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Ploidy distribution and DNA content variations of Lonicera caerulea (Caprifoliaceae) in Japan

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Title: Ploidy distribution and DNA content variations of *Lonicera caerulea* (Caprifoliaceae) in Japan

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Abstract

Ploidy level and geographical distribution were investigated in Japanese Lonicera caerulea L. Flow cytometric analysis revealed the presence of DNA diploid and DNA tetraploid plants in Japan. Chromosome observation confirmed that diploid and tetraploid plants showed $2n = 2x = 18$ and $2n = 4x = 36$, respectively. The DNA diploid populations were found only in lowland mires, Betsukai, Bekenbeushi, Kushiro and Kiritappu located in eastern Hokkaido. On the other hand, DNA tetraploid populations were distributed in a wide area of Hokkaido, and mainland of Japan. The habitats of DNA tetraploid plants were lowland to alpine region. The DNA content measurement with flow cytometry revealed significant differences in the relative DNA contents among DNA tetraploid populations. The relative DNA content within DNA tetraploid populations varied 1.157-fold at maximum, and might correlate with altitude indicating that DNA contents were smaller as altitude increases. The wide area of distribution in various environments of DNA tetraploid plants suggested the adaptability of the tetraploid plants. Although diploid and tetraploid populations were found, no triploid was detected, indicating crossing difficulty between diploid and tetraploid as confirmed by crossing experiment.

Key words: chromosome number; DNA ploidy level; flow cytometry; genome size variation; geographical distribution; Lonicera caerulea
Introduction

Polyploidy has played a key role in plant evolution. Generally, polyploids differ from their progenitors in morphological, ecological, physiological and/or cytological characteristics. Consequently, polyploid may show exploitation of a new niche and reproductive isolation (reviewed by Ramsey and Schemske 2002). Thus, investigation of ploidy distribution is important issue to understand evolutionary adaptation and species specification in polyploid plant.

The genus *Lonicera* belongs to the family Caprifoliaceae, and comprises more than 200 species (Naugţemys et al. 2007). Some species in this genus are used as ornamental and edible plants. One of them, namely, *Lonicera caerulea* L., is generically known as blue honeysuckle; it is a shrub with edible fruits. *L. caerulea* is distributed in northern Eurasia, Japan and North America. The fruits of wild *L. caerulea* are traditionally used in Japan, Russia, and northern China (Anetai et al. 1996; Thompson and Chaovanalikit 2003). It has been cultivated as a berry crop, particularly in Russia and Japan, because the fruits are rich in nutrients such as polyphenolics and anthocyanins (Chaovanalikit et al. 2004; Terahara et al. 1993), and its cultivation is economical.

In natural populations of *L. caerulea*, diploid (*2n = 2x = 18*) and tetraploid (*2n = 4x = 36*) have been observed in Europe to Asia (Ammal and Saunders 1952). Limited information about ploidy levels of *L. caerulea* has been noted in Japanese populations. Solovyeva and Plehanova (2003) and Naugţemys et al. (2007) indicated taxonomic obscurity in *Lonicera* subsection *caeruleae* Rehder. So far, although different scientific names were used in published manuscripts, following findings were obtained. Hara (1983) investigated Caprifoliaceae of Japan and described tetraploid (*2n = 36*) in *L.*
caerulea subsp. edulis (Turcz.) Hultén var. edulis. Plekhanova et al. (1992) examined chromosome numbers and distribution area of Lonicera subsection caeruleae in far east of Asia to northern regions of Eurasia. They described L. emphyllocalyx Maxim. of Hokkaido, Japan as tetraploid (2n = 36). Plekhanova (2000) referred large population of L. caerulea L. (2n = 36) existing on Hokkaido, Japan. Thus, only tetraploid plants have been reported in Japan. However, little is known about the geographical distribution and ploidy of L. caerulea in Japan.

In the present study, we determined DNA ploidy levels by flow cytometry in the population of Japanese L. caerulea, and their chromosome numbers were observed. In addition, unexpected DNA content variations were found by flow cytometry in DNA tetraploid plants. Furthermore, geographical patterns of ploidy distribution and DNA content variations were discussed.

Materials and Methods

Plant materials

The sources of plant materials (Lonicera caerulea L.) used in this study are shown in Table 1. Lonicera caerulea is native to Japan and is called as ‘blue honeysuckle’ or locally ‘Haskap’. In Japan, this species is distributed on mountain areas in mainland, and in Hokkaido. Basic chromosome number is x = 9.

In order to evaluate ploidy levels of Japanese L. caerulea, we have collected wild species in Hokkaido. Sampling locations of natural populations could be entirely covered Hokkaido. One location was added from mainland of Japan. For this study, 450 wild L. caerulea individual plants from 21 populations were collected (Table 1, Fig. 1) in Japan between 2008 and 2009. Plants from natural populations of L. caerulea in the
eight locations in Hokkaido, namely, Betsukai, Bihoro Pass, Mt. Daisetsu, Mt. Piyashiri, Mt. Yokotsudake, Taiki, Kiritappu mire, and Yufutsu mire, were provided by the Hokkaido Forestry Research Institute (HFRI). The plants from Senjogahara mire in Tochigi Prefecture on the mainland were obtained from collections at Nikko Botanical Garden (NBG), Graduate School of Science, The University of Tokyo. Two plants from Kiritappu mire were obtained from collections at Hokkaido University. We also collected wild *L. caerulea* plants from Betsukai; Mt. Yokotsudake; Mt. Eboshidake; the Bekanbeushi, Kiritappu, Kushiro, Shibetsu, and Yufutsu mires in Hokkaido. We sampled leaves to analyze ploidy with a flow cytometry. For chromosome observations in root tips, we harvested scions from plants obtained from HFRI and the Bekanbeushi, Kushiro, Senjogahara, and Yufutsu mires, and propagated individual plants by cutting.

**DNA ploidy analysis and DNA content measurements using flow cytometry**

DNA ploidy level was analyzed using 4’, 6-diamidino-2-phenylindole (DAPI) stained flow cytometry (Partec PA; Partec GmbH, Münster, Germany). According to the concept of Suda et al. (2006), we use the term “DNA ploidy level”. The prefix ‘DNA’ indicates that ploidy levels are mostly inferred from nuclear DNA content without knowledge of the exact chromosome numbers. Fresh young leaf was chopped with a 0.2 ml drop of nuclei extraction buffer (CyStain UV precise P; Partec, Münster, Germany). After filtration through a 30-μm nylon mesh, crude nuclear samples were stained with 0.8 ml DAPI solution containing 10 mM Tris, 50 mM sodium citrate, 2 mM MgCl₂, 1% (w/v) PVP K-30, 0.1% (v/v) Triton X-100, and 2 mg l⁻¹ DAPI (pH 7.5) (Mishiba et al. 2000). After incubation for five minutes at room temperature, DNA ploidy level was analyzed with a flow cytometry. A total of more than 5000 cells were analyzed. Leaves
of Yufutsu No. 47 (2n = 4x = 36) were used as the internal standard. Totally, 450 plants from 21 populations were examined. For each sample, two replicates were performed.

We also performed the measurement of relative DNA content by flow cytometry as described above, but with slight modifications. The volume of nuclei extraction buffer applied was increased up to 0.4 ml. Fresh leaves of Capsicum annuum L. (cv. ‘Kyonami’) were used as the internal standard. Totally, 14 diploid plants from 3 populations (the Bekenbeushi and Kushiro mires, and Betsukai) and 38 tetraploid plants from 9 populations (Bihoro Pass, Mt. Daisetsu, Mt. Eboshidake, Mt. Yokotsudake, Taiki, and the Kiritappu, Senjogahara, Shibetsu, and Yufutsu mires) were investigated. Fresh young leaves were sampled at flowering season in May, 2009. The leaves of five individuals from Mt. Yokotsudake and Mt. Eboshidake were exceptionally sampled at fruiting season in August, 2009. A total of more than 5000 cells were analyzed in each measurement. Basically, when the coefficient of variation (CV) was within 5%, the analysis was accepted. Exceptionally, in measurement of some individuals from Mt. Yokotsudake, Mt. Eboshidake, Kushiro and Bekenbeushi mire, CV of more than 5% was accepted because of limited sample amount. For each sample, three replicates were performed within 1 day.

Relative DNA content was estimated by the ratio between the fluorescence intensity of the sample and internal standard: sample relative DNA content = sample peak mean / standard peak mean. The relative DNA content was then calculated for each sample as the average of the three replications.

Statistical tests were performed using the SPSS 16.0J program. In DNA diploid populations, differences of the relative DNA content was analyzed using non-parametric Mann-Whitney’s U-test, with P < 0.05 as the level of statistical significance. The
differences in relative DNA content among DNA tetraploid populations were analyzed using one-way analysis of variance (ANOVA) followed by Bonferroni’s test, with $P < 0.01$ as the level of statistical significance. The relationship between the relative DNA content and altitude was investigated by Pearson’s correlation coefficient test. The altitude is shown in Table 1. When the altitude ranged widely, the mean values were used for analysis.

**Chromosome analysis**

For counting the chromosome number, actively growing root tips were utilized for chromosome observation. Root tips were pretreated with ice water for 24 h and fixed with acetic acid:ethanol (1:3) at 4°C overnight. The fixed root tips were treatment with an enzyme mixture of 2% Cellulase Onozuka RS (Yakult Pharmaceutical Co. Ltd., Japan) and 0.5% Pectolyase Y-23 (Seishin Pharmaceutical Co. Ltd., Japan) (Shibata and Hizume 2002) in the citrate buffer (0.01 M citric acid and 0.01 M trisodium citrate dehydrate), pH 4.5, at 37°C for 20 min. After treatment with the enzyme mixture, the root tips were rinsed with distilled water. Then, the root tips were squashed with forceps in a drop of 45% acetic acid on slide glass, covered with cover glass, and squashed again. Cover glasses were removed by freezing glass slides in liquid nitrogen, slides were dried at 37°C. For staining, a drop of DAPI solution [0.233 g 1,4-diazabicyclo(2.2.2)-octane, 1 ml 0.2 M Tris-HCl, pH 8.0, 9 ml glycerol, 0.5 μg ml$^{-1}$ of DAPI] (Sahara et al. 2003) was put on the slide glass, and covered with cover glass. Preparations were observed under a fluorescence microscope (Aixo imager M1, Carl Zeiss, Oberkochen, Germany). Chromosome number was counted for 1–2 representative individuals from 11 populations. For each sample, 5–10 measurements
were recorded.

Results

DNA ploidy level analysis by flow cytometry, and chromosome counts

We measured the DNA ploidy levels of 450 plants from 21 locations with flow cytometry, and we obtained two distinct groups with regard to the relative DNA content (Table 1). All plants harvested in the Bekanbeushi and Kushiro mires showed 2Cx DNA content (Table 1, Fig. 2). In this study, description of genome size is expressed as C_x-value, which is designated as DNA content of a monoploid genome with chromosome number x, according to Greilhuber et al. (2005). Of the plants from Kiritappu mire and Betsukai, two types of DNA ploidy levels were obtained. One strain from Kiritappu mire from the collection at Hokkaido University had 2Cx DNA content, whereas other plants from Kiritappu mire had 4Cx DNA content (Table 1). In addition, one strain from Betsukai (population code, BT-1) and one strain from Betsukai from the collection at HFRI (population code, BT-2) had 2Cx DNA content, whereas other Betsukai strains from HFRI had 4Cx DNA content (Table 1). Plants of all strains with 4Cx DNA content were found in Bihoro Pass; Mt. Daisetsu; Mt. Eboshidake; Mt. Piyashiri; Mt. Yokotsudake; Taiki; and the Yufutsu, Senjogahara, and Shibetu mires (Table 1). Flow cytometric analysis revealed that 65 plants had 2Cx DNA content, while 385 had 4Cx DNA content (Table 1).

The results of the flow cytometric analysis were confirmed by determining the chromosome number. The chromosome number was counted for representative individuals from 11 populations. This analysis confirmed that diploid (2n = 2x = 18) and tetraploid (2n = 4x = 36) populations existed in the natural populations of L. caerulea
(Table 1, Fig. 3). The populations from the Bekanbeushi and Kushiro mires and one strain from Kiritappu mire obtained from Hokkaido University were diploid (Table 1, Fig. 3a). Other strains from Kiritappu mire from Hokkaido University and HFRI were tetraploid (Table 1, Fig. 3b). In addition, all the plants obtained from other populations were tetraploid (Table 1).

**DNA content variation**

Next, we measured of relative DNA content in *L. caerulea* by using fresh leaves of *Capsicum annuum* (cv. ‘Kyonami’) as an internal standard (Fig. 4). The relative DNA content in DNA diploid and DNA tetraploid populations is shown in Table 2. The mean relative DNA content in 14 individuals from three DNA diploid populations (the Bekanbeushi and Kushiro mires and Betsukai) was 0.257. The mean relative DNA content in 38 individuals from 9 DNA tetraploid populations (Bihoro Pass, Mt. Daisetsu, Mt. Eboshidake, Mt. Yokotsudake, Taiki, and the Kiritappu, Senjogahara, Shibetsu, and Yufutsu mires) was 0.528. The variation of relative DNA content within same DNA ploidy level was larger in DNA tetraploid populations than that of DNA diploid populations. The coefficient of variation of relative DNA content was 0.88 and 4.55 in DNA diploid and DNA tetraploid populations, respectively. The relative DNA content within DNA diploid populations varied 1.024-fold (from 0.254 to 0.260) at maximum, while those of DNA tetraploid populations varied 1.157-fold (from 0.492 to 0.569).

The relative DNA content of individual DNA tetraploid populations is shown in Table 3 and Figure 5. The relative DNA content within DNA tetraploid populations varied 1.157-fold at maximum (Table 2), however, only slight variation (1.002 to 1.052-fold) was observed within a single population (Table 3). The coefficient of
variation of relative DNA content was 0.2–1.8 within a single population (Table 3). The differences in relative DNA content among five DNA tetraploid populations (Mt. Daisetsu, Mt. Yokotsudake, and the Kiritappu, Shibetsu, and Yufutsu mires) were analyzed using one-way ANOVA followed by Bonferroni’s test. The populations from Bihoro Pass, Mt. Eboshidake, Senjogahara mire, and Taiki were excluded from the statistical analysis because of low sample number. The Bonferroni test ($P < 0.01$) showed significant differences among DNA tetraploid populations (Table 3, Fig. 5). DNA tetraploid populations could divide into three groups; 1) Mt. Daisetsu, 2) Mt. Yokotsudake and Yufutsu mire, 3) Kiritappu mire and Shibetsu mire. In DNA diploid populations, no significant differences between Bekanbeushi mire and Kushiro mire were found from the statistical analysis by Mann-Whitney’s U-test. Population from Betsukai was excluded from the statistical analysis because of low sample number.

The relationship between the relative DNA content of the DNA tetraploid populations and altitude was determined with in the 38 individuals from 9 DNA tetraploid populations. The relative DNA content in the DNA tetraploids was significantly correlated with altitude ($r = -0.684$, $P < 0.001$, $n = 38$), indicating that DNA contents were smaller as altitude increases (Fig. 6).

**Discussion**

**Distribution of diploid and tetraploid plants**

In the present study, ploidy study revealed that diploid ($2n = 2x = 18$) and tetraploid ($2n = 4x = 36$) plants of *L. caerulea* were distributed in Japan. Worldwide, the diploid and tetraploidal of *Lonicera* subsection *Caeruleae* were found in Eurasia to North America (Plekhanova et al. 1992; Solovyeva and Plehanova 2003). So far, diploid species were
found in only four locations, such as Central Asian, Transbaikalian, South-Ussurian (Plekhanova 2000; Plekhanova et al. 1992), and Alberta, Canada (Solovyeva and Plehanova 2003). In addition to previous reports, our study found out distribution of diploid *L. caerulea* in Hokkaido, Japan.

In previous reports (Plekhanova 2000; Plekhanova et al. 1992), dominant tetraploid populations of *Lonicera* subsection *Caeruleae* were found in Eurasia. It has also been shown that tetraploid plants spread to more northern and alpine habitats than diploid plants do. Diploid plants occurred on territories unaffected by glaciers, and are located in proximity to Central China, the center of origin of bushy *Lonicera* (Plekhanova 2000; Plekhanova et al. 1992). In the case of Japan, DNA tetraploid plants were distributed in wide area of Hokkaido, and mainland of Japan, while DNA diploid plants occurred only in Betsukai, Bekanbeushi mire, Kiritappu mire and Kushiro mire located in eastern Hokkaido. The habitats of DNA tetraploid plants were lowland mires, around lowland mires, developed mires, and mountain regions (mountain mire or rocky area) (Table 1). On the other hand, DNA diploid plants were found only in lowland mire (Table 1). In many plant species, the geographical distributions of diploid and tetraploid plants differ (Lewis 1980). In the present study, the populations in the mountainous habitats (Bihoro Pass, Mt. Daisetsu, Mt. Eboshidake, Mt. Piyashiri, Mt. Yokotsudake, and Senjogahara mire) were only tetraploid, suggesting that these populations prefer in mountainous regions. In addition, the wide area of distribution in various environments, similar to that shown in previous studies in Eurasia (Plekhanova 2000; Plekhanova et al. 1992), suggested the adaptability of the tetraploid plants.

Natural triploid populations of *Lonicera* subsection *Caeruleae* have not been reported. Similarly, no triploid plant was observed in the present study. In general,
triploid plants are produced by the crossing between diploid and tetraploid or by the union of unreduced (2n) and reduced (n) gametes in diploid plants (reviewed by Ramsey and Schemske 1998). However, crosses between diploid and tetraploid plants are often unsuccessful because of abnormal seed development (reviewed by Haig and Westoby 1991; reviewed by Ramsey and Schemske 1998). To investigate the possibility of triploid induction, we attempted to cross between diploid and tetraploid L. caerulea plants. The number of mature seeds obtained from the reciprocal cross was substantially less than that of 2x-2x and 4x-4x plants (unpublished data). Low fruit and seed setting as well as the low viability of seedlings from crosses between diploid and tetraploid individuals of Lonicera subsection Caeruleae were also reported by Plekhanova (2000). The lack of a triploid population may be because of the low success of seed production and reduced survival of triploid seedlings.

**DNA content variations among DNA tetraploid populations**

In the present study, variations in the relative DNA content were detected by flow cytometry among the DNA tetraploid populations (Table 3, Fig. 5). Genomic size variation was first mentioned by Evans et al. (1966). It has been reported at the interspecific and intraspecific levels (Knight et al. 2005). Genomic size variations within the same species have been reported in some species such as those belonging to Poaceae (Ceccarelli et al. 1992; Creber et al. 1994; Reeves et al. 1998; Sugiyama and Yamada 2003; Šmarda and Bureš 2006) and Asteraceae (Suda et al. 2007). These studies have shown the correlation of genomic size with geographical condition, climate, or vegetation. The correlation between genomic size and altitude was reported in tetraploid populations of Dactylis glomerata L. (Creber et al. 1994; Reeves et al. 1998),
i.e., the genomic size decreases with increasing altitude. Similarly, the relative DNA contents in the DNA tetraploids were smaller as altitude increases in *L. caerulea* (Fig. 6). Murray (2005) stated that varying DNA content could be correlated with eco-geographic variations, suggesting that the varying DNA content is adaptive and may be an example of incipient speciation. The genome downsizing following polyploid formation may be a widespread phenomenon (Leitch and Bennett 2004). The changes of DNA content could be caused by activation of transposons, homoeologous recombination, or elimination of specific DNA sequences (Leitch and Bennett 2004). Solovyeva and Plehanova (2003) examined the karyotype of tetraploid *Lonicera* subsection *Caeruleae* plants and observed variations in chromosome size, number of satellites, and telomeres, which might confirm our results regarding the DNA content variability in DNA tetraploids.

However, DNA content variation could be explain by other factors, such as ancient introgression with related species (Suda et al., 2007), or dissimilar levels of secondary metabolites that may have interfered with DNA fluorochromes (Kubátová et al., 2008). Aneuploidy is also one of the possible explanation about DNA content variation, although no aneuploidy was observed in the present study as well as in previous reports of *Lonicera* subsection *Caeruleae*. Further studies including phylogenetic analysis and analyses of biochemical compounds affecting DNA staining are required for reliable conclusions about relative DNA content variation in *L. caerulea*.

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nutraceutical values of blue honeysuckle (*Lonicera caerulea*) in Oregon, USA. Acta Hortic 626:65–72
Table 1. Geographical locations and ploidy levels of *Lonicera caerulea* in Japan.

<table>
<thead>
<tr>
<th>Population codes</th>
<th>Populations</th>
<th>Habitat</th>
<th>Geographical coordinates</th>
<th>DNA Ploidy level</th>
<th>Chromosome number (2n)</th>
<th>Voucher number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Latitude (°N)</td>
<td>Longitude (°E)</td>
<td>Altitude (m)</td>
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<tr>
<td>Mainland SE'</td>
<td>Senjogahara mire</td>
<td>Area surrounding Senjogahara mire</td>
<td>36.46</td>
<td>139.26</td>
<td>1390–1400</td>
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<td>Hokkaido BK-1</td>
<td>Bekenbeushi mire</td>
<td>Alder forest</td>
<td>43.10</td>
<td>144.51</td>
<td>0–10</td>
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<td>BK-2</td>
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<td>BK-3</td>
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<td>Developed area along R44</td>
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<td>Betsukai</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>3</td>
<td>1</td>
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<tr>
<td>BKp</td>
<td>Bihoro Pass</td>
<td>Rocky area</td>
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<td>144.15</td>
<td>500–525</td>
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<td>KI-1</td>
<td>Kiritappu mire</td>
<td>High moor near estuary of Biwase River</td>
<td>43.03</td>
<td>145.04</td>
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<td>Kiritappu mire</td>
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<td>KI-3p</td>
<td>Kiritappu mire</td>
<td>—</td>
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<td>145.01–145.06</td>
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<tr>
<td>KI-4p</td>
<td>Kiritappu mire</td>
<td>—</td>
<td>43.02–43.08</td>
<td>145.01–145.06</td>
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<td>KU</td>
<td>Kushiro mire</td>
<td>High moor, Alder forest adjacent to high moor</td>
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<td>Mt. Daisetsu</td>
<td>Around mountainous mire</td>
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<td>142.53</td>
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<td>ME</td>
<td>Mt. Eboshidake</td>
<td>Rocky area around moutaintop</td>
<td>41.55</td>
<td>140.47</td>
<td>1070–1078</td>
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<td>MP</td>
<td>Mt. Pyashiri</td>
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<td>142.33–142.36</td>
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<td>MY-1</td>
<td>Mt. Yokotsudake</td>
<td>Rocky area around moutaintop</td>
<td>41.56</td>
<td>140.46</td>
<td>1120–1150</td>
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<td>MY-2p</td>
<td>Mt. Yokotsudake</td>
<td>Rocky area around moutaintop</td>
<td>41.56</td>
<td>140.46</td>
<td>1120–1150</td>
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<td>SH</td>
<td>Shibetsu mire</td>
<td>High moor in Pho River Historic &amp; Nature Park</td>
<td>43.41</td>
<td>145.06</td>
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<td>TAp</td>
<td>Taiki</td>
<td>Alder forest, <em>Quercus crispula</em> and <em>Q. dentata</em> forest</td>
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<td>143.28–143.30</td>
<td>0–70</td>
<td>4</td>
</tr>
<tr>
<td>YU-1</td>
<td>Yufutsu mire</td>
<td>Area drained by industrial development</td>
<td>42.40</td>
<td>141.45</td>
<td>5</td>
<td>200</td>
</tr>
<tr>
<td>YU-2p</td>
<td>Yufutsu mire</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>11</td>
<td>0</td>
</tr>
</tbody>
</table>

*The plants were provided by Nikko Botanical Garden (NBG). The habitat, geographical coordinates, and altitude were determined from...*
sample recordings from NBG.

b All the plants were provided by the Hokkaido Forestry Research Institute (HFRI). These plants were collected by T. Sato. The habitat, geographical coordinates, and altitude were determined from Sato’s report (1985).

c All the plants were obtained from collections at Hokkaido University. These plants were collected by T. Sato. The geographical coordinates and altitude were determined from Sato’s report (1985).

Dashes indicate no data.
Table 2. Relative DNA content in DNA diploid and DNA tetraploid populations estimated by flow cytometry.

<table>
<thead>
<tr>
<th>Ploidy level</th>
<th>Number of population</th>
<th>n</th>
<th>Mean ± SD</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Difference within population (fold)$^a$</th>
<th>Coefficient of variation$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA diploid</td>
<td>3</td>
<td>14</td>
<td>0.257 ± 0.002</td>
<td>0.254</td>
<td>0.260</td>
<td>1.024</td>
<td>0.88</td>
</tr>
<tr>
<td>DNA tetraploid</td>
<td>9</td>
<td>38</td>
<td>0.528 ± 0.024</td>
<td>0.492</td>
<td>0.569</td>
<td>1.157</td>
<td>4.55</td>
</tr>
</tbody>
</table>

Relative DNA content was estimated by the ratio between the fluorescence intensity of a sample and internal standard: sample relative DNA content = sample peak mean / standard peak mean.

$^a$ Differences of relative DNA content (fold) within population was calculated as Maximum relative DNA content / Minimum relative DNA content.

$^b$ Coefficient of variation of relative DNA content was calculated as (SD / Mean) × 100.
### Table 3. Relative DNA content in DNA tetraploid populations as estimated by flow cytometry.

<table>
<thead>
<tr>
<th>Populations</th>
<th>n</th>
<th>Mean ± SD</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Difference within population (fold)</th>
<th>Coefficient of variation(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Senjogahara mire</td>
<td>1</td>
<td>0.511</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Bihoro Pass</td>
<td>1</td>
<td>0.496</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Kiritappu mire</td>
<td>6</td>
<td>0.564 ± 0.005</td>
<td>c</td>
<td>0.557</td>
<td>0.569</td>
<td>1.022</td>
</tr>
<tr>
<td>Mt. Daisetsu</td>
<td>5</td>
<td>0.497 ± 0.004</td>
<td>a</td>
<td>0.492</td>
<td>0.501</td>
<td>1.018</td>
</tr>
<tr>
<td>Mt. Eboshidake</td>
<td>2</td>
<td>0.510 ± 0.001</td>
<td>–</td>
<td>0.510</td>
<td>0.511</td>
<td>1.002</td>
</tr>
<tr>
<td>Mt. Yokotsudake</td>
<td>6</td>
<td>0.518 ± 0.007</td>
<td>b</td>
<td>0.513</td>
<td>0.531</td>
<td>1.035</td>
</tr>
<tr>
<td>Shibetsu mire</td>
<td>6</td>
<td>0.557 ± 0.010</td>
<td>c</td>
<td>0.537</td>
<td>0.565</td>
<td>1.052</td>
</tr>
<tr>
<td>Taiki</td>
<td>1</td>
<td>0.535</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Yufutsu mire</td>
<td>10</td>
<td>0.518 ± 0.004</td>
<td>b</td>
<td>0.513</td>
<td>0.529</td>
<td>1.031</td>
</tr>
</tbody>
</table>

Relative DNA content was estimated by the ratio between the fluorescence intensity of a sample and internal standard: sample relative DNA content = sample peak mean / standard peak mean.

\(a\) Differences of relative DNA content (fold) within population was calculated as Maximum relative DNA content / Minimum relative DNA content.

\(b\) Coefficient of variation of relative DNA content was calculated as \((\text{SD} / \text{Mean}) \times 100\).

Means ± SD followed by the same letter are not significantly different (Bonferroni’s test, \(P < 0.01\)).

Dashes indicate no data.
Figure Legends

**Fig. 1.** Map of Japan showing sampling locations of *Lonicera caerulea* described in Table 1. The Circles indicate sampling locations. The white and black circles show the diploid and tetraploid populations, respectively. The letters indicate the population codes described in Table 1. The squares indicate other habitats of *L. caerulea* described in previously reports (Hara 1983; Sato 1985).

**Fig. 2.** Histograms of the relative fluorescence intensity of nuclei isolated from the leaves of *L. caerulea*. Peak of 2Cx and 4Cx were from Kushiro mire strain and internal standard, respectively. The leaves of Yufutsu No. 47 (2n = 4x = 36) were used as the internal standard.

**Fig. 3.** Metaphase chromosomes of *L. caerulea* plants obtained from individual locations. (a) Kushiro mire (2n = 2x = 18). (b) Kiritappu mire (2n = 4x = 36). Bars = 5 μm.

**Fig. 4.** Histograms of the relative fluorescence intensity of nuclei isolated from the leaves of *L. caerulea* and internal standard. (a) DNA diploid plants from Kushiro mire (b) DNA tetraploid plants from Shibetsu mire. The peak of *L. caerulea* is marked with an asterisk. Fresh leaves of *Capsicum annuum* (cv. ‘Kyonami’) were used as internal standard.

**Fig. 5.** Box plot graph of relative DNA content of DNA tetraploid plants from different populations. Thick line across the box indicates the median, the box contains the values
between the 25% to 75%, and the whiskers extend to the highest and lowest values, excluding the outliers. The asterisk represents the outlier. The same letters above the box-plots indicate that there is no significant differences (Bonferroni’s test, $P < 0.01$). Dashes indicate no data.

**Fig. 6.** Relationship of relative DNA contents with altitude. The samples were 38 individuals from nine DNA tetraploid populations described in Table 3. The relative DNA contents were negatively correlated with altitude ($r = -0.684$, $P < 0.001$, $n = 38$).
Figure 1

Mainland

Hokkaido
Figure 2

<table>
<thead>
<tr>
<th>Ploidy</th>
<th>Fluorescence intensity</th>
<th>CV</th>
</tr>
</thead>
<tbody>
<tr>
<td>2Cx</td>
<td>23.35</td>
<td>3.21</td>
</tr>
<tr>
<td>4Cx</td>
<td>49.11</td>
<td>2.55</td>
</tr>
</tbody>
</table>
**Figure 4**

(a) Histogram showing the number of nuclei against fluorescence intensity for **Capsicum annuum**. The peak fluorescence intensity and CV for sample (*) and **Capsicum** are listed in the table below.

<table>
<thead>
<tr>
<th>Peak</th>
<th>Fluorescence intensity</th>
<th>CV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample (*)</td>
<td>45.67</td>
<td>3.83</td>
</tr>
<tr>
<td><strong>Capsicum</strong></td>
<td>181.49</td>
<td>2.89</td>
</tr>
</tbody>
</table>

(b) Similar histogram and table as above, but for a different sample.

<table>
<thead>
<tr>
<th>Peak</th>
<th>Fluorescence intensity</th>
<th>CV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample (*)</td>
<td>56.18</td>
<td>3.12</td>
</tr>
<tr>
<td><strong>Capsicum</strong></td>
<td>100.14</td>
<td>2.75</td>
</tr>
</tbody>
</table>
Figure 5
Figure 6