

Ploidy, Relative Genome Size, and Inheritance of Spotted Foliage in *Aucuba* Species (Garryaceae)

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Abstract. *Aucuba* have been cultivated for centuries and are valued as adaptable, broad-leaved, evergreen shrubs that also can have attractive, spotted variegations on the foliage. Improved understanding of the cytogenetics and heritability of specific traits, for specific clones and cultivars, can provide basic information to help facilitate the breeding and improvement of *aucuba*. The objectives of this study were to determine ploidy level and relative genome size of a diverse collection of species and cultivars of *aucuba* using flow cytometry and cytology and to make additional observations on heritability of spotted leaf variegation. Chromosome counts were $2n = 2x = 16$ for *Aucuba chinensis* (*A. omeiensis*), $2n = 4x = 32$ for *A. japonica* ‘Rozannie’, and $2n = 6x = 48$ for *A. sp.* ‘Hosoba’. Relative 2C genome size for the 57 taxa varied from 13.8 pg for *A. obtusata* to 42.0 pg for *A. ‘Hosoba’* and fell within three discrete groups consistent with cytotype. Genome size for diploid taxa (*A. chinensis* and *A. obtusata*) ranged from 13.8 to 21.0 pg, tetraploids (*A. himalaica* var. *oblanceolata*, *A. japonica*, and *A. japonica* var. *borealis*) ranged from 28.8 to 31.2 pg, and the first-ever reported hexaploids (*A. ‘Hosoba’* and *A. sp.* – Vietnam) ranged from 40.5 to 42.0 pg. Unlike prior reports that indicated inheritance of spotted variegations were extranuclear genes that were maternally inherited, we found that the spotted leaf trait expressed in *A. japonica* ‘Shilpot’ appears to be a nuclear gene that is inherited in a quantitative fashion and not strictly maternal. These data provide an enhanced foundation for breeding improved *aucuba*.

Aucuba (*Aucuba*) is a relatively small genus of about nine species (The Plant List, 2018). Formerly placed in the Cornaceae and Aucubaceae families, the genus now resides in the Garryaceae (Stevens, 2018). Although currently extant in Eastern Asia (ranging from the Eastern Himalayas east to China, Korea, and Japan and south to Myanmar, Taiwan, and Vietnam), fossil leaves of *aucuba* from the Eocene have been found in Washington State, suggesting a

much broader past distribution (Wehr and Hopkins, 1994).

Cultivated for centuries, *aucuba* are revered as showy, adaptable, evergreen shrubs (Creech, 1984). *Aucuba japonica* is commonly grown around shrines and temples in Japan, thriving in the understory of *Cryptomeria japonica*. John Graeffler is credited with introducing *A. japonica* to the Western World in 1783, where it has become an important and valuable landscape plant (The International Dendrology Society, 2018). Of particular merit are many cultivars that have showy, variegated foliage ranging from gold flecking and spots to variegated leaf margins. Although the flowers are inconspicuous and dioecious, female plants can have outstanding displays of red fleshy fruit. In addition to ornamental merit, *A. japonica* are notoriously tolerant of dry shade, pruning, pollution, and general neglect (Lehrer, 2009). Large specimens frequently are found on old estates and homesteads in the Southeastern United States, where they have long outlived their original gardeners.

Knowledge of ploidy is important for plant breeders, as it can influence crossability, fertility of progeny, segregation of traits, and gene expression. There is limited information on the cytogenetics of *aucuba*. Some species, including *A. chinensis*,

eriobotryifolia, *himalaica*, and *omeiensis*, have been reported to be diploid with $2n = 2x = 16$ (Hara, 1966; Kurosawa, 1971; 1981; The Chromosome Counts Database, 2018). *Aucuba japonica* are more variable and well-studied, with three varieties (*japonica*, *borealis*, and *ovoidea*) that have been classified based on a combination of morphology, ploidy, and geographic distribution (Ohi et al., 2003). Variety *borealis* is the most northerly distributed and found in southwestern Hokkaido and the Sea of Japan (Western) side of Honshu and is tetraploid ($2n = 4x = 32$). Variety *japonica* is present on the Pacific Ocean (Eastern) side of Honshu and Shikoku and is also tetraploid. Variety *ovoidea* is distributed further south, ranging from southern Honshu, western Shikoku, and Kyushu to Okinawa, and is diploid. Variety *borealis* is considered to be the most cold-hardy of the group with a smaller stature, decumbent shoots, smaller leaves, and pubescent young buds and inflorescences. Varieties *japonica* and *ovoidea* are larger growing and more similar to one another in appearance but clearly differentiated by ploidy. Karyomorphological studies on a diverse sampling of diploid *A. japonica* var. *ovoidea* showed the presence of XY sex chromosomes where the fourth-longest pair of chromosomes was heterogeneous in male individuals and homogeneous in female individuals (Tsusaka et al., 2007). Despite the work that has been completed on chromosome numbers and ploidy in *aucuba*, there is extremely little documentation of ploidy for specific cultivars and clones.

Minimal work has been published on genetics and heritability of specific traits in *aucuba*. Hagedoorn (1950) first reported that the spotted leaf trait was maternally inherited and that all seedlings from spotted-leaved females had spotted leaves regardless of the male parent and that seedlings from green-leaved female parents always had green leaves even when the pollen parent had spotted leaves. Allen (1990) further substantiated this observation by collecting seed from green-leaved and spotted-leaved plants that all produced seedlings with the same phenotype as the female parent.

There is considerable potential to breed and improve *aucuba* as nursery crops. The diversity in desirable foliage characteristics (leaf size and variegations), plant size, cold hardiness, and potentially disease resistance and regional adaptability provides substantial opportunities to combine desirable traits. Additional cytogenetic data, with specific information on ploidy of specific cultivars and clones, and information on the mode of inheritance of specific traits, would provide valuable information to better enable plant breeding programs. The objectives of this study were to determine ploidy and relative genome size for a diverse collection of species and cultivars of *aucuba* using flow cytometry and cytology and to make

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additional observations on the heritability of spotted leaf variegation of specific cultivars.

Materials and Methods

Plant material. Plants and/or tissue from 57 taxa of aucuba were obtained from the JC Raulston Arboretum (JCRA), Raleigh, NC; the Mountain Crop Improvement Laboratory, Mills River, NC; and the United States National Arboretum, Washington, DC. Current nomenclature for aucuba was adopted from The Plant List (2018) and Xiang and Boufford (2005). Synonyms from accession records and collection data were included when present.

Cytology. Chromosome counts were determined for three taxa, including *A. chinensis* (*A. omeiensis*, JCRA 120820), *A. japonica* ‘Rozannie’ (MCIL 2015-077), and *A. ‘Hosoba’* (JCRA 120812) to determine chromosome numbers and calibrate genome size measurements with ploidy. Actively growing root tips were harvested from container-grown plants in mid-July and placed into a solution of 2 mM 8-hydroxyquinoline and 0.248 mM cycloheximide and incubated for 3 h at 23 °C, then moved to 4 °C for an additional 3 h. Roots were then rinsed in cold distilled water and fixed in Carnoy’s solution (six parts 95% ethanol: three parts chloroform: one part glacier acetic acid) at 21 °C for 18 hours before storage in 70% ethanol. Fixed roots were hydrolyzed in a 3:1 95% ethanol:12 M HCl for ≈15 min and stained with a modified carbol fuchsin stain for at least 5 min (Kao, 1975; Singh, 2003). Root tips were then placed onto a microscope slide with a drop of stain, overlaid with a cover slip. Tissue was squashed underneath a cover slip, and chromosomes were counted at ×1000 magnification.

Flow cytometry. Tissue from vegetative buds (≈0.3 cm²) was collected and placed in a plastic petri dish with ≈0.3 cm² of expanding leaf tissue from *Pisum sativum* ‘Ctirad’ that served as an internal standard with a known genome size of 2C = 8.75 pg (Greilhuber et al., 2007). Samples were finely chopped with a razor blade in 0.4 mL of nuclei extraction buffer (CyStain ultraviolet Precise P Nuclei Extraction Buffer; Sysmex Partec, Görlitz, Germany) and filtered through a 50-μm nylon mesh filter. Nuclei were stained with 1.6 mL 4′,6-diamidino-2-phenylindole (DAPI) before analysis with a flow cytometer (Partec PA II, Munster, Germany). Samples were run until at least 5000 nuclei were counted, and two subsamples were analyzed for each accession. Genome size (2C) of samples was calculated as: 2C = genome size of standard × (mean fluorescence value of sample ÷ mean fluorescence value of standard). The 1C_x monoploid genome size (i.e., DNA content of one base set of chromosomes) was calculated as the 2C genome size ÷ ploidy level and subjected to analysis of variance by taxa, and means were separated using LSD_{0.05} (Proc GLM; SAS Version 9.3; SAS Institute, Cary, NC).

Heritability of spotted variegation. Cross pollinations were completed between the female, green-leaved cultivar *A. japonica* ‘Petite Jade’ and the male, spotted-leaved cultivar *A. japonica* ‘Shilpot’ in Mar. 2015. Fruit were collected in the Fall of 2015, seeds were extracted, given cold/moist stratification for 90 d at 6 °C, and germinated and grown under standard production practices. Seedlings were evaluated for expression of spotted variegations in May 2018.

Results and Discussion

The chromosome count for *A. chinensis* (*A. omeiensis*, JCRA 120820) was diploid with $2n = 2x = 16$ (Fig. 1), consistent with a previous report for *A. chinensis* (Kurosawa, 1981). *Aucuba japonica* ‘Rozannie’ (MCIL 2015-077) was confirmed to be tetraploid with $2n = 4x = 32$ (Fig. 1), consistent with other reports of tetraploid forms of *A. japonica* var. *japonica* and var. *borealis* (Ohi et al., 2003). Surprisingly, *A. ‘Hosoba’* (JCRA 120812) was found to be hexaploid with $2n = 6x = 48$ (Fig. 1), being the first report of a hexaploid aucuba. The species designation of this cultivar has not been determined yet, though records at the JCRA indicate it was of Chinese origin. Confirmation of these cytotypes also provides calibration points to allow estimation of ploidy of other taxa from genome size data.

Relative 2C genome size for the 57 taxa varied from 13.8 pg for *A. obtusata* to 42.0 pg for *A. ‘Hosoba’* (Table 1). These values fell within three discrete groups consistent with cytotype. Genome size for diploid taxa ranged from 13.8 to 21.0 pg, tetraploids ranged from 28.8 to 31.2 pg, and hexaploids ranged from 40.5 to 42.0 pg. These values substantiate previous reports that *A. chinensis* and *himalaica* are diploid and that *A. japonica* (including *A. japonica* var. *borealis*) are tetraploid. The lack of any diploid *A. japonica* var. *ovoidea* present in our sampling may reflect their more southern nativity and possibly limited cold hardiness restricting their cultivation. Our finding that *A. obtusata* is diploid is newly reported. In addition to the cytological confirmation that *Aucuba* ‘Hosoba’ is hexaploid, genome size measurements indicate that the two *Aucuba* accessions from Vietnam (JCRA 150090 and 150090) are also hexaploid; however, it would be desirable to confirm this with actual chromosome counts. Although the correct species designations for these Vietnam taxa have not yet been determined, both were wild-collected from Fansipan (Phan Xi Păng) Mountain in North Vietnam at elevations between 2096 and 2310 m.

Overall, these genome size values were similar but greater than those values reported by Zonneveld et al. (2005) for (tetraploid) *A. japonica*, which ranged from 24.9 to 25.7 pg [determined with propidium iodide (PI) stain and *Agave americana* as an internal standard]. Different fluorochrome stains can give different estimates of absolute genome size, although both PI and DAPI have been found

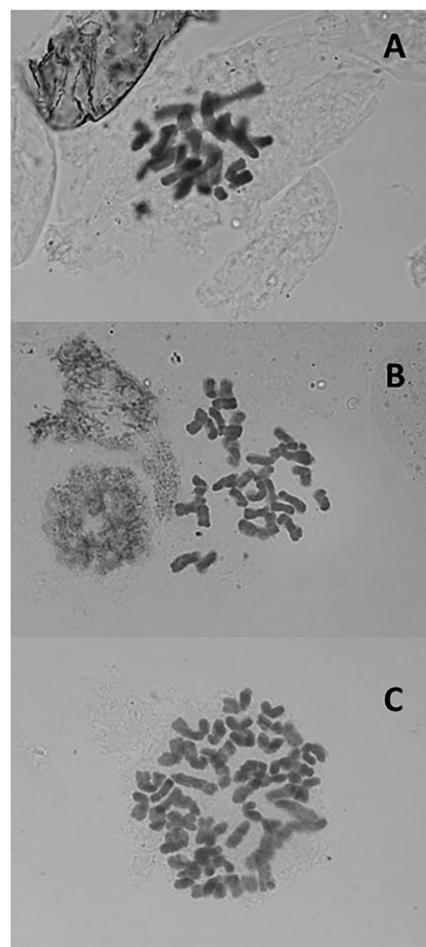


Fig. 1. Somatic chromosomes of (A) *Aucuba chinensis* ($2n = 2x = 16$), (B) *A. japonica* ‘Rozannie’ ($2n = 4x = 32$), and (C) *A. sp.* ‘Hosoba’ ($2n = 6x = 48$).

to be effective and consistent for determining and comparing relative genome size and ploidy among closely related taxa (Parris et al., 2010). Furthermore, DAPI is less expensive, faster, and typically provides more precise and repeatable results as it is specific to double-stranded DNA and is not influenced by variable chromatin structure (Doležel and Bartoš, 2005; Greilhuber et al., 2007).

Base 1C_x genome size was fairly similar for most aucuba species, ranging from 6.8 to 7.5 pg with the notable outlier of *A. chinensis* (*A. omeiensis*, JCRA 120820) that had a considerably higher value of 10.6 pg. Xiang and Boufford (2005) currently treat *A. chinensis* subsp. *omeiensis* and *A. omeiensis* as synonyms for *A. chinensis*. However, the larger and unique base genome size of *A. chinensis* (*A. omeiensis*, JCRA 120820) indicates that this accession/taxon has undergone considerably more within-ploidy genome expansion than other taxa evaluated and may warrant additional study and treatment as a separate species apart from *A. chinensis*.

Cross pollinations of *A. japonica* ‘Petite Jade’ by *A. japonica* ‘Shilpot’ yielded 14 seedlings that segregated with three plants

Table 1. Relative genome size and estimated ploidy for aucuba taxa.

Taxon	Source ^z	Accession	Relative 2C genome size (pg) ± SEM	Estimated Ploidy (x) ^y	Relative 1C _x genome size (pg) ± SEM by species ^x
<i>A. chinensis</i>	JCRA	090886	14.73 ± 0.04	2	7.37 ± 0.00 BC
<i>A. chinensis</i> (<i>A. omeiensis</i>)	JCRA	120820	21.20 ± 0.14	2*	10.60 ± 0.00 D
<i>A. himalaica</i> var. <i>oblanceolata</i> (<i>A. japonica</i> f. <i>longifolia</i>)	JCRA	140141	28.70 ± 0.50	4	7.19 ± 0.10 BC
<i>A. himalaica</i> var. <i>oblanceolata</i> ‘Salicifolia’ (<i>A. japonica</i> f. <i>longifolia</i> ‘Salicifolia’)	JCRA	130580	28.78 ± 0.02	4	
<i>A. japonica</i> ‘Big Mamma’	JCRA	110779	29.49 ± 0.04	4	7.36 ± 0.02 BC
<i>A. japonica</i> ‘Cecil-Alice’	JCRA	031019	29.44 ± 0.14	4	
<i>A. japonica</i> ‘Cho Dai Ji’	JCRA	100377	29.24 ± 0.61	4	
<i>A. japonica</i> ‘Crotonifolia’	JCRA	011565	28.80 ± 0.04	4	
<i>A. japonica</i> ‘Daisuke’s Tiger’	JCRA	120384	29.44 ± 0.01	4	
<i>A. japonica</i> ‘Eclipse’	JCRA	120706	29.46 ± 0.13	4	
<i>A. japonica</i> ‘Emily Rose’	MCIL	2015-079	29.83 ± 0.15	4	
<i>A. japonica</i> ‘Fructu Albo’	JCRA	xx0160	28.48 ± 0.00	4	
<i>A. japonica</i> ‘Fujikawa’	JCRA	100401	29.63 ± 0.16	4	
<i>A. japonica</i> ‘Goldilocks’	JCRA	041654	29.30 ± 0.04	4	
<i>A. japonica</i> ‘Hime Kikufurin’	JCRA	110300	28.47 ± 0.17	4	
<i>A. japonica</i> ‘Hosoba Hoshifu’	JCRA	100347	29.98 ± 0.08	4	
<i>A. japonica</i> ‘Kameba’	JCRA	120385	30.59 ± 0.14	4	
<i>A. japonica</i> ‘Koba-no-aoki-fuiki’	JCRA	140005	28.35 ± 0.35	4	
<i>A. japonica</i> ‘Linda Eggins’	JCRA	130661	29.40 ± 0.57	4	
<i>A. japonica</i> ‘Little Angel’	JCRA	120338	29.00 ± 0.03	4	
<i>A. japonica</i> ‘Marmorata’	MCIL	2015-045	28.89 ± 0.12	4	
<i>A. japonica</i> ‘Meigetsu’	JCRA	031020	29.26 ± 0.13	4	
<i>A. japonica</i> ‘Mr. Goldstrike’	MCIL	2015-046	29.38 ± 0.05	4	
<i>A. japonica</i> ‘Natsu-no-kumo’	JCRA	110303	29.82 ± 0.02	4	
<i>A. japonica</i> ‘Pacman’	JCRA	120678	29.91 ± 0.32	4	
<i>A. japonica</i> ‘Peachie’	JCRA	130131	29.20 ± 0.02	4	
<i>A. japonica</i> ‘Petite Jade’	JCRA	140140	29.22 ± 0.27	4	
<i>A. japonica</i> ‘Rozannie’	MCIL	2015-077	29.09 ± 0.12	4*	
<i>A. japonica</i> ‘Shilpot’ Pepperpot™	MCIL	2015-042	28.98 ± 0.21	4	
<i>A. japonica</i> ‘Shuugetsu’	JCRA	110313	28.78 ± 0.06	4	
<i>A. japonica</i> ‘Subaru’	JCRA	110305	28.88 ± 0.18	4	
<i>A. japonica</i> ‘Sulphurea Marginata’	JCRA	xx0159	29.49 ± 0.16	4	
<i>A. japonica</i> ‘Suruga Benten’	JCRA	110307	29.45 ± 0.02	4	
<i>A. japonica</i> ‘Tatsumaki’	JCRA	110301	29.71 ± 0.00	4	
<i>A. japonica</i> (chicory form)	JCRA	111032	29.30 ± 0.14	4	
<i>A. japonica</i> (cream-white, contorted)	JCRA	110318	30.81 ± 0.23	4	
<i>A. japonica</i> (gold with green center)	JCRA	131402	28.67 ± 0.88	4	
<i>A. japonica</i> (dwarf)	JCRA	120704	30.64 ± 0.26	4	
<i>A. japonica</i> (seedling #6)	JCRA	041598	30.26 ± 0.23	4	
<i>A. japonica</i> (thin white margin)	JCRA	110306	29.68 ± 0.19	4	
<i>A. japonica</i> (USNA #3 – female)	JCRA	040065	29.99 ± 0.45	4	
<i>A. japonica</i> (USNA #4)	JCRA	040066	29.30 ± 0.44	4	
<i>A. japonica</i> (USNA #5 – female)	JCRA	040067	30.02 ± 0.15	4	
<i>A. japonica</i>	USNA	2016-034	30.52 ± 0.19	4	
<i>A. japonica</i> var. <i>borealis</i> ‘Bored Female’	MCIL	2015-043	29.65 ± 0.20	4	7.53 ± 0.07 C
<i>A. japonica</i> var. <i>borealis</i> ‘Honshu’	JCRA	140004	29.23 ± 0.18	4	
<i>A. japonica</i> var. <i>borealis</i> ‘Sea of Japan’	MCIL	2015-067	28.78 ± 0.01	4	
<i>A. japonica</i> var. <i>borealis</i>	USNA	2016-035	31.22 ± 0.46	4	
<i>A. japonica</i> var. <i>borealis</i>	USNA	2016-036	30.54 ± 0.35	4	
<i>A. japonica</i> var. <i>borealis</i>	USNA	2016-037	30.61 ± 0.07	4	
<i>A. japonica</i> var. <i>borealis</i>	USNA	2016-038	30.34 ± 0.00	4	
<i>A. japonica</i> var. <i>borealis</i>	USNA	2016-039	30.49 ± 0.41	4	
<i>A. obcordata</i>	JCRA	121186	13.84 ± 0.00	2	7.02 ± 0.10 AB
<i>A. aff. obcordata</i>	JCRA	140899	14.23 ± 0.08	2	
<i>A. sp.</i> (Vietnam, female – Hinkley, DJHV 8035)	JCRA	150091	40.46 ± 0.06	6	6.76 ± 0.02 A
<i>A. sp.</i> (Vietnam, male – Hinkley and Wynn Jones, HWJ 1006)	JCRA	150090	40.67 ± 0.16	6	
<i>A. sp.</i> ‘Hosoba’	JCRA	120812	42.01 ± 0.00	6*	7.00 ± 0.00 AB

^zSource codes: MCIL = Mountain Crop Improvement Laboratory, Mills River, NC; JCRA = JC Raulston Arboretum, Raleigh, NC; USNA = United States National Arboretum, Washington, DC.

^yValues followed by an asterisk were confirmed with cytology.

^xValues followed by different letters within a column are significantly different, least significant difference, $P \leq 0.05$.

with homogeneous green leaves, three plants with faint and infrequent spots, and eight plants moderate-to-heavy variegation (Fig. 2). Considering that these plants are tetraploids, the population size was not large enough to adequately categorize the phenotypes and model the exact mode of inheritance, but it is clear that the spotted variegation trait of

‘Shilpot’ is quantitative and not strictly maternally inherited. This result is inconsistent with previous reports that the mode of inheritance in aucuba is strictly maternal/extranuclear (Allen, 1990; Hagedoorn, 1950). Although these earlier reports did not identify the particular clones used in their crosses, our combined results indicate that

there are probably multiple genes with different modes of inheritance that can give rise to variegated aucuba plants. The spotted-leaved trait expressed in *A. japonica* ‘Shilpot’ is most likely a nuclear gene with multiple alleles and quantitative inheritance. Alternatively, if it is an extranuclear gene, it would have to be inherited

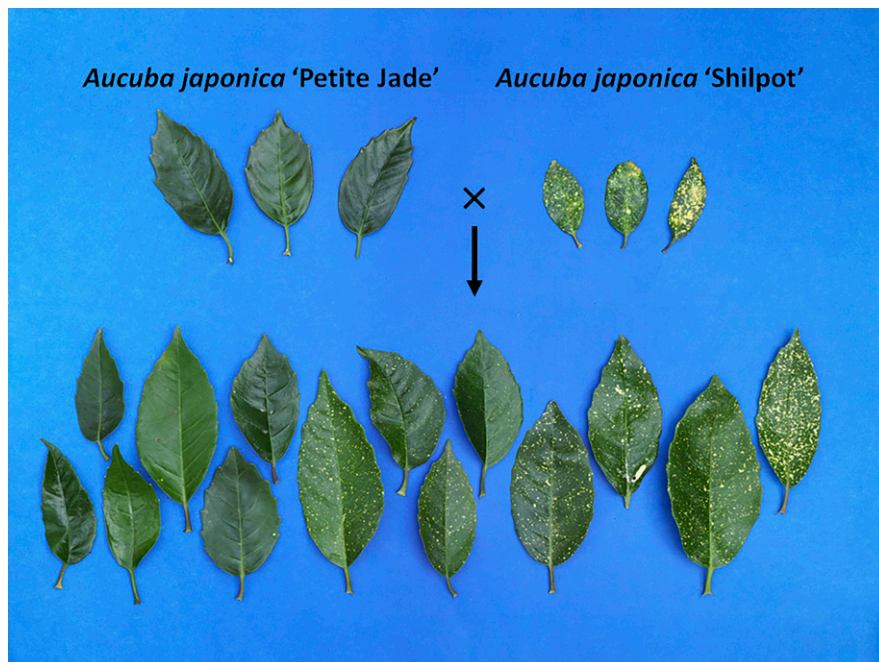


Fig. 2. Representative leaves from parents and 14 F₁ hybrids between *Aucuba japonica* 'Petite Jade' (female) and *A. j.* 'Shilpot' (male).

paternally/biparentally. Additional observations of seedlings from open-pollinated, green-leaved plants of *A. japonica* var. *borealis* 'Honshu' also produced seedlings with spotted leaves (data not presented), further corroborating that spotted variegation is not always maternally inherited.

These results document relative genome size and ploidy of a diverse collection of aucuba species and cultivars including cultivated and wild-collected specimens. Additional data are presented on a newly described quantitative mode of inheritance for a desirable spotted variegation trait. These data provide basic information on cytogenetics and heritability for specific clones and cultivars to aid in future breeding and improvement of aucuba. Expanded sampling of species and unclassified taxa has shown a wider range of ploidy and base genome size than previously

reported that provides additional insights into the evolution and systematics of this genus.

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