97:715–722 (2006).
- Cerbah M, Coulaud J, Siljak-Yakovlev S: rDNA organization and evolutionary relationships in the genus *Hypochaeris* (Asteraceae). *J Hered* 89:312–318 (1998).
- Cota JH, Philbrick CT: Chromosome number variation and polyploidy in the genus *Echinocereus* (Cactaceae). *Am J Bot* 81:1054–1062 (1994).
- Cota JH, Wallace RS: Karyotypic studies in the *Echinocereus* (Cactaceae) and their taxonomic significance. *Caryologia* 48:105–122 (1995).
- Das AB, Mohanty S: Karyotype analysis and in situ nuclear DNA content in seven species of *Echinopsis* Zucc. of the family Cactaceae. *Cytologia* 71:75–79 (2006).
- Dhar Mk, Bern F, Kaul S, Gill B: Characterization and physical mapping of ribosomal RNA gene families in *Plantago*. *Ann Bot* 97:541–548 (2006).
- Fregonezi JN, Fernandes T, Domingues Torezan JM, Vieira O, Vanzela ALL: Karyotype differentiation of four *Cestrum* species (Solanaeae) based on the physical mapping of repetitive DNA. *Genet Mol Biol* 29:97–104 (2006).
- Galasso I, Saponetti LS, Pignone D: Cytotaxonomic in *Vigna* III. Chromosomal distribution and reacting properties of the heterochromatin in five wild species of the section *Vigna*. *Caryologia* 49:311–319 (1996).
- Gerlach WL, Bedbrook JL: Cloning and characterization of ribosomal RNA genes from wheat and barley. *Nucleic Acids Res* 7:1869–1885 (1979).
- Gitai J, Horres R, Benko-Iseppon AM: Chromosomal features and evolution of Bromeliaceae. *Plant Syst Evol* 253:65–80 (2005).
- Guerra M: Patterns of heterochromatin distribution in plant chromosomes. *Genet Mol Biol* 23:1029–1041 (2000).
- Hall KJ, Parker JS: Stable chromosome fission associated with rDNA mobility. *Chromosome Res* 3:417–422 (1995).
- Hanson RE, Islan-Faridi MN, Persival EA, Crane CF: Distribution of 5S and 18S-28S rDNA loci in a tetraploid cotton (*Gossypium hirsutum* L.) and its putative diploid ancestors. *Chromosome Res* 10:55–61 (1996).
- Hasterok R, Wolny E, Hosiawa M: Comparative analysis of rDNA distribution in chromosomes of various species of Brassicaceae. *Ann Bot* 97:205–216 (2006).
- Hemleben V, Kovarik A, Torres-Ruiz RA, Volkov RA, Beridz RA: Plant highly repeated satellite DNA: molecular evolution, distribution and use for identification of hybrids. *Syst Biodivers* 5:277–289 (2007).
- Heslop-Harrison JS, Brandes A, Schwarzacher T: Tandemly repeated DNA sequences and centromeric chromosomal regions of *Arabidopsis* species. *Chromosome Res* 11:241–253 (2003).
- Hunt D, Taylor N, Charles G: *The New Cactus Lexicon* (DH books, Milborne Port 2006).
- Johnson MA: Further cytological investigations in *Mammillaria prolifera* and other *Mammillaria* species. *Cact Succ J Gr Brit* 42:43–47 (1980).
- Jong J: *Laboratory Manual of Plant Cytological Techniques* (Royal Botanical Garden, Edinburgh 1997).
- Las Peñas ML, Bernardello G, Kiesling R: Karyotypes and fluorescent chromosome banding in *Pyrrhocactus* (Cactaceae). *Plant Syst Evol* 272:211–222 (2008).
- Leitch AR, Heslop-Harrison JS: Physical mapping of the 18S-5.8S-26S rRNA genes in barley by in situ hybridization. *Genome* 35:1013–1018 (1992).
- Leitch IJ, Bennett MD: Genome downsizing in polyploid plants. *Bot J Linn Soc* 82:651–663 (2004).
- Leuenberger BE: *Pereskia* (Cactaceae). *Mem New York Bot Gard* 41:1–141 (1986).
- Leuenberger BE: *Maihuenia* – monograph of a Patagonian genus of Cactaceae. *Bot Jahrb Syst* 119:1–92 (1997).
- Levan A, Sandberg A, Fredga K: Nomenclature for centromeric position on chromosomes. *Hereditas* 52:201–220 (1964).
- Lim KY, Skalicka K, Koukalova B, Volkov RA, Matyasek R, et al: Dynamic changes in the distribution of a satellite homologous to intergenic 26-18S rDNA spacer in the evolution of *Nicotiana*. *Genetics* 166:1935–1946 (2004).
- Lombello RA, Forni-Martins ER: Cytological studies in climbers of Brazilian forest reserve. *Cytologia* 63:415–420 (1998).
- Macas J, Navrátilová A, Meiszáros T: Sequence subfamilies of satellite repeats related to rDNA intergenic spacer are differentially amplified on *Vicia sativa* chromosomes. *Chromosoma* 112:152–158 (2003).
- Marcon BA, Guerra M: Variation in chromosome numbers, CMA bands and 45S rDNA sites in species of *Selaginella* (Pteridophyta). *Ann Bot* 95:271–276 (2005).
- Melo NF, Guerra M: Variability of the 5S and 45S rDNA sites in *Passiflora* L. species with distinct base chromosome numbers. *Ann Bot* 92:309–316 (2003).
- Miranda M, Ikeda F, Endo T, Morigucki T, Omura M: Comparative analysis on the distribution of heterochromatin in *Citrus*, *Poncirus* and *Fortunella* chromosomes. *Chromosome Res* 5:86–92 (1997).
- Moraes AP, Santos Soares Filho W, Guerra M: Karyotype diversity and the origin of grapefruit. *Chromosome Res* 15:115–121 (2007).
- Morawetz W: Remarks on karyological differentiation patterns in tropical woody plants. *Plant Syst Evol* 152:49–100 (1986).
- Moscone EA, Loidl J, Ehrendorfer F, Hunziker AT: Analysis of active nucleolus organizing regions in *Capsicum* (Solanaceae) by silver staining. *Am J Bot* 82:276–287 (1995).
- Moscone EA, Lambrou M, Ehrendorfer F: Fluorescent chromosome banding in the cultivated species of *Capsicum* (Solanaceae). *Plant Syst Evol* 202:37–63 (1996).
- Moscone EA, Klein F, Lambrou M, Fuchs J, Schweizer D: Quantitative karyotyping and dual-color FISH mapping of 5S and 18S-25S rDNA probes in the cultivated *Phaseolus* species (Leguminosae). *Genome* 42:1224–1233 (1999).
- Moscone EA, Samuel R, Schwarzacher T, Schweizer D: Complex rearrangements are involved in *Cephalanthera* (Orchidaceae) chromosome evolution. *Chromosome Res* 15:931–943 (2007).
- Palomino G, Socorro ZL, Scheinvar L: Estudios citogenéticos de dos especies y una variedad del género *Nyctocereus* (Cactaceae). *Bol Soc Bot México* 48:75–80 (1988).
- Parfitt BD: *Echinocereus nicholii* (L. Benson) Cactaceae. *Phytologia* 63:157–158 (1987).
- Pinkava DJ: On the evolution of the North American Opuntioideae, in Hunt D, Taylor N (eds): *Studies in the Opuntioideae*, pp 78–99 (Royal Botanic Gardens, Kew 2002).
- Pinkava DJ, McGill LA, Reeves T: Chromosome number in some cacti of western North America. *Bull Torrey Bot Club* 104:105–110 (1977).
- Pinkava DJ, Baker MA, Parfitt BD: Chromosome number in some cacti of western North America V. *Syst Bot* 10:471–483 (1985).
- Pinkava DJ, Parfitt BD, Baker MA, Worthington RD: Chromosome numbers in some cacti of western North America VI. *Madroño* 39:98–113 (1992).

- Ran Y, Murray BG, Hammett KRW: Karyotype analysis of the genus *Clivia* by Giemsa and fluorochrome banding and in situ hybridization. *Euphytica* 106:139–147 (1999).
- Romero Zarco C: A new method for estimating karyotype asymmetry. *Taxon* 35:556–530 (1986).
- Ross R: Chromosome counts, cytology, and reproduction in the Cactaceae. *Am J Bot* 68: 463–470 (1981).
- Schubert I, Wobus U: In situ hybridization confirms jumping nucleolus organizing regions in *Allium*. *Chromosoma* 92:143–148 (1985).
- Schwarzacher T: DNA, chromosomes, and in situ Hybridization. *Genome* 46:953–962 (2003).
- Schwarzacher T, Heslop-Harrison P: Practical in situ Hybridization (Bios Scientific Publishers Limited, Oxford 2000).
- Schwarzacher T, Ambros P, Schweizer D: Application of Giemsa banding to orchid karyotype analysis. *Plant Syst Evol* 134:293–297 (1980).
- Schweizer D: Reverse fluorescent chromosome banding with chromomycin and DAPI. *Chromosoma* 58:307–324 (1976).
- Schweizer D, Ambros P: Chromosome banding, in Gosden JR (ed): *Methods in Molecular Biology. Chromosome Analysis Protocols* (Humana Press, Totowa 1994).
- Sinclair JH, Brown DD: Retention of common nucleotide sequences in the ribosomal deoxyribonucleic acid of eukaryotes and some of their physical characteristics. *Biochemistry* 10:10–20 (1971).
- Souza M, Benko-Iseppon AN: Cytogenetics and chromosome banding patterns in Caesalpinioideae and Papilionioideae species of Pará, Amazonas, Brazil. *Bot J Linn Soc* 144: 181–191 (2004).
- Stupard J, Song J, Tek AL, Cheng Z, Dong F, Jiang J: Highly condensed potato pericentromeric heterochromatin contains rDNA-related tandem repeats. *Genetics* 162:1435–1444 (2002).
- Susanna A, Garnatje T, Garcia-Jacas N: Molecular phylogeny of *Cheirolophus* (Asteraceae: Cardueae-Centaureinae) based on ITS sequences of nuclear ribosomal DNA. *Plant Syst Evol* 214:147–160 (1999).
- Taketa S, Harrison G, Heslop-Harrison JS: Comparative physical mapping of the 5S and 18S-26S in nine wild *Hordeum* species and cytotypes. *Theor Appl Genet* 98:1–9 (1999).
- Tel-Zur N, Abbo S, Bar-Zvi D, Mizrahi Y: Genetic relationships among *Hylocereus* and *Selenicereus* vine cacti (Cactaceae): Evidence from hybridization and cytological studies. *Ann Bot* 94:527–534 (2004).
- Unfried K, Schiebel K, Hemleben V: Subrepeats of rDNA intergenic spacer present as prominent independent satellite DNA in *Vigna radiata* but not in *Vigna angularis*. *Gene* 99: 63–68 (1991).
- Urdampilleta JD, Ferrucci MS, Torezan JMD, Vanzela ALL: Karyotype relationships among four South American species of *Urvillea* (Sapindaceae; Paullinieae). *Plant Syst Evol* 258:85–95 (2006).
- Vaio M, Speranza P, Valls JFM, Guerra M, Mazzella C: Localization of the 5S and 45S rDNA sites and cpDNA sequence analysis in species of the *Quadrifaria* group of *Paspalum* (Poaceae, Paniceae). *Ann Bot* 96:191–200 (2005).
- Vanzela ALL, Cuadrado A, Guerra M: Localization of 45S rDNA and telomeric sites on holocentric chromosomes of *Rhynchospora tenuis* Link (Cyperaceae). *Genet Mol Biol* 26: 199–201 (2003).
- Wallace RS: Phylogenetic analysis. *Succ Plant Res* 1:125–130 (1994).
- Zhang D, Sang T: Chromosomal structural rearrangement of *Paeonia brownii* and *P. californica* revealed by fluorescence in situ hybridization. *Genome* 41:848–853 (1998).