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Occurrence of Ploidy Variation in *Camellia* × *vernalis*

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Maternal inheritance of chloroplast DNA (cpDNA) in the genus *Camellia* was confirmed using the polymorphism of *atpH–atpI* region. CpDNAs of all *C. × vernalis* cultivars showed the same type as those of *C. sasanqua*, and all the progenies from *C. × vernalis*, either open pollinated or crossed, had the same cpDNA type as their maternal plants. Flow cytometry subjected to estimate the ploidy level of *C. japonica*, *C. sasanqua* and *C. × vernalis* revealed that there is a very strong positive linear correlation ($r^2 = 0.981$) between fluorescent intensity and ploidy level, suggesting that the method is useful to investigate the ploidy level in *Camellia*. Natural occurrence of ploidy variation was shown in the progenies of tetraploid *C. × vernalis* cultivars, and zygote patterns of the progenies are discussed.

INTRODUCTION

Camellia × *vernalis* is considered to be a hybrid species between *C. sasanqua* ($2n = 6x = 90$) and *C. japonica* ($2n = 2x = 30$) (Tanaka *et al.*, 1986). It has excellent characters like a long bloom time, a large number of flowers and large variations of flower colors and shapes. Tanaka *et al.* (1986) classified *C. × vernalis* cultivars into four groups; eleven cultivars of the ‘Gaisen’ type tetraploid, thirty-five of the triploid, seven of the ‘Egao’ type tetraploid and seven of the pentaploid groups, and postulated the origin of each group as follows; 1) the ‘Gaisen’ type tetraploids ($2n = 4x = 60$) were resulted from the primary hybrid between *C. sasanqua* and *C. japonica*, 2) the triploid ($2n = 3x = 45$) and pentaploid ($2n = 5x = 75$) groups originated

from the backcross generations of a tetraploid ‘Gaisen’ type with *C. japonica* and *C. sasanqua*, respectively and 3) the ‘Egao’ type tetraploids ($2n = 4x = 60$) were derived from the second backcross generation between an unreduced triploid gamete of the triploid *C. × vernalis* and a normal haploid gamete of *C. japonica* (Fig. 1).

Polymorphism of chloroplast DNA (cpDNA) has been often adopted in phylogenetic study because cpDNA shows maternal inheritance (Corriveau and Coleman, 1988) and slower evolution speed than nuclear DNA (Small *et al.*, 1998). Shibata *et al.* (2000) investigated the polymorphism of *atpH–atpI* region of cpDNA in the genus *Camellia*, and reported two types of PCR products (types A and B for approx. 800 bp and 1,200 bp, respectively). They also reported that *C. sasanqua*, *C. japonica* ssp. *rusticana* and *C. chrysantha* were classified into type A and *C. japonica* and *C. sinensis* were into type B, suggesting that the polymorphism can be used as a genetic marker to distinguish the species among genus *Camellia*. Tanaka *et al.* (2005) revealed that the products of this region in some *C. × vernalis* cultivars were type A, demonstrating that *C. sasanqua* was a maternal ancestor of *C. × vernalis*.

Ploidy levels have been determined by counting the somatic chromosomes, and also been estimated by measuring the length of stomata in many plant species. The procedures are still difficult and/or time consuming in *Camellia*, particularly in high ploidy species, so that the efficient analytical methods for rapid and suitable determination of ploidy levels have been long desired. Recently, flow cytometry has become a useful tool for rapid and efficient estimation of genome size and ploidy levels in some crops (Eaton *et al.*, 2004; Hcini *et al.*, 2006).

We investigated *atpH–atpI* region in *C. × vernalis* cultivars, their open pollinated seedlings and artificially crossed progenies with *C. japonica* to confirm their maternal inheritance, and examined their ploidy variation by flow cytometric analysis.

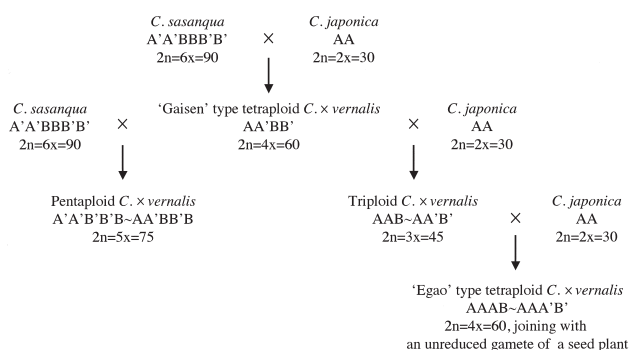


Fig. 1. Outline of introgressive hybrid formation between *C. sasanqua* and *C. japonica* (Tanaka, 1985, modified by N. T.).

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MATERIALS AND METHODS

Chloroplast DNA analysis*Plant materials*

Seven cultivars including four diploids ('Hatsu-arashi', 'Kanto-hatsuwari', 'Kariginu' and 'Shiratama') and three triploids ('Akashigata', 'Akebono' and 'Kumagai') of *C. japonica*, two cultivars ('Choji-guruma' and 'Inu-hariko') and four accessions of *C. sasanqua* from Kurokamiyama, Saga Pref., Hirado, Nagasaki Pref., Funakakushi, Goto Is., Nagasaki Pref. and Osuzuyama, Miyazaki Pref., and eight cultivars ('Egao', 'Gaisen', 'Omi-goromo', 'Sado', 'Sayo-hime', 'Takarazuka', 'Tamuke-yama' and 'Ume-ga-ka') of *C. × vernalis* were supplied to investigate their cpDNA. Twenty-one plants obtained from the crosses between 'Gaisen' and *C. japonica*, and between 'Ume-ga-ka' and *C. japonica*, and the seedlings from the open pollination of 'Gaisen' and 'Ume-ga-ka' were also used.

DNA extraction

Total genomic DNA was extracted from 100 mg of frozen young leaf or leaf/flower buds by the modified CTAB method (Kobayashi *et al.*, 1998). Frozen samples were powdered with a mortar and a pestle, and washed three times with 1 ml of 50 mM Tris-HCl buffer (pH 8.0) containing 5 mM of EDTA, 350 mM of sorbitol, 0.1% of mercaptoethanol and 10% polyethylene-glycol 6,000. The pellet was suspended in 500 μ l of CTAB buffer and separated with chloroform-isoamyl alcohol (24:1). After centrifugation at 12,000 rpm for 5 min at room temperature, the aqueous layer was transferred into 1.5 ml micro tubes. The DNA precipitated by isopropanol was washed with 70% ethanol and dried. The DNA was, then, dissolved in Tris-EDTA buffer.

PCR analysis

AtpH-atpI region was amplified by PCR by using a pair of primers, *atpH* (5'-TTGACCAACTCCAGGTC-CAA-3') and *atpI* (5'-CCGCAGCTTATATAGGC-GAA-3'). PCR amplification was performed in a total volume of 25 μ l solution containing 25 ng template DNA, 0.5 μ M of each primer, 1.25 mM of each dNTP, 2.5 μ l of 10 \times reaction buffer and 0.5 unit of *Taq* DNA polymerase (Roche). The amplification was carried out by using Program Temp Control System PC-701 (ASTECC) with one cycle of 30 sec at 94°C, followed by 45 cycles of 30 sec at 94°C, 2 min at 60°C and 3 min at 72°C and one cycle of 7 min at 72°C. The PCR products were separated by electrophoresis in 1.5% agarose gel, and visualized under UV illumination after staining with ethidium bromide.

Flow cytometric analysis*Plant materials*

Five cultivars including four diploids ('Hatsu-arashi', 'Kanto-hatsuwari', 'Kariginu' and 'Shiratama') and one triploid ('Akebono') of *C. japonica*, two accessions of *C. sasanqua* from Hirado, Nagasaki Pref. and Harumaki, Yakushima Is., Kagoshima Pref., and seven cultivars

('Egao', 'Omi-goromo', 'Sado', 'Sayo-hime', 'Takarazuka', 'Tamuke-yama' and 'Ume-ga-ka') of *C. × vernalis* were screened for flow cytometry analysis. Twenty-five plants (14 triploids, one pentaploid, one hexaploid and nine unknown) obtained from the crosses between 'Gaisen' and *C. japonica* and between 'Ume-ga-ka' and *C. japonica*, and the seedlings of the open pollination of 'Gaisen' and 'Ume-ga-ka' were also provided for the analysis.

Flow cytometry

Approximately 1 cm² petal was chopped with a sharp razor blade in nuclei extraction buffer (High resolution DNA kit, Partec), and the suspension containing the released nuclei was passed through a 50 μ m nylon mesh filter. Then the nuclei in filtrate were stained with four times volumes of staining solution (High resolution DNA kit, Partec) containing 4'-6-diamidino-2-phenylindole (DAPI). After shaking the solution gently, samples were analyzed with a flow cytometer (PA Ploidy Analyzer, Partec). Relative DNA content was estimated according to the fluorescent intensity of the prominent peak in each measurement. Correlation between fluorescent intensity of the prominent peak and ploidy level was evaluated and statistical analysis of the intensity of each cultivar/accession was carried out by Tukey's HSD test.

RESULTS AND DISCUSSION

Chloroplast DNA analysis

Two types of PCR amplified products were observed in *atpH-atpI* region; types A (approx. 800 bp) and B (approx. 1,200 bp) (Fig. 2), corresponding with the previous studies (Shibata *et al.*, 2000; Tanaka *et al.*, 2005). 'Egao', 'Gaisen', 'Sayo-hime', 'Takarazuka' and 'Ume-ga-ka' of *C. × vernalis* had A type cpDNA in accordance with Tanaka (2005)'s result (Table 1). Other *C. × vernalis* cultivars, 'Omi-goromo', 'Sado' and 'Tamuke-yama' also had A type cpDNA suggesting that their maternal ancestors were also *C. sasanqua*. All the artificially crossed progenies and open pollinated seedlings showed the same cpDNA type (Type A) as

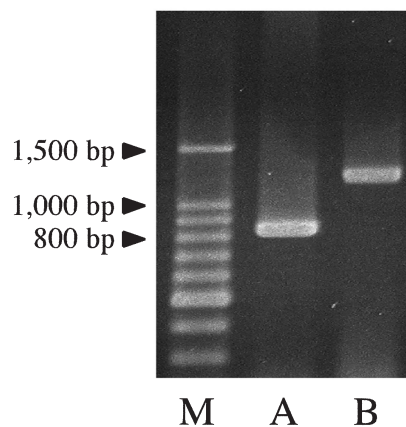


Fig. 2. Profiles of PCR products of *atpH-atpI* region in cpDNA. A: *C. sasanqua* (approx. 800 bp) B: *C. japonica* (approx. 1,200 bp) M: 100 bp DNA ladder marker.

Table 1. PCR products type of *atpH-atpI* region of cpDNA in *C. japonica*, *C. sasanqua*, *C. × vernalis* and their progenies

	Cultivar, accession	<i>atpH-atpI</i>
<i>C. japonica</i>	'Hatsu-arashi'	B
	'Kanto-hatsuwara'	B
	'Kariginu'	B
	'Shiratama'	B
	'Akashigata'	B
	'Akebono'	B
	'Kumagai'	B
<i>C. sasanqua</i>	'Choji-guruma'	A
	'Inu-hariko'	A
	Kurokamiyama, Saga Pref.	A
	Hirado, Nagasaki Pref.	A
	Funakakushi, Goto Is., Nagasaki Pref.	A
<i>C. × vernalis</i>	Osuzuyama, Miyazaki Pref.	A
	'Egao'	A
	'Gaisen'	A
	'Omi-goromo'	A
	'Sado'	A
	'Sayo-hime'	A
	'Takarazuka'	A
	'Tamuke-yama'	A
	'Ume-ga-ka'	A
	Crosses	<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 2
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 3		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 4		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 5		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 6		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 7		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 9		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 10		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 11		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 12		A
<i>C. × vernalis</i> 'Gaisen' × <i>C. japonica</i> No. 13		A
<i>C. × vernalis</i> 'Ume-ga-ka' × <i>C. japonica</i> No. 1		A
Open pollination		<i>C. × vernalis</i> 'Gaisen' open seedling No. 1
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 3	A
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 5	A
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 7	A
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 8	A
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 9	A
	<i>C. × vernalis</i> 'Gaisen' open seedling No. 10	A
	<i>C. × vernalis</i> 'Ume-ga-ka' open seedling No. 1	A
<i>C. × vernalis</i> 'Ume-ga-ka' open seedling No. 2	A	

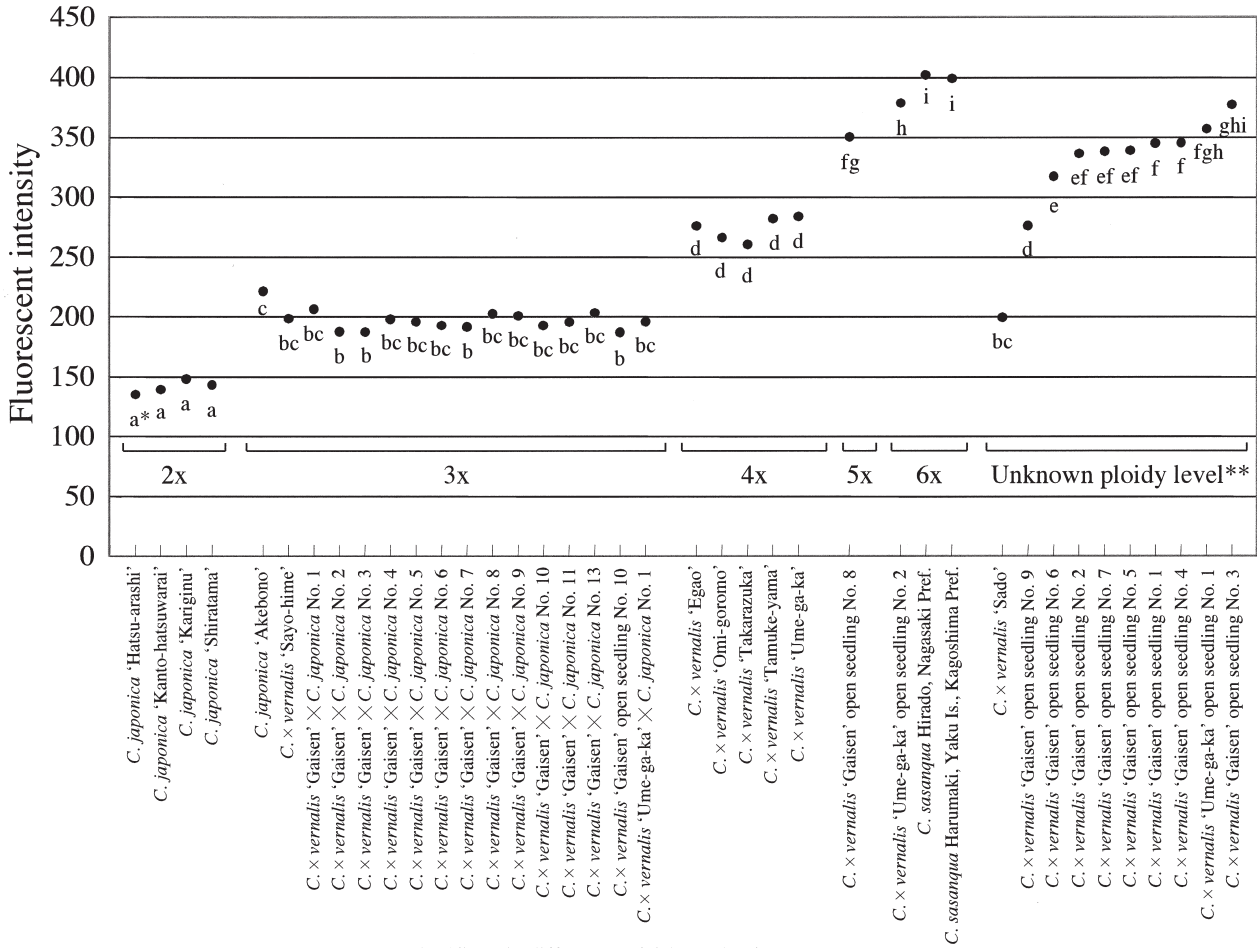
their maternal parents, confirming the maternal inheritance of cpDNA in *Camellia*.

Flow cytometric analysis

When the fluorescent intensity of the prominent peak of *C. sasanqua* (hexaploid) was adjusted to 400, the values of diploids, triploids, tetraploids and pentaploids were around 140, 200, 270 and 350, respectively (Fig. 3). Significant differences ($P < 0.05$) of the fluorescent intensities were found between different ploidy cultivars/accessions. The strong positive linear correlation ($r^2 = 0.981$) between chromosome number and fluorescent intensity was recognized (Fig. 4). These results suggest that flow cytometry is useful for distinguishing ploidy levels in the genus *Camellia*. Ploidy levels of the accessions whose chromosome number were unknown were estimated from the results of statistical analysis (Fig. 3). It was suggested that, for example, *C. × vernalis* 'Sado' and 'Gaisen' open seedling No. 9 were

triploid and tetraploid, respectively. Pentaploidy was estimated in 'Gaisen' open seedlings Nos. 2, 7, 5, 1 and 4. 'Gaisen' open seedlings Nos. 6 and 3 and 'Ume-ga-ka' open seedling No. 1 were considered to be aneuploids.

Ploidy variation was recognized in open pollinated progenies from 'Gaisen' and 'Ume-ga-ka'. The triploid progeny ('Gaisen' open seedling No. 10) might be derived from the cross between 'Gaisen' and *C. japonica*, and the tetraploid progeny ('Gaisen' open seedling No. 9) might be from the cross with other tetraploids of *C. × vernalis*. The following two possibilities are considered for the pentaploid progenies ('Gaisen' open seedlings Nos. 8, 2, 7, 5, 1 and 4); 1) progenies from the zygote between an unreduced tetraploid gamete of 'Gaisen' and normal haploid gamete of *C. japonica* or 2) progenies from the zygote between normal diploid gamete of 'Gaisen' and normal triploid gamete of *C. sasanqua*. The former hypothesis seems unlikely because all the progenies of 'Gaisen' × *C. japonica* were



*Values with same letters are not significantly different at 5% by Tukey's HSD test.
 **Chromosome number was unknown.

Fig. 3. Fluorescent intensity of the prominent peak in *C. japonica*, *C. sasanqua*, *C. × vernalis* and their progenies.

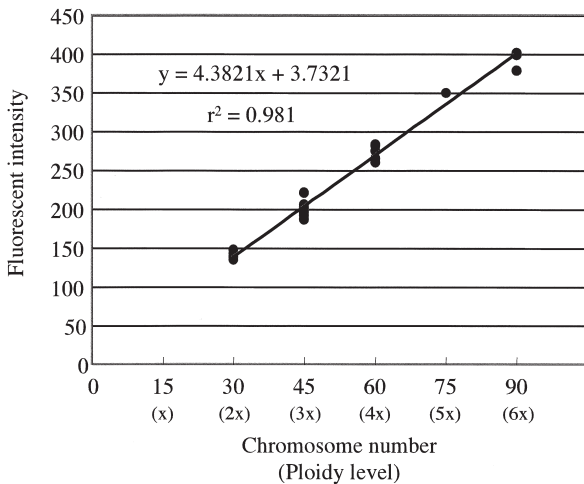


Fig. 4. Correlation between chromosome number and fluorescent intensity in *Camellia* species.

triploids. Hexaploid progeny ('Ume-ga-ka' open seedling No. 2) might be derived from a zygote between an unreduced tetraploid gamete and normal diploid gamete of tetraploid *C. × vernalis*. It is impossible to explain the mechanism of the appearance of aneuploids in this study. The ploidy variation in the progenies of 'Gaisen'

and 'Ume-ga-ka' might be established by natural crossing, contributing to a ploidy variation of *C. × vernalis*.

A significant difference ($P < 0.05$) of fluorescent intensities was found between triploid *C. japonica* 'Akebono' and some triploid *C. × vernalis* progenies, 'Gaisen' × *C. japonica* Nos. 2, 3 and 7 and 'Gaisen' open seedling No. 10. The three times value of fluorescent intensity of diploid *C. japonica* and two times value of that of triploid *C. japonica* were approximately 420 and 440, respectively. Although these values are theoretically to be equal to that of hexaploid, the values of actual hexaploid *C. sasanqua* were approximately 400. Tanaka (1985) inferred from Kondo (1975, 1978)'s observation that the genome structure of *C. japonica*, *C. sasanqua* and 'Gaisen' type *C. × vernalis* were AA, A'A'BBB'B' and AA'BB', respectively. The genome structure of triploid *C. japonica* 'Akebono', triploid *C. × vernalis* and 'Egao' type tetraploid *C. × vernalis* are thought to be AAA, AA⁰B⁰ and AAA⁰B⁰, respectively from Tanaka's supposition. It was, therefore, suggested that DNA contents of genomes A and B are different. The fluorescent intensity of single A genome was estimated to be 70–75 and that of B genome 55–60. The fluorescent intensity of 'Egao' type tetraploid *C. ×*

vernalis (AAA[○]B[○]) and hexaploid *C. sasanqua* (A'A'BBB'B') was calculated to be 285 and 390 in maximum, respectively, based on our estimation, and the values are both in accordance with those of the actual fluorescent intensity of them.

Conclusion

We confirmed the maternal inheritance of cpDNA in *Camellia* using the polymorphism of *atpH-atpI* region in some *C. × vernalis* cultivars, their open pollinated seedlings and artificially crossed progenies with *C. japonica*. Our study also demonstrated that flow cytometry makes rapid and efficient determination of ploidy levels possible in *Camellia* and gave some hypotheses to the mechanism causing ploidy variation in the progenies of *C. × vernalis*.

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