

Cytogenetics, morphology and viability of *Borreria spinosa* (Rubiaceae) pollen grains

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ABSTRACT. Introduction: *Borreria spinosa* is a herbaceous plant considered a weed in agricultural crops, although it also has potential in apiculture. Despite its agronomic importance, its cytogenetic and palynological traits are poorly known. **Objective:** To do the cytogenetic characterization of *B. spinosa*, describe the morphology of its pollen grains, and estimate their potential viability. **Methods:** We collected plant material in Santiago del Estero, Argentina, in May 2017 and used classical methods (8-hydroxyquinoline and Farmer for mitosis and meiosis; viability with Müntzing; pollen morphology with nonacetolysed and acetolysed techniques, and SEM). **Results:** *B. spinosa* has a chromosome number of $2n = 56$, with a karyotype of $46 m + 10 sm$. Meiotic abnormalities mainly include chromosomes outside the equatorial plate, lagging chromosomes, and cytomixis. Pollen grains are small to medium-sized, suboblate to oblate-spheroidal, with a viability of 70%. This is a tetraploid species, most chromosomes are metacentric or submetacentric; and it has irregular meiotic behavior and cytomixis, with concomitant low viability in pollen grains. **Conclusion:** In *B. spinosa*, pollen grains have similar characteristics to other species of the genus *Borreria*.

Keywords: Cytomixis, karyotype, microsporogenesis, pollen morphology, weeds, palynology.

RESUMEN. “Citogenética, morfología y viabilidad de los granos de polen de *Borreria spinosa* (Rubiaceae)”. **Introducción:** *Borreria spinosa* es una planta herbácea considerada maleza en cultivos agrícolas, aunque también presenta potencial en la producción apícola. A pesar de su importancia agronómica, sus rasgos citogenéticos y palinológicos son poco conocidos. **Objetivo:** Caracterizar citogenéticamente a *B. spinosa*, describir la morfología de sus granos de polen, y estimar su viabilidad potencial. **Métodos:** Recolectamos material vegetal en Santiago del Estero, Argentina, en mayo de 2017 y utilizamos métodos clásicos (8-hidroxiquinolina y Farmer para mitosis y meiosis; viabilidad con Müntzing; morfología del polen con técnicas no acetolizadas y acetolizadas, y MEB). **Resultados:** *B. spinosa* tiene un número cromosómico de $2n = 56$, con un cariotipo de $46 m + 10 sm$. Las anomalías meióticas incluyen principalmente cromosomas fuera de la placa ecuatorial, cromosomas rezagados y citomixis. Los granos de polen son de pequeños a medianos, suboblato a oblato-esferoidales, con viabilidad del 70%. Es tetraploide, la mayoría de los cromosomas son metacéntricos o submetacéntricos; y tiene comportamiento meiótico irregular y citomixis, con baja viabilidad concomitante del polen. **Conclusión:** En *B. spinosa*, los granos de polen tienen características similares a otras especies del género *Borreria*.

Palabras clave: Citomixis, cariotipo, microesporogénesis, morfología polínica, malezas, palinología.

Rubiaceae Juss is the fourth largest angiosperm family, consisting of approximately 611 genera, with herbaceous and woody arboreal species that have flowers adapted to different pollinators (The Angiosperm Phylogeny Group et al., 2016). Globally, it is distributed in tropical and subtropical regions, and 30% of its species are found in Latin America (Dutra et al., 2020). Within this family, the genera *Borreria* G. stands out with approximately 80 species, distributed from the south of the United States to the center of Argentina and Uruguay and some adventitious species in Africa, Asia and Oceania (Miguel et al., 2022). In South America, 30 species are recorded, which grow preferably in open fields with sandy or lateritic soils and less frequently in humid rocky environments with shrubby or larger elements. (Zuloaga et al., 2019).

Borreria spinosa Cham. & Schltdl. ex. DC., is an herbaceous or sub shrubby plant, 0.1 to 1 meter high. It has a tetragonal, glabrous or pubescent stem. Leaves are pseudo verticillate due to the presence of brachyblasts, sessile to pseudo speciolate. It has elliptic leaves, acute apex, attenuated base, with glabrescent abaxial and adaxial faces, sometimes with papillae of thick base on the abaxial face. It also has terminal and axillary bilateral inflorescences, with one, two or five glomeruli per branch, hemispherical. It is characterized by oblongoid capsules, brown, hairy in the upper third, dehiscent in two valves united at the base. Seeds are oblongoid to subcylindrical and brown (Nepomuceno et al., 2018).

In Argentina, *B. spinosa* has been reported as a weed of agricultural crops that covers a wide region between the provinces of Salta and Buenos Aires and stands out for its low sensitivity to glyphosate (Luna & Druetta, 2018). Despite this, Scandaliaris et al., (2020) mentioned the apicultural potential of this species in the province of Córdoba. Salgado et al., (2014) reported the presence of traces of pollen of species of the genus *Borreria* in honeys from the province of Chaco. The presence of pollen of species of this genus has also been observed very frequently in honeys from northeastern Brazil (de Borges, 2020; Silva & Novais, 2020). These reports suggest the apicultural potential of *B. spinosa*.

In *Borreria* three ploidy levels 2x, 4x and 6x were recognized (Daviña & Cabral, 1991) and for the genus sporophytic and gametophytic chromosome counts of $2n=8$; 40; 64 - 56 and $n=14$; 26–27; 28 were recorded, respectively (Goldblatt & Johnson, 1990).

There are no references on the cytogenetic characteristics and palynological studies of *B. spinosa*. The first aspect contributes to the understanding of its reproductive biology, of great relevance in weed control. Palynology studies have been very useful in issuing allergy risk warnings, in honey production certification and in establishing taxonomic relationships (Marcos et al., 2015).

The aim of this work was to do the cytogenetic characterization of *B. spinosa*, describe the morphology of its pollen grains and estimate the potential viability of these grains.

MATERIALS AND METHODS

Study area and plant material: The collection of adult plants and seeds of *B. spinosa* was performed in the locality El Zanjón, province of Santiago del Estero, Argentina (27°55'00" S, 64°15'00" W), in May 2017. The village is located in the phytogeographic region of Western Chaco and has subtropical climate with dry season. The average annual rainfall is 579mm, and the average annual temperature is 19,8°C (Landi et al., 2021).

The reference material was deposited in the herbarium of the Facultad de Agronomía y Agroindustrias de la Universidad Nacional de Santiago del Estero (Fig. 1).



Fig. 1. *Borreria spinosa*. Argentina. Province of Santiago del Estero

Characterization of mitosis: The roots used were obtained from seeds germinated in a humid chamber at 35°C (daytime) and 20°C (nighttime), with a 12-hour photoperiod. The roots were pretreated in 0,002M 8-hydroxyquinoline for 8 h at room temperature. They were then fixed in Farmer's solution (ethyl alcohol-acetic acid, 3:1) for 24 h and preserved in 70° ethanol at 4°C until use. The material was hydrolyzed in 1N HCl at 60°C for 15 min, and transient preparations were made and stained with a drop of 2% propionic hematoxylin. Observations were made under a ZeissAxio LAB.A1 optical microscope, Olympus, and chromosome length measurements were performed using MicroMeasure 3.3 software (Reeves, 2001).

The following variables were evaluated: total chromosome length (c), long arm length (l), short arm length (s), centromeric index (ci), total complement length (TCL), mean chromosome length (cme), maximum and minimum chromosome length (c max, c min). Chromosomes were classified according to the methodology proposed by Levan et al., (1964), inter- and intrachromosomal asymmetries were calculated according to Romero (1986), and the criterion of Bataglia (1955) was used for the description of satellites. For the elaboration of the karyotype and idiogram, seven metaphase plates with well extended chromosomes were considered.

Characterization of meiosis and pollen viability: Flower buds were immersed in Farmer's solution (ethanol-acetic acid 3:1) for 24 h and subsequently preserved in 70° ethanol at 4°C. Anthers were stained with a drop of 2% acetic orcein acetic acid and observed under a ZeissAxio LAB.A1 light microscope, Olympus.

Viability estimation was performed using the Müntzing, with a count of at least 1000 pollen grains (Maryam et al., 2015).

Pollen morphology: The material was prepared following the techniques for nonacetolysed and acetolysed pollen proposed by Wodehouse (1935) and Erdtman (1960), respectively. Preparations were mounted in glycerol gelatin. For scanning electron microscopy (SEM) analysis, acetolysed pollen grains were metallized with a gold-palladium bath. Pollen grains were described taking into account the number, type and position of apertures. The following parameters proposed by Faegri & Iversen (1966) were measured in 20 pollen grains: polar axis (P), equatorial diameter (E), shape (P/E), width and length of the colpi and thickness of the exine. Grain size was defined according to the size classes of Erdtman (1952). The terminology used in the description of pollen type corresponds to Punt et al., (1994).

Observations were made with a ZeissAxio LAB.A1, Olympus, optical microscope and a Zeiss Supra 55VP scanning electron microscope.

RESULTS

The study evidenced that *B. spinosa* has a sporophytic number $2n = 56$ chromosomes and karyotypic formula $46m + 10sm$ (Table 1).

TABLE 1
Morphometric parameters of *Borreria spinosa* chromosomes

Chromosome pair	c(μm) — $\bar{x} \pm SD$	l(μm) — $\bar{x} \pm SD$	s(μm) — $\bar{x} \pm SD$	Ci%	Chromosome type
1	2,70±0,63	1,43±0,26	1,27±0,22	47,14	m-sat
2	2,61±0,54	1,44±0,27	1,17±0,12	44,75	m
3	2,47±0,40	1,32±0,15	1,15±0,10	46,54	m
4	2,38±0,31	1,32±0,15	1,05±0,01	44,33	m
5	2,32±0,25	1,26±0,09	1,06±0,01	45,63	m
6	2,27±0,20	1,24±0,07	1,04±0,01	45,62	m
7	2,24±0,17	1,20±0,03	1,05±0,01	46,67	m
8	2,18±0,11	1,23±0,06	0,94±0,11	43,36	m
9	2,14±0,07	1,17±0,01	1,00±0,05	46,94	m
10	2,12±0,05	1,14±0,03	0,98±0,07	46,36	m
11	2,06±0,01	1,10±0,07	0,96±0,14	46,76	m
12	2,02±0,05	1,11±0,06	0,91±0,14	44,86	m
13	1,98±0,09	1,08±0,09	0,90±0,15	45,24	m
14	1,91±0,16	1,06±0,11	0,85±0,20	44,31	m
15	1,87±0,20	1,05±0,12	0,82±0,23	43,78	m
16	1,79±0,28	0,96±0,21	0,82±0,23	46,08	m
17	1,76±0,31	0,97±0,20	0,79±0,26	44,98	m
18	1,73±0,34	0,92±0,25	0,81±0,24	46,79	m
19	1,68±0,39	0,92±0,25	0,77±0,28	45,60	m
20	1,59±0,48	0,91±0,26	0,68±0,37	42,85	m
21	1,50±0,57	0,81±0,36	0,69±0,36	45,74	m
22	1,41±0,66	0,76±0,41	0,64±0,41	45,72	m
23	2,49±0,42	1,55±0,38	0,94±0,11	37,99	m
24	2,45±0,38	1,69±0,52	0,76±0,29	31,17	sm
25	2,46±0,39	1,59±0,42	0,86±0,19	35,12	sm
26	2,26±0,19	1,47±0,30	0,79±0,26	35,03	sm
27	1,93±0,14	1,29±0,12	0,64±0,41	33,01	sm
28	1,81±0,26	1,14±0,03	0,67±0,38	37,19	sm

Mean total chromosome length (c), mean long arm length (l), mean short arm length (s), mean centromeric index (ci), metacentric (m), submetacentric (sm), satellite (sat) and standard deviation (SD).

In the first metacentric chromosome pair, the presence of a satellite in a terminal position on the short arm was observed (Fig. 2A).

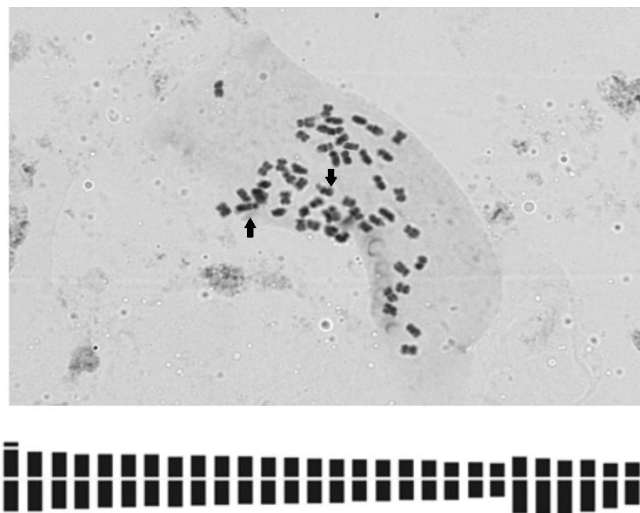


Fig. 2. Mitotic metaphase and idiogram of *Borreria spinosa*. A) Metaphase $2n = 56$ chromosomes. B) Idiogram (46m + 10sm). Arrows indicate the chromosome pair with satellite. Scale 10µm

The total length of the haploid chromosome complement was 58,12µm, the mean chromosome length was 2,07µm and the range of variation in mean chromosome lengths was 1,81-2,70 µm (Table 2).

TABLE 2

Karyotypic parameters of *Borreria spinosa*

Parameter	<i>B. spinosa</i>
<u>2n</u>	<u>56</u>
Karyotype formula	46m + 10sm
TCL (µm)	58,12µm
cme (µm)	2,07µm
cmax. (µm)	2,70µm
c min. (µm)	1,41µm
A ₁	0,20
A ₂	0,18

Total complement length (TCL), mean chromosome length (cme), maximum and minimum chromosome length (c max, c min), mean centromeric index (ci), intrachromosomal asymmetry index (A1), interchromosomal asymmetry index (A2). m: metacentric, sm: submetacentric.

Chromosomes showed a gradual size decrease (Fig. 2B). The karyotype is unimodal, since the chromosome size was similar; this is evidenced by the value of the interchromosomal asymmetry index $A_2 = 0,18$. The intrachromosomal asymmetry index indicates a symmetrical karyotype with $A_1 = 0,20$.

B. spinosa has a gametophytic number of $n = 28$ II (Fig. 3A). Numerous irregularities could be observed at both stages of meiotic division. Pollen mother cells (PMCs) in MI exhibited chromosomes outside the equatorial plate in 10% of them (Fig. 3B), AI with lagging chromosomes in 17% of them (Fig. 3C) and multiple spindle formation in about 90% of them (Fig. 3D). Cells in TI

were regular, and lagging chromosomes were occasionally observed in 2% of them. During the first stage of meiotic division, cytomixis was observed with the formation of cytoplasmic channels between two or more cells in the stages of diakinesis (Fig. 3E), MI and TI (Fig. 3F).

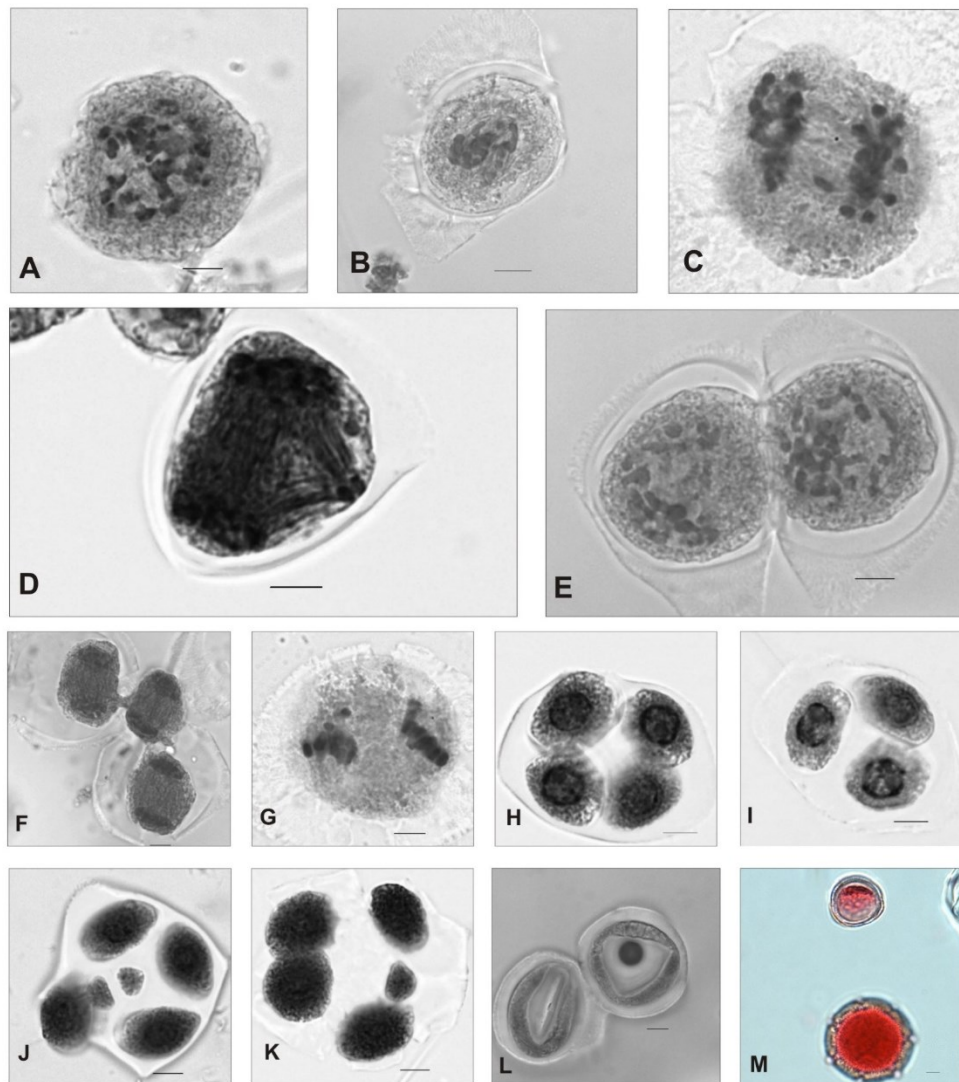


Fig. 3. Meiosis in *Borreria spinosa*. A) Diakinesis $n = 28$. B) MI. C) AI with lags. D) Multiple polar spindles in AI. E) Diakinesis with cytoplasmic channels. F) PMCs in AI with formation of cytoplasmic channels. G) MII with lagging chromosome. H) Tetrad. I) Triad. J) Hexad. K) Pentad. L) Cells without cellular content. M) Viable and nonviable pollen grain. Scale 5 μm .

In the second stage of meiotic division, PMCs in MII presented chromosomes outside the equatorial plate in 8% of them (Fig. 3G) and the formation of perpendicular spindles in 2% of them, whereas in AI and TI lagging chromosomes were observed in 2% of them. During spore formation, 1,47% triads (Fig. 3I), 81,54% tetrads (Fig. 3H), 15,86% pentads (Fig. 3K), 1,10% hexads with microspores and micronuclei (Fig. 3J) were recorded. Cytoplasmic channel formation was observed between spores. PMCs without cellular content were recorded throughout the division (Fig. 3L). Seventy percent of the pollen grains were viable; the rest were smaller and nonviable (Fig. 3M).

The nonacetolysed pollen grain (Fig. 4A-D) is small to medium-sized, suboblate to oblate spheroidal, with P 21 ($23\pm1,58$) $25\mu\text{m}$ and E 24 ($26\pm1,32$) $28\mu\text{m}$ and colpi5 ($5,92\pm0,67$) $6,60\mu\text{m}$ long to 1,70 ($2,35\pm0,47$) $3\mu\text{m}$ wide.

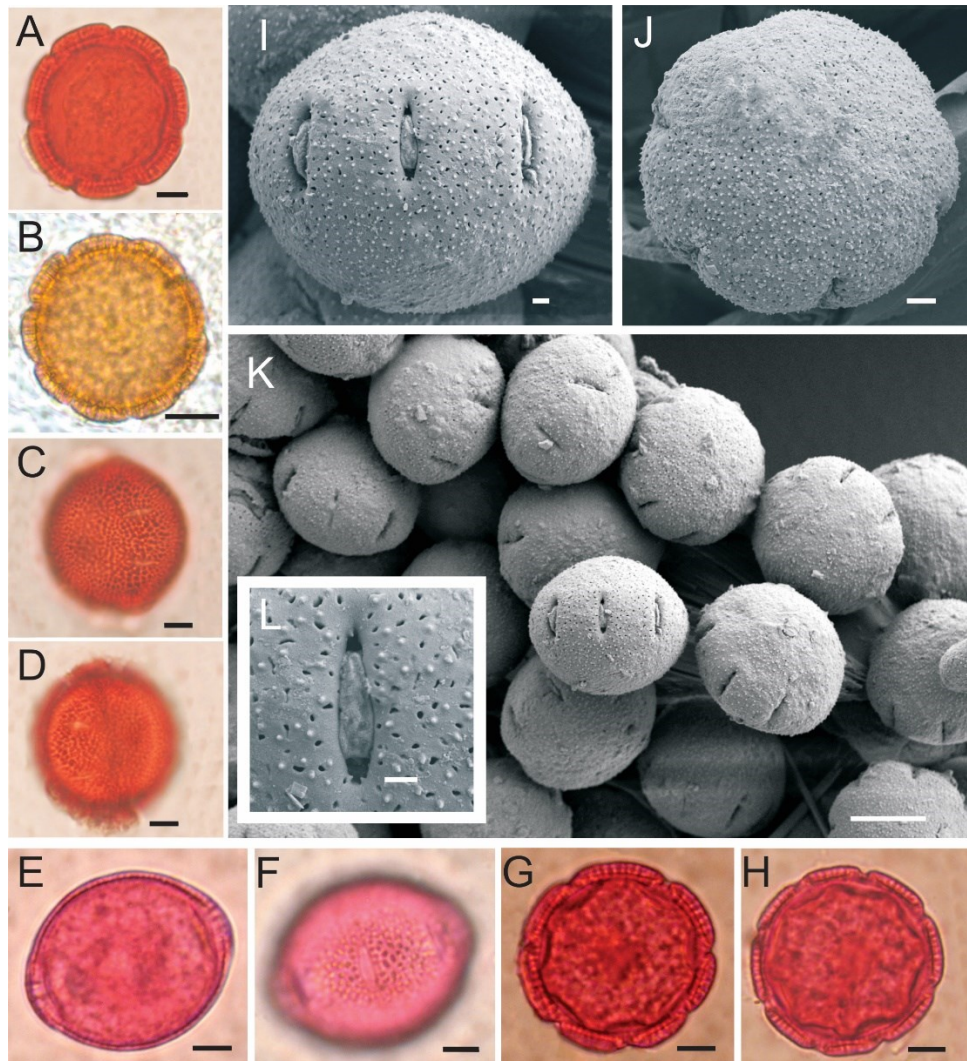


Fig. 4. Pollen grains on *Borreria spinosa*. A-D Nonacetolysed pollen. A) Equatorial view in optical section. B) Equatorial view on surface. C) Polar view in 8-colporate optical section. D) Polar view in 7-colporate surface. E-H Acetolysed pollen. E) Polar view in 9-colporate optical section. F) Polar view in 7-colporate optical section. G and H) Equatorial view on surface. I-L MEB. I) Equatorial view. J) Polar view. K) General view of the grains. L) Detail of aperture and surface. Scale 5μm

Acetolysed pollen grains (Fig. 4 E-H) are medium-sized, suboblate to oblate spheroidal, with P 24 ($25,40\pm1,26$) $27\mu\text{m}$, E 27 ($28,81\pm1,38$) $31\mu\text{m}$, circular in scope, 7-8-9 zonocolporate (52% 8-zonocolporate, 35% 7-zonocolporate, 13% 9-zonocolporate). Colpi are short ($5,5\text{--}6\mu\text{m}$ long), and their width varies from one line, when fully folded, to $1,5\mu\text{m}$, if expanded. The longitudinal end aperture is difficult to determine if it forms an endocingulum. The mesocolpium is $8,50$ ($10,19\pm1,07$) $11,50\mu\text{m}$, and the apocolpium is 21 ($24,31\pm1,87$) $26\mu\text{m}$. Perforations are observed in the tectum. The exine, 3 to $4\mu\text{m}$ thick, decreases towards the colpi. In the sexine, thicker than the nexine, simple columellae are observed.

Scanning electron microscopic observation corroborates that the grains are 7-8-9 zonocolporate (Fig. 4I-J), with microperforations of various shapes, from 0,20 to 0,50 μm in diameter (Fig. 4K-L). The positive elements observed with optical microscopy are nano-spinules ranging from 0,25 to 0,30 μm in height with a sharp apex. Perforations, as well as nanospinules, are densely and regularly arranged.

DISCUSSION

The basic chromosome number proposed for the tribe Spermacoceae is $x = 14$; thus, *B. spinosa* would be a tetraploid species with $2n = 56$.

In accordance with the results obtained for *B. spinosa*, Selvaraj (1987) performed chromosome counts and karyotypes in *B. articularis* and *B. pusilla* and reported the same chromosome number ($2n = 56$) with ranges of variation from 1 to 3,5 and 0,5 to 3,5 μ , respectively. The chromosome variation recorded in *B. spinosa* is more closely related to that of *B. articularis*. In addition, this author reported karyotypic formulas $30m + 18sm + 4st + 2B$ (B chromosomes) for *B. articularis* and $30m + 22m + 4st$ for *B. pusilla* that do not coincide with those recorded in the species under study ($46m + 10sm$). Selvaraj (1987) observed the presence of satellites in several chromosomes, which agrees with what was observed in *B. spinosa*, although in the latter satellites were only present in the largest chromosome pair.

Meiosis is a decisive process for sexual reproduction in plants. Abnormalities are occasionally observed that can lead to sterility of gametes as well as variation in their genetic constitution (Thangavel et al., 2023; Wang et al., 2023). Extensive chromosomal rearrangement and the formation of a single, transient spindle during mitosis and meiosis ensure proper chromosome segregation (Felismino et al., 2015). In plants, the formation of a bipolar spindle is the rule and allows for viable gametes and balanced genetic constitution. However, the formation of multipolar spindles can lead to meiotic irregularities (Shamina et al., 2003). *Borreria spinosa* exhibited multipolar spindle formation consisting of three or more poles; its chromosomes, according to Sidorchuk & Deineko (2014), would be randomly positioned in metaphases and then distributed in three or more directions during anaphase. This irregular spindle activity would result in the random misorientation of chromosomes in PMCs and subsequent formation of independently oriented chromosome subgroups. In the taxon studied, irregular spindle activity could be responsible for dyads, triads, polyads and micronuclei during microsporogenesis and subsequently result in pollen grains with reduced viability, as observed by Shamina et al., 2003.

During the first stage of meiotic division, a little-known phenomenon was recorded: cytomixis. It involves the transfer of chromatin or nuclei among plant cells through intercellular channels, frequently observed during microsporogenesis (Mursalimov et al., 2014; Mursalimov & Deineko, 2017). Whether cytomixis is a spontaneous or induced process, it can have severe genetic consequences, which combined with other meiotic abnormalities significantly affects pollen fertility (Mursalimov & Deineko, 2017; Mandal & Nandi, 2017).

The anomalies observed during meiosis are related to pollen sterility and low seed production characteristic of this species, which has the ability to reproduce sexually and asexually (Luna & Druetta, 2018). Thus, the presence of meiotic irregularities may imply that the process of sexual reproduction is not fully effective for its propagation.

According to the literature, the pollen morphology of *B. spinosa* has not yet been studied; on the contrary, different authors have found considerable variations in the pollen characters of this genus, thus defining a Euro-pollen genus. Gonçalves-Esteves et al., (2020) described the morphology of pollen grains of Rubiaceae from the Atlantic Forest of the state of Rio de Janeiro (Brazil), including *B. brachystemoides*, *B. capitata*, *B. latifolia*, *B. palustris*, *B. scabioisoides* and *B. verticillata*. In these

species, the pollen grains were small to medium-sized, oblate-spheroid and suboblate, colporate, with varying numbers of openings (5-12). Polar diameter was highly variable among the species studied, with a minimum value of $24,1 \pm 0,2 \mu\text{m}$ in *B. brachystemoides* and a maximum of $43,6 \pm 0,3$ in *B. latifolia*. The equatorial diameter ranged between values of $25,7 \pm 0,1$ in *B. brachystemoides* and $48,7 \pm 0,3 \mu\text{m}$ in *B. latifolia*. Dutra et al., (2020) described *B. verticillata* (L.) G. Mey pollen grains collected in the Brazilian cerrado as medium-sized, prolate spheroidal, with a polar diameter ranging from $27,5$ to $32,5 \mu\text{m}$ and an equatorial diameter ranging from $25,0$ to $27,5 \mu\text{m}$, 7-colpate and tectate-microechinate exine.

Borreria spinosa is a tetraploid, and most of its chromosomes are metacentric or submetacentric. It exhibits irregular meiotic behavior and cytomixis, with concomitant low viability in pollen grains. In this taxon, palynology showed that pollen grains have similar characteristics to other *Borreria* species.

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ETHICAL, CONFLICT OF INTEREST AND FINANCIAL STATEMENTS

The authors declare that they have fully complied with all pertinent ethical and legal requirements, both during the study and in the production of the manuscript; that there are no conflicts of interest of any kind; that all financial sources are fully and clearly stated in the acknowledgements section; and that they fully agree with the final edited version of the article. A signed document has been filed in the journal archives.

The statement of each author's contribution to the manuscript is as follows: P.B.P, V. de los A.P, and D.A.M.: Study design, data collection, analysis, result interpretation and manuscript writing. R.A.A. and M.L.E.: Data collection and final approval of the manuscript. All co-authors: preparation and final approval of the manuscript.

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