1	Genome sequence of Hydrangea macrophylla and its application in analysis of the double
2	flower phenotype
3	
4	Authors
5	Nashima K ^{*1} , Shirasawa K ^{*2} , Ghelfi A ² , Hirakawa H ² , Isobe S ² , Suyama T ³ , Wada T ³ , Kurokura T ⁴ ,
6	Uemachi T ⁵ , Azuma M ¹ , Akutsu M ⁶ , Kodama M ⁶ , Nakazawa Y ⁶ , Namai K ⁶
7	
8	1. College of Bioresource Sciences, Nihon University, Kameino 1866, Fujisawa, Kanagawa, 252-
9	0880 Japan
10	2. Kazusa DNA Research Institute, Kazusa-Kamatari 2-6-7, Kisarazu, Chiba, 292-0813 Japan
11	3. Fukuoka Agriculture and Forestry Research Center, Yoshiki 587, Chikushino, Fukuoka, 818-8549
12	Japan
13	4. Faculty of Agriculture, Utsunomiya University, Mine 350, Utsunomiya, Tochigi, 321-8505 Japan
14	5. School of Environmental Science, University of Shiga Prefecture, Hassakacho 2500, Hikone,
15	Shiga, 522-0057 Japan
16	6. Tochigi Prefectural Agricultural Experimental Station, Kawarayacho 1080, Utsunomiya, Tochigi,
17	320-0002 Japan
18	
19	*equally contributed as first author
20	Corresponding author: Nashima K
21	College of Bioresource Sciences, Nihon University, Kameino 1866, Fujisawa, Kanagawa, 252-0880
22	Japan
23	Tel: +81-466-84-3507
24	Mail: nashima.kenji@nihon-u.ac.jp

25 Abstract

26	Owing to its high ornamental value, the double flower phenotype of hydrangea (Hydrangea
27	macrophylla) is one of its most important traits. In this study, genome sequence information was
28	obtained to explore effective DNA markers and the causative genes for double flower production in
29	hydrangea. Single molecule real-time sequencing data followed by a HiC analysis was employed. The
30	resultant haplotype-phased sequences consisted of 3,779 sequences (2.256 Gb in length and N50 of
31	1.5 Mb), and 18 pseudomolecules comprising 1.08 Gb scaffold sequences along with a high-density
32	SNP genetic linkage map. Using the genome sequence data obtained from two breeding populations,
33	the SNPs linked to double flower loci (D_{jo} and D_{su}), were discovered for each breeding population.
34	DNA markers J01 linked to D_{jo} and S01 linked to D_{su} were developed, and these could be used
35	successfully to distinguish the recessive double flower allele for each locus respectively. The <i>LEAFY</i>
36	gene was suggested as the causative gene for D_{su} , since frameshift was specifically observed in double
37	flower accession with d_{su} . The genome information obtained in this study will facilitate a wide range
38	of genomic studies on hydrangea in the future.
39	
40	Keywords:
41	Hydrangea, double flower, de novo genome sequencing, DNA marker
42	
43	
44	
45	
46	
47	
48	

49 **1. Introduction**

50 Hydrangea macrophylla (Thunb.) Ser., commonly known as hydrangea, originated in Japan, 51 and since it is the place of origin, there are rich genetic resources for this plant in Japan. Wild 52 hydrangea accessions with superior characteristics have been bred to create attractive cultivars, and it 53 has a long history of use as an ornamental garden plant in temperate regions. There are both decorative 54 and non-decorative flowers in an inflorescence. Decorative flowers have large ornamental sepals that 55 attract pollinators, whereas non-decorative flowers have inconspicuous perianths that instead play a major role in seed production¹⁻³. In hydrangea, there are two types of decorative flower phenotype: 56 57 single flower and double flower. Single flowers generally have four petaloid sepals per decorative 58 flower, while this number in double flowers is approximately fourteen. Double flowers do not have 59 stamens or petals⁴. Therefore, petals and stamens would be converted to petaloid sepals since number 60 of petaloid sepals are increased and stamens and petals are lost. Because of their high ornamental value, 61 producing double flower is an important breeding target in hydrangea cultivation.

62 To obtain double flower progenies, the double flower cultivars 'Sumidanohanabi' (Figure 63 1A) and 'Jogasaki' (Figure 1B) were crossbred in Japan⁴. Previous studies have suggested that double 64 flower phenotype is a recessive characteristic controlled by a single major gene^{4,5}. Suyama et al.⁴ 65 found that crosses between the progeny of 'Sumidanohanabi' and the progeny of 'Jogasaki' produced 66 only single flower descendants. Thus, it was also suggested that genes controlling the double flower 67 phenotype are different⁴. While Suyama et al.⁴ suggested that a single locus with different double 68 flower alleles controls the phenotype, Waki et al.⁵ speculated that two different loci control double 69 flower production individually. Therefore, it is not clear whether a single locus or two loci control the 70 phenotype. We term the double flower locus D_{su} as the locus controlling the double flower phenotype 71 of 'Sumidanohanabi' and the double flower locus D_{io} as the locus controlling the double flower 72 phenotype of 'Jogasaki.' Waki et al.⁵ identified D_{su} on the genetic linkage map. They also found that

the DNA marker STAB045 was the nearest marker to D_{su} , and that STAB045 could help in distinguishing flower phenotype with a 98.6% fitting ratio⁵. Contrarily, D_{jo} has not been identified, and the DNA marker linked to D_{jo} has not been developed. It is still not known whether D_{jo} and D_{su} are at the same loci.

77 The mechanisms and genes controlling double flower phenotype in hydrangea have not been 78 clarified. Waki et al.⁵ hypothesized that the mutation of C-class genes could be associated with the 79 double flower phenotype of 'Sumidanohanabi', since the C-class gene mutant of Arabidopsis thaliana 80 and C-class gene-repressed petunias produce double flowers⁶. However, the double flower phenotype 81 of hydrangea is morphologically different from that of A. thaliana and petunia-petals and stamens 82 would be converted to petaloid sepals, while stamens converted to petals in A. thaiana and petunia. 83 This suggests that the genes controlling double flower production in hydrangea are different from 84 corresponding genes in other plant species. Identification of the genes controlling double flower 85 production in hydrangea could reveal novel regulatory mechanisms of flower development.

86 Genomic information is essential for DNA marker development and identification of genes controlling specific phenotypes. However, no reference genome sequence is publicly available for 87 88 hydrangea so far. Although a genome assembly of hydrangea (1.6 Gb) using only short-read data has 89 been reported⁷, the resultant assembly is so fragmented that it comprises 1,519,429 contigs with an 90 N50 size of 2,447 bp and has not been disclosed. Improved, advanced long-read technologies and 91 bioinformatics methods would make it possible to determine the sequences of complex genomes. An 92 assembly strategy for single molecule real-time sequencing data followed by a HiC analysis has been 93 developed to generate haplotype-phased sequences in heterozygous regions of diploid genomes⁸. 94 Genome sequences at the chromosome level could be obtained with a HiC clustering analysis⁹ as well as with a genetic linkage analysis¹⁰. Such genomic sequence will provide basic information to identify 95 96 genes and DNA markers of interest, and to discover allelic sequence variations. In this study, we

97	constructed the genomic DNA sequence, obtained SNPs information, and performed gene prediction.
98	We also developed DNA markers linked to D_{jo} using SNP information obtained by double digest
99	restriction site associated DNA sequence (ddRAD-Seq) analysis of breeding population 12GM1,
100	which segregated double flower phenotypes of D_{jo} . In addition, we attempted to identify the causative
101	genes for D_{io} and D_{su} .

102

103 **2. Materials and Methods**

104 **2.1. De novo assembly of the hydrangea genome**

105 For genomic DNA sequencing, H. macrophylla 'Aogashima-1,' collected from Aogashima 106 island of the Izu Islands in Tokyo Prefecture, Japan, was used. Genomic DNA was extracted from the 107 young leaves with Genomic-Tip (Qiagen, Hilden, Germany). First, we constructed a sequencing 108 library (insert size of 500 bp) with TruSeq DNA PCR-Free Library Prep Kit (Illumina, San Diego, CA, 109 USA) to sequence on HiSeqX (Illumina). The size of the 'Aogashima-1' genome was estimated using 110 Jellyfish v2.1.4¹¹. After removing adapter sequences and trimming low-quality reads, high-quality reads were assembled using Platanus¹². The resultant sequences were designated HMA r0.1. 111 112 Completeness of the assembly was assessed with sets of BUSCO v.1.1b¹³.

113 Next, a SMRT library was constructed with SMRTbell Express Template Prep Kit 2.0 114 (PacBio, Menlo Park, CA, USA) in accordance with the manufacture's protocol and sequenced with 115 SMRT Cell v2.1 on a Sequel System (PacBio). The sequence reads were assembled using FALCON 116 $v.1.8.8^{14}$ to generate primary contig sequences and to associate contigs representing alternative alleles. 117 Haplotype - resolved assemblies (i.e. haplotigs) were generated using FALCON-Unzip v.1.8.8¹⁴. 118 Potential sequence errors in the contigs were corrected twice with ARROW v.2.2.1 implemented in 119 SMRT Link v.5.0 (PacBio) followed by one polishing with Pilon¹⁵. Subsequently, a HiC library was 120 constructed with Proximo Hi-C (Plant) Kit (Phase Genomics, Seattle, WA, USA) and sequenced on HiSeqX (Illumina). After removing adapter sequences and trimming low-quality reads, high-quality
HiC reads were used to generate two haplotype-phased sequences from the primary contigs and
haplotig sequences with FALCON-Phase⁸.

124 To validate the accuracy of the sequences, we developed a genetic map based on SNPs, 125 which were from a ddRAD-Seq analysis on an F2 mapping population (n = 147), namely 12GM1, 126 maintained at the Fukuoka Agriculture and Forestry Research Center, Japan. The 12GM1 population 127 was generated from a cross between 'Posy Bouquet Grace' (Figure 1C) and 'Blue Picotee Manaslu' 128 (Figure 1D). Genomic DNA was extracted from the leaves with DNeasy Plant Mini Kit (Qiagen). A ddRAD-Seq library was constructed as described in Shirasawa et al.¹⁶ and sequenced with HiSeq4000. 129 Sequence reads were processed as described by Shirasawa et al.¹⁶ and mapped on the HMA r1.2 as a 130 131 reference. From the mapping alignment, high-confidence biallelic SNPs were obtained with the 132 following filtering options: --minDP 5 --minQ 10 --max-missing 0.5. The genetic map was constructed 133 with Lep-Map3¹⁷.

Potential mis-jointed points in the phase 0 and 1 sequences of HMA_r1.2 were cut and rejoined, based on the marker order in the genetic map, for which we employed ALLMAPS¹⁸. The resultant sequences were named HMA_r1.3.pmol, as two haplotype-phased pseudomolecule sequences of the 'Aogashima-1' genome. Sequences that were unassigned to the genetic map were connected and termed chromosome 0.

139

140 **2.2 Gene prediction**

For gene prediction, we performed Iso-Seq analysis. Total RNA was extracted from 12 samples of 'Aogashima-1': flower buds (2 stages); decorative flowers (2 stages); colored and colorless non-decorative flowers; fruits; shoots; roots; buds, and one-day light-intercepted leaves and buds. In addition, the 29 samples listed in Supplementary Table S1 were included. Iso-Seq libraries were

145 prepared with the manufacture's Iso-Seq Express Template Preparation protocol, and sequenced on a 146 Sequel System (PacBio). The raw reads obtained were treated with ISO-Seq3 pipeline, implemented 147 in SMRT Link v.5.0 (PacBio) to generate full-length, high-quality consensus isoforms. In parallel, 148 RNA-Seq data was also obtained from the 16 samples listed in Supplementary Table S1. Total RNA 149 extracted from the samples was converted into cDNA and sequenced on HiSeq2000, Hiseq2500 150 (Illumina), and NovaSeq6000 (Illumina). The Iso-Seq isoform sequences and the RNA-Seq short-151 reads were employed for gene prediction.

152 To identify putative protein-encoding genes in the genome assemblies, ab-initio-, evidence-, 153 and homology-based gene prediction methods were used. For this prediction, unigene sets generated 154 from 1) the Iso-Seq isoforms; 2) de novo assembly of the RNA-Seq short-reads with Trinity-v2.4.0¹⁹; 155 3) peptide sequences predicted from the genomes of Arabidopsis thaliana, Arachis hypogaea, 156 Cannabis sativa, Capsicum annuum, Cucumis sativus, Populus trichocarpa, and Quercus lobata; and 157 4) *ab-initio* genes, were predicted with Augustus-v $3.3.1^{20}$. The unigene sequences were aligned onto 158 the genome assembly with BLAT²¹ and genome positions of the genes were listed in general feature 159 format version 3 with blat2gff.pl (https://github.com/vikas0633/perl/blob/master/blat2gff.pl). Gene 160 annotation was performed with Hayai-annotation Plants²². Completeness of the gene prediction was 161 assessed with sets of BUSCO v4.0.6¹³.

162

163 2.3 Detection of SNPs linked to the double flower phenotype

164 For identification of SNPs linked to double flower loci D_{jo} and D_{su} , ddRAD-Seq data 165 analysis was performed. ddRAD-Seq data of the 12GM1 population described above was used to 166 identify D_{io} . For identification of SNPs linked to double flower locus D_{su} , KF population⁵—93 F2 167 specimens of 'Kirakiraboshi' (Figure 1E) and 'Frau Yoshimi' (Figure 1F)-were used for ddRAD-Seq 168 analysis. The KF population was maintained at Tochigi Prefectural Agricultural Experimental Station,

Japan. ddRAD-Seq analysis of the KF population was performed using the same method used for the12GM1 population.

171 ddRAD-Seq data of the 12GM1 and KF populations were processed as follows: Low-quality sequences were removed and adapters were trimmed using Trimmomatic-0.36²³ (LEADING:10, 172 173 TRAILING:10, SLIDINGWINDOW:4:15, MINLEN:51). BWA-MEM (version 0.7.15-r1140) was 174 used for mapping onto genome sequence. The resultant sequence alignment/map format (SAM) files 175 were converted to binary sequence alignment/map format files and subjected to SNP calling using the 176 mpileup option of SAMtools²⁴ (version 1.4.1) and the view option of BCFtools (parameter -vcg). If 177 the DP of called SNP in individuals was under 5%, the genotype was treated as missing. SNPs with 178 5% or more of missing genotype were filtered out. Each SNP was evaluated, fitting ratios with the 179 flower phenotype.

180

181 **2.4 DNA marker development and analysis for D**_{jo}

182 A CAPS marker was designed based on SNP (Scaffold:0008F-2, position: 780104) that was 183 completely linked to the double flower locus D_{io} . Primers were designed using Primer3²⁵ under 184 conditions with product size ranging from 150 to 350 bp, primer size from 18 to 27 bp, and primer 185 TM from 57 to 63°C. Primer sequences of the designed CAPS marker named J01 were: Forward: 5'-186 CTGGCAGATTCCTCCTGAC-3' and Reverse: 5'-TATTTCCTTGGGGAGGCTCT-3'. PCR assays 187 were done in a total volume of 10 μ L, containing 5 μ L of GoTaq Master Mix (Promega, Mdison, WI, 188 USA), 1 mM each of forward and reverse primer, and 5 ng of template DNA. The PCR conditions 189 were 94°C for 2 min, 35 cycles of denaturation at 94°C for 1 min, annealing at 55°C for 1 min, and 190 extension at 72°C for 1 min; and a final extension step at 72°C for 3 min. Then, restriction enzyme 191 assay was done in a total volume of 10 µL, containing 5 µL of PCR product, ten units of restriction 192 enzyme TaqI (New England Biolabs, Ipswich, MA, USA), and 1 µL of cut smart buffer. Restriction

194 GRRED (Biocraft, Tokyo, Japan) and separated in 1.5% (w/v) agarose gel in TAE buffer. Designed 195 CAPS marker J01 was applied to the 12GM1 population, 14GT77 population (64 F2 specimens of 196 'Posy Bouquet Grace' × 'Chibori') and the 15IJP1 population (98 F1 specimens of 'Izunohana' × 197 03JP1) that segregate the double flower locus D_{io} .

198

199 2.5 Resequencing and comparison of LEAFY gene sequence and DNA marker development

200 To compare sequences, resequencing of genomic DNA was performed for accessions of 201 'Kirakiraboshi,' 'Frau Yoshimi,' 'Posy Bouquet Grace,' and 'Blue Picotee Manaslu.' Sequencing 202 libraries (insert size of 500 bp) for the four lines were constructed with TruSeq DNA PCR-Free Library 203 Prep Kit (Illumina) to sequence on a HiSeqX (Illumina). From the sequence reads obtained, low-204 quality bases were deleted with PRINSEQ v.0.20.4²⁶ and adaptor sequences were trimmed with fastx 205 clipper (parameter, AGATCGGAAGAGC) in FASTX-Toolkit v.0.0.13 -a 206(http://hannonlab.cshl.edu/fastx toolkit). High-quality reads were aligned on the HMA r1.2 with 207 Bowtie 2^{27} v.2.2.3 to detect sequence variant candidates by with the mpileup command in SAMtools 208 v.0.1.19²⁴. High-confidence variants were selected using VCFtools²⁹ v.0.1.12b with parameters of --209 minDP 10, --maxDP 100, --minO 999, --max-missing 1.

For comparison of *LEAFY (LFY)* sequence in 'Kirakiraboshi,' 'Frau Yoshimi,' 'Posy Bouquet Grace,' and 'Blue Picotee Manaslu,' BLAST analysis using genomic sequence of *LFY* (Scaffold 0577F, position 678200-684639) as query, and genomic DNA sequence of each cultivar as database, was performed to confirm detected sequence variants. These data analyses were performed using CLC main workbench (Qiagen). INDEL marker S01 that amplifies the second intron of *LFY*, was designed by visual inspection (Forward: 5'-CATCATTAATAGTGGTGACAG-3', Reverse: 5'-CACACATGAATTAGTAGCTC-3'). The PCR conditions were 94°C for 2 min, 35 cycles of

denaturation at 94°C for 1 min, annealing at 55°C for 1 min, extension at 72°C for 1 min; and a final
extension step at 72°C for 3 min. The PCR product was stained with 1x GRRED (Biocraft) and
separated in 2.5% (w/v) agarose gel in TAE buffer.

220

221 2.6 Cloning and sequence determination of LFY gene of 'Kirakiraboshi' and 'Frau Yoshimi'

222 Total RNA was isolated from the flower buds of 'Kirakiraboshi,' and 'Frau Yoshimi' using RNAiso

223 Plus (TaKaRa, Japan), and reverse transcribed using PrimeScript II 1st strand cDNA Synthesis Kit

224 (TaKaRa, Japan). The sequence of the LFY gene was amplified by PCR in 50-μL reaction mixture by

225 using TaKaRa Ex Taq Hot Start Version (TaKaRa Bio, Shiga, Japan) and the LFY specific primer

226 (Forward: 5'-ATGGCTCCACTACCTCCACC-3' and Reverse: 5'-CTAACACCCTCTAAAAGCAG-

227 3'). These PCR products were purified, and inserted into a pMD20-T vector using the Mighty TA-

228 cloning kit (TaKaRa Bio). The sequence of *LFY* coding sequence (CDS) in pMD20-T vector was

analyzed by 3130xl DNA sequencer (Applied Biosystems, Foster City, CA, USA). Sequence
alignments were obtained by using CLC main workbench (Qiagen).

231

232 2.7 DNA marker assessment across hydrangea accessions

233 For assessment of DNA markers for the double flower phenotype, 35 H. macrophylla 234 accessions were used. Genotyping for J01 was performed as described above. Genotyping for S01 was 235 performed by fragment analysis as follows. PCR amplification was performed in a 10-µL reaction 236mixture containing 5 µL of GoTaq Master Mix (Promega), 5 pmol FAM-labeled universal primer (5' 237 - FAM-gctacggactgacctcggac -3'), 2.5 pmol forward primer with universal adapter sequence (5' -238 gctacggactgacctcggacCATCATTAATAGTGGTGACAG -3'), 5 pmol reverse primer, and 5 ng of 239 template DNA. DNA was amplified in 35 cycles of 94°C for 1 min, 55°C for 1 min, and 72°C for 2 240 min; and a final extension of 5 min at 72°C. The amplified PCR products were separated and detected

in a PRISM 3130xl DNA sequencer (Applied Biosystems, USA). The sizes of the amplified bands
were scored against internal-standard DNA (400HD-ROX, Applied Biosystems, USA) by
GeneMapper software (Applied Biosystems, USA).

244

245 **3. Results and Discussion**

246 **3.1 Draft genome assembly with long-read and HiC technologies**

The size of the hydrangea genome was estimated by k-mer-distribution analysis with the short-read of 132.3 Gb data. The resultant distribution pattern indicated two peaks, representing homozygous (left peak) and heterozygous (right peak) genomes, respectively (Figure 2). The haploid genome of hydrangea was estimated to be 2.2 Gb in size. The short reads were assembled into 612,846 scaffold sequences. The total length of the resultant scaffolds, i.e. HMA_r0.1, was 1.7 Gb with an N50 length of 9.1 kb (Supplementary Table S2). Only 72.2% of complete single copy orthologues in plant genomes were identified in a BUSCO analysis (Supplementary Table S2).

254Next, we employed long sequence technology to extend the sequence contiguity and to 255 improve the genome coverage. A total of 106.9 Gb of reads (49.4×) with an N50 read length of 28.8 256 kb was obtained from 14 SMRT Cells. The long-reads were assembled, followed by sequence error 257 corrections into 15,791 contigs consisting of 3,779 primary contigs (2.178 Gb in length and N50 of 258 1.4 Mb), and 12,012 haplotig sequences (1.436 Gb in length and N50 of 184 kb). To obtain two 259 haplotype-phased complete-length sequences, 697 M reads of HiC data (105.3 Gb) were obtained and 260 subjected to FALCON-Phase. The resultant haplotype-phased sequences consisted of 3,779 sequences 261 (2.256 Gb in length and N50 of 1.5 Mb) for "phase 0," and 3,779 sequences (2.227 Gb in length, and 262N50 of 1.4 Mb) for "phase 1."

264 **3.2 Pseudomolecule sequences based on genetic mapping**

265 To detect potential errors in the assembly and to assign the contig sequences onto the hydrangea 266 chromosomes, we established an F2 genetic map based on SNPs derived from a ddRAD-Seq 267 technology. Approximately 1.8 million high-quality ddRAD-Seq reads per sample were obtained from 268 the mapping population and mapped to either of the two phased sequences with alignment rates of 269 88.4% and 88.7%, respectively. A set of SNPs detected from the alignments were classified into 18 270 groups and ordered to construct two genetic maps for the two phased sequences (2,849.3 cM in length 271 with 3,980 SNPs, and 2,944.5 cM in length with 4,071 SNPs). The nomenclature of the linkage groups 272 was named in accordance with the previous genetic map based on SSRs⁵. The phased sequences were 273 aligned on each genetic map to establish haplotype-phased, chromosome-level pseudomolecule 274sequences. During this process, one contig was cut due to possible mis-assembly. The resultant 275sequences for phase 0 had 730 contigs with a total length of 1,078 Mb and the other for phase 1 had 276 743 contigs spanning 1,076 Mb.

277

278 **3.3. Transcriptome analysis followed by gene prediction**

279 In the Iso-Seq analysis, Circular Consensus Sequence (CCS) reads were generated from the raw 280 sequence reads. The CCS reads were classified in full-length and non-full length reads and the full-281 length reads were clustered to produce consensus isoforms. In total, 116,634 high-quality isoforms 282 were used for gene prediction. In the RNA-Seq analysis, on the contrary, a total of 80.7 Gb reads were 283 obtained and assembled into 12,265 unigenes. The high-quality isoforms and unigenes together with 284 gene sequences predicted from the Arabidopsis thaliana, Arachis hypogaea, Cannabis sativa, 285Capsicum annuum, Cucumis sativus, Populus trichocarpa, and Quercus lobate genomes were aligned 286 onto the assembly sequence of the hydrangea genome. By adding ab-initio on genes, 32,205 and 287 32,222 putative protein-encoding genes were predicted from the phase 0 and phase 1 sequences,

respectively. This gene set included 91.4% complete BUSCOs. Out of the 10,108 genes, 16,725, and 21,985 were assigned to Gene Ontology slim terms in the biological process, cellular component, and molecular function categories, respectively. Furthermore, 4,271 genes had assigned enzyme commission numbers.

292

293 **3.4 Identification of SNPs tightly linked to double flower phenotype**

294 To identify SNPs tightly linked to the double flower phenotype of 'Jogasaki,' ddRAD-Seq 295 analysis was performed on the 12GM1 population, which segregates the double flower phenotype of 296 'Jogasaki.' As a result, 14,006 of SNPs were called by ddRAD-Seq analysis of the 12GM1 population. 297 In this population, the double flower phenotype was expected when the plant was homozygous for the 298 'Posy Bouquet Grace' genotype, and the single flower phenotype was expected when the plant was 299 homozygous for 'Blue Picotee Manaslu' or was heterozygous. Each SNP was tested for its fitting rate 300 to this model. As a result, ten SNPs were found to have more than a 95% fitting rate, and six SNPs 301 were completely co-segregated with flower phenotype (Table 1).

302 CAPS marker J01 was developed based on SNP at scaffold 0008F-2_780104. J01 CAPS 303 marker amplified 167 bp of fragment by PCR, and digestion with Taq I restriction enzyme generated 304 50 bp and 117 bp fragments in the double flower allele (Figure 3). J01 marker was fitted with flower 305 phenotype at 99.3% in the 15IJP1 and 14GT77 populations, which segregated the double flower 306 phenotype of 'Jogasaki' (Supplementary Table S3, S4). This indicated that J01 marker was tightly 307 linked to the D_{jo} locus. Thus, D_{jo} is suggested to be located adjacent to J01, which is located at position 308 46,326,384 in CHR17, (Figure 4).

309 For identification of SNPs linked to the double flower phenotype of 'Sumidanohanabi,' the 310 KF population that segregates the double flower phenotype derived from 'Sumidanohanabi' were used. 311 First, we tried to find co-segregated scaffolds with the double flower phenotype by ddRAD-Seq

312 analysis of the KF population. As a result of ddRAD-Seq analysis, 15,102 of SNPs were called. In this 313 population, the double flower phenotype was expected when the plant was homozygous for the 314 'Kirakiraboshi' genotype, and the single flower phenotype was expected when the plant was 315 homozygous for 'Frau Yoshimi' or was heterozygous. Each SNP was tested for its fitting rate to this 316 model. As a result, five SNPs on three scaffolds were found to have more than a 95% fitting rate with 317 the model (Table 2). Since SNPs on scaffold 3145F all had the same genotype across the KF population, 318 three loci—on scaffold 0577F, 3145F, 0109F—were detected. According to genotypes of the KF 319 population, these three loci were tightly linked within 5 cM; 0109F (0 cM) - 3145F (3.9 cM) - 0577F 320 (5.0 cM). Since the SNP at position 868569 in 0109F was found at the position 57,436,162 in CHR04, 321 locus D_{su} , which controls the double flower phenotype of 'Sumidanohanabi,' was suggested to be 322 located on terminal of CHR04 (Figure 4).

323

324 **3.5 Prediction of genes controlling double flower**

325 To find the gene controlling D_{su} and D_{jo} , we searched the homeotic genes on scaffolds shown 326 in Table 1 and Table 2. We did not find any notable homeotic gene controlling flower phenotype for 327 D_{jo} . For D_{su} , the g182220 gene, which encoded a homeotic gene LFY, was found on scaffold 0577F. 328 To investigate the possibility that it was the causative gene for D_{su} , sequence variants on LFY genomic 329 sequence were searched to identify 'Kirakiraboshi' specific mutation, using resequencing data of 330 'Kirakiraboshi,' 'Frau Yoshimi,' 'Posy Bouquet Grace,' and 'Blue Picotee Manaslu.' As a result, five 331 INDELs and six sequence variants were found as 'Kirakiraboshi' specific mutations (Figure 5). 332 Cloning and sequencing of LFY CDS was performed on 'Kirakiraboshi' and 'Frau Yoshimi.' 333 From 'Frau Yoshimi,' a single CDS comprising three exons was obtained. From 'Kirakiraboshi,' two

334 CDSs with splice variants were obtained. While splicing 1 CDS resulted in three exons, splicing 2

335 CDS resulted in only two exons, corresponding to the first and third splice products of splicing 1 CDS

336 (Supplementary Figure S1). The deduced amino acid sequences were aligned using CDSs of 'Frau 337 Yoshimi' and 'Kirakiraboshi,' g182220 sequence, protein LFY of Arabidopsis thaliana, and protein FLO of Antirrhium majos. While the deduced amino acid sequences of 'Frau Yoshimi' and g182220 338 339 showed sequence similarity in the entire region, frameshift occurred in the two CDSs obtained from 340 'Kirakiraboshi' and the resulting products had no sequence similarity across the latter half (Figure 6). 341 Frameshift observed in splicing 1 CDS was due to one bp of DNA insertion in the second exon, at 342 position 1,931 (Figure N3A). On the contrary, frameshift observed in splicing 2 CDS was due to the 343 complete loss of the second exon (Figure 6). 344 To develop a DNA marker for distinguishing the d_{su} allele from the D_{su} alleles in the LFY 345 genomic sequence, we focused and designed a DNA marker on 'Kirakiraboshi' specific 14 bp deletion 346 at position 3,617 from initiation codon (Figure 5). We developed INDEL S01 marker amplified 236

347 bp fragment for the double flower allele of 'Kirakiraboshi,' and 250 bp and 280 bp fragments for the 348 single flower allele of 'Frau Yoshimi' (Figure 7A). Three types of alleles resulted from the presence 349 or absence of a 30 bp deletion at position 3,482 in addition to the 14 bp INDEL. These were both 30 350 bp and 14 bp deletions on the 236 bp allele, 30 bp deletion on the 250 bp allele, and no deletion on the

351 280 bp allele (Figure 7B).

352

353 3.6 Genotyping of hydrangea accessions using J01 and S01 markers

354 Since the J01 marker could distinguish D_{jo}/d_{jo} alleles and the S01 marker could distinguish 355 D_{su}/d_{su} alleles, a combined use of J01 and S01 DNA markers was expected to reveal the origin of the 356 double flower phenotype, d_{io} or d_{su} , in various accessions. Therefore, DNA marker genotyping on H. 357 macrophylla accessions were performed using two DNA markers, J01 and S01. All tested double 358 flower accessions showed homozygous genotypes of J01 or S01; ten of the double flower accessions 359 were homozygous of 117 50 in J01, and four were homozygous of 236 in S01 (Table 3). Contrarily, 360 all single flower accessions showed other genotypes.

361 Previously, the double flower phenotype has been revealed to be controlled by a single locus 362 with the inheritance of single flower dominant and double flower recessive genes^{4,5}. It was also 363 suggested that genes controlling the double flower phenotype were different between 'Jogasaki' and 364 'Sumidanohanabi' based on confirmation of the segregation ratio of crossed progenies⁴. Our study 365 revealed that the double flower phenotype of 'Jogasaki' was controlled by a single D_{io} locus on CHR17, 366 and the double flower phenotype of 'Sumidanohanabi' was controlled by a single D_{su} locus on CHR04. 367 In addition, all double flower accessions showed homozygosity for the double flower allele at one 368 locus, D_{jo} or D_{su} . Contrarily, all single flowers have dominant single flower alleles on both D_{jo} and D_{su} 369 loci. This indicated that each locus independently controls flower phenotype.

370 Developed DNA markers J01 and S01 could successfully identify recessive double flower 371 alleles for D_{jo} and D_{su} , respectively. Both markers showed high fitting ratio with phenotype and were 372 applicable to the examined H. macrophylla accessions. The S01 marker is superior to the DNA marker 373 STAB045 linked to D_{su} and which was discovered by Waki et al.⁵ because the former has a wide range 374 of applicability. While the S01 marker genotype completely fitted with the phenotype in all tested 375 accessions, STAB045 did not (data not shown). Because both J01 and S01 showed a wide range of 376 applicability, it is advantageous to use them in combination to reveal the existence of the double flower 377 allele in H. macrophylla accessions. This information will help in selection of candidate parents with 378 heterozygous recessive double flower alleles to obtain double flower progenies. In addition, these 379 DNA markers should be useful in marker assisted selection (MAS) of double flower progenies. To 380 obtain double flower progenies, at least the paternal parent should be of the single flower phenotype 381 because very few or none at all pollen grains are produced in double flower individuals. In addition, 382 it requires approximately 2 years to confirm the flower phenotype from the time of crossing. 383 Identification of flower phenotype at the seedling stage by MAS would enable the discarding of single

flower individuals and allow the growth of double flower individuals. The developed DNA markers should accelerate the breeding of double flower phenotypes.

386 In the genomic sequence of 'Kirakiraboshi,' an insertion was detected in the second exon of 387 the LFY gene. This insertion actually resulted in frameshift of cloned mRNA in both splice variants. 388 Therefore, it was speculated that the function of the LFY gene was suppressed or lost in 'Kirakiraboshi'. 389 The LFY gene and its homologue FLO have been identified in many plants, such as Arabidopsis 390 thaliana and Antirrhinum majus, and are known as transcription factors for major flowering signals²⁹⁻ 391 ³¹. Additionally, many types of phenotypes in *Arabidopsis lfy* mutants have been reported^{32,33}. In the 392 *lfy* strong phenotype, most organs are sepal-like, or mosaic sepal/carpels organs, and the sepal-like 393 organs are characteristic of wild-type cauline leaves³³. Therefore, the flowers of the *lfy* mutant 394 appeared to be double flowers that are formed from leaves or sepals. Additionally, a similar phenotype 395 has been reported in LFY homologue mutants or transgenic plants such as the flo mutant of 396 Antirrhinum majus³⁴, uni mutant of pea³⁵, and co-suppressed NFL transgenic plant of tobacco³⁷. 397 Therefore, generally, when the LFY gene function is lost, petal, stamens, and a carpel are likely to be 398 replaced by sepal-like organs. In decorative flowers of hydrangea, sepals show petaloid characteristics 399 including pigmentation and enlarged organ size. It is possible that sepal-like organs in decorative 400 flowers show petaloid characteristics and form double flowers. Therefore, we assumed that LFY is a 401 causative gene of the double flower phenotype of 'Sumidanohanabi'.

402 However, there remain several unexplained observations in this study. The double flower of 403 'Kirakiraboshi' did not exhibit the exact same phenotype of the *lfy* mutant. Generally, the flowers of 404 *lfy* or its orthologous gene mutants have only leaf-like or sepal-like organs that have chlorophyll, 405 stomata, and trichome, and these organs have almost no petal identity^{33,34}. When flowering signals in 406 *lfy* mutant were lost completely, floral organs were not fully formed³³⁻³⁵. In the double flowers of 407 'Kirakiraboshi', the floral organs keep their petal identity, have papilla cells, and are pink or blue.

408 These phenotypes of 'Kirakiraboshi' might reflect partial remaining of LFY function. Additionally, it 409 has been reported that *lfy* mutants with an intermediate or weak phenotype sometimes develop petaloid 410 organs³³. According to the genomic sequence of *H. macrophylla*, no other *LFY* gene was observed. It

411 could be considered that the double flowers of 'Kirakiraboshi' were induced via partial repression of412 the LFY function.

413 On the contrary, we could not find any candidate gene that controls the double flower 414phenotype for the D_{jo} locus. One possible reason was that SNPs were not called in scaffold with 415 causative gene. In pseudomolecules, about half of the total scaffolds length was not included since 416 relevant SNPs were not called. Improvement of SNP density would be effective for discovering 417 additional scaffolds that are tightly linked to D_{io} . Although candidate gene for D_{io} could not be 418 identified from the linkage information, we predicted several candidate genes. In hydrangea, stamens 419 and petals were absent from decorative flowers of the double flower plant, and there was an increased 420 number of sepals⁴. Since causative genes should explain the changes in formation, the B-class genes 421 of the ABC model, PI and AP3, were predicted as candidate genes. In A. thaliana, the B-class gene pi 422 or ap3 mutants showed an increase in the number of sepals converted from petals³⁷. If these genes 423 were mutated in hydrangea, an increase in sepals would be expected. In hydrangea, HmPI, HmAP3, 424 and HmTM6 were identified as B-class genes^{38,39}. As HmAP3 was located on CHR13, it was not 425 considered as a causative gene for D_{io} . In this study, HmPI and HmTM6 were not included in the 426 pseudomolecule. Ascertaining the loci of these genes might reveal the causative gene for D_{io} .

In this study, we report DNA markers and possible causative genes for the double flower phenotype observed in two hydrangea cultivars. For this analysis, we established a reference sequence for the hydrangea genome using advanced sequencing technologies including the long-read technology (PacBio) and the HiC method⁹, bioinformatics techniques for the diploid genome assembly¹⁴, and haplotype phasing⁸. To the best of our knowledge, this is the first report on the

432	chromosome-level haplotype-phased sequences in hydrangea at the level of the species (H.
433	macrophylla), genus (Hydrangea), family (Hydrangeaceae), and order (Cornales). The genomic
434	information from this study based on NGS technology is a significant contribution to the genetics and
435	breeding of hydrangea and its relatives. It will serve to accelerate the knowledge base of the evolution
436	of floral characteristics in Hydrangeaceae.
437	
438	Acknowledgments: We thank Ohama A, Ono M, Seki A and Kitagawa A (Nihon University) and
439	Sasamoto S, Watanabe A, Nakayama S, Fujishiro T, Kishida Y, Kohara M, Tsuruoka H, Minami C,
440	and Yamada M (Kazusa DNA Research Institute) for their technical help.
441	
442	Funding: This study was partially supported by the Nihon University College of Bioresource Sciences
443	Research Grant for 2018, and by the JSPS KAKENHI Grant, Number JP18K14461.
444	
445	Supporting information:
446	Supplementary Table S1. RNA samples used for Iso-Seq and RNA-Seq
447	Supplementary Table S2. Statistics of the genome sequences of Hydrangea macrophylla
448	'Aogashima-1'
449	Supplementary Table S3. J01 marker genotypes and double flower phenotypes of 15IJP1 population.
450	Supplementary Table S4. J01 marker genotypes and double flower phenotypes of 14GT77
451	population.
452	Supplementary Figure S1. Alignment of LFY genomic sequence and CDS.
453	
454	Data availability:
455	The sequence reads are available from the DNA Data Bank of Japan (DDBJ) Sequence Read Archive

- 456 (DRA) under the accession numbers DRA010300, DRA010301, and DRA010302. The assembled
- 457 sequences are available from the BioProject accession number PRJDB10054. The genome information
- 458 is available at Plant GARDEN (https://plantgarden.jp).
- 459
- 460 **References**
- 461 1. Uemachi, T., Kato, Y., and Nishio, T. 2004, Comparison of decorative and non-decorative flowers
- 462 in Hydrangea macrophylla (Thunb.) Ser., Sci. Hortic., 102, 325–334
- 463
- 464 2. Uemachi, T., Kurokawa, M., and Nishio, T. 2006, Comparison of inflorescence composition and
- 465 development in the lacecap and its sport, hortensia Hydrangea macrophylla (Thunb.) Ser., J. Japan.
- 466 Soc. Hort. Sci., 75, 154–160.
- 467
- 468 3. Uemachi, T. and Okumura, A. 2012, The inheritance of inflorescence types in *Hydrangea*469 *macrophylla, J. Japan. Soc. Hort. Sci.*, 81, 263–268.
- 470
- 4. Suyama, T., Tanigawa, T., Yamada, A. et al. 2015, Inheritance of the double-flowered trait in
 decorative hydrangea flowers, *Hortic. J.*, 84, 253-260.
- 473
- 474 5. Waki, T., Kodama, M., Akutsu, M. et al. 2018, Development of DNA markers linked to double-
- 475 flower and hortensia traits in *Hydrangea macrophylla* (Thunb.) Ser., *Hortic J.*, 87, 264-273.
- 476
- 6. Heijmans, K., Ament, K., Rijpkema, A.S. et al. 2012, Redefining C and D in the petunia ABC, *Plant Cell*, 24, 2305-2317.
- 479

- 480 7. Tränkner, C., Krüger, J., Wanke, S., Naumann, J., Wenke, T. and Engel, F. 2019, Rapid identification
- 481 of inflorescence type markers by genotyping-by-sequencing of diploid and triploid F1 plants of
- 482 *Hydrangea macrophylla*, *BMC Genet.*, 20, 60.
- 483
- 484 8. Kronenberg, Z. N., Hall, R. J., Hiendleder, S., et al. 2018, FALCON-Phase: Integrating PacBio
- 485 and Hi-C data for phased diploid genomes, *BioRxiv*, 327064.
- 486
- 487 9. Dudchenko, O., Batra, S. S., Omer, A. D., et al. 2017, De novo assembly of the Aedes aegypti

488 genome using Hi-C yields chromosome-length scaffolds. *Science*, 356, 92-95.

- 489
- 490 10. Mascher, M. and Stein, N. 2014, Genetic anchoring of whole-genome shotgun assemblies,
 491 *Front Genet*, 5, 208.
- 492
- 493 11. Marcais, G. and Kingsford, C. 2011, A fast, lock-free approach for efficient parallel counting
 494 of occurrences of k-mers, *Bioinformatics*, 27, 764-770.
- 495
- 12. Kajitani, R., Toshimoto, K., Noguchi, H., et al. 2014, Efficient de novo assembly of highly
- 497 heterozygous genomes from whole-genome shotgun short reads, *Genome Res*, 24, 1384-1395.
- 498
- 499 13. Simao, F. A., Waterhouse, R. M., Ioannidis, P., Kriventseva, E. V., and Zdobnov, E. M. 2015,
- 500 BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs,
- 501 *Bioinformatics*, 31, 3210-3212.
- 502
- 503 14. Chin, C. S., Peluso, P., Sedlazeck, F. J., et al. 2016, Phased diploid genome assembly with

single-molecule real-time sequencing, *Nat Methods*, 13, 1050-1054.

- 505
- 506 15. Walker, B. J., Abeel, T., Shea, T., et al. 2014, Pilon: an integrated tool for comprehensive
- 507 microbial variant detection and genome assembly improvement, *PLoS One*, 9, e112963.
- 508
- 509 16. Shirasawa, K., Hirakawa, H., and Isobe, S. 2016, Analytical workflow of double-digest
- 510 restriction site-associated DNA sequencing based on empirical and in silico optimization in
- 511 tomato, DNA Res, 23, 145-153.
- 512
- 513 17. Rastas, P. 2017, Lep-MAP3: robust linkage mapping even for low-coverage whole genome 514 sequencing data, *Bioinformatics*, 33, 3726-3732.
- 515
- 516 18. Tang, H., Zhang, X., Miao, C., et al. 2015, ALLMAPS: robust scaffold ordering based on
 517 multiple maps, *Genome Biol*, 16, 3.
- 518
- 519 19. Grabherr, M. G., Haas, B. J., Yassour, M., et al. 2011, Full-length transcriptome assembly

520 from RNA-Seq data without a reference genome, *Nat Biotechnol*, 29, 644-652.

- 521
- 522 20. Stanke, M., Keller, O., Gunduz, I., Hayes, A., Waack, S., and Morgenstern, B. 2006,
- 523 AUGUSTUS: ab initio prediction of alternative transcripts, *Nucleic Acids Res*, 34, W435-439.
- 524
- 525 21. Kent, W. J., 2002, BLAT the BLAST-like alignment tool, Genome Res, 12, 656-664.
- 526
- 527 22. Ghelfi, A., Shirasawa, K., Hirakawa, H., and Isobe, S. 2019, Hayai-Annotation Plants: an

528	ultra-fast a	ind c	comprehensive	functional	gene	annotation	system	in j	plants,	Bioinformatics,	35,

529 4427-4429.

530

531 23. Bolger, A.M., Lohse, M., and Usadel, B. 2014, Trimmomatic: a flexible trimmer for Illumina
532 sequence data, *Bioinformatics*, 30, 2114-2120.

533

- 534 24. Li, H., Handsaker, B., Wysoker, A., et al. 2009, The Sequence Alignment/Map format and
- 535 SAMtools, *Bioinformatics*, 25, 2078-2079.

536

- 537 25. Untergasser, A., Cutcutache, I., Koressaar, T. et al. 2012, Primer3--new capabilities and interfaces.
- 538 Nucleic Acids Res., 40, e115.

539

540 26. Schmieder, R. and Edwards, R. 2011, Quality control and preprocessing of metagenomic
541 datasets, *Bioinformatics*, 27, 863-864.

542

543 27. Langmead, B. and Salzberg, S. L. 2012, Fast gapped-read alignment with Bowtie 2, *Nat*544 *Methods*, 9, 357-359.

545

546 28. Danecek, P., Auton, A., Abecasis, G., et al. 2011, The variant call format and VCFtools,

547 *Bioinformatics*, 27, 2156-2158.

- 549 29. Jaeger, K.E., Pullen, N., Lamzin, S., Morris, R.J., and Wigge, P.A. 2013, Interlocking feedback
- 550 loops govern the dynamic behavior of the floral transition in Arabidopsis, *Plant Cell*, 25, 820-
- 551 833.

- 553 30. Krizek, B.A. and Fletcher, J.C. 2005, Molecular mechanisms of flower development: an
- armchair guide, *Nat. Rev. Genet.*, 6, 688.
- 555
- 556 31. William, D.A., Su, Y., Smith, M.R., Lu, M., Baldwin, D.A., and Wagner, D. 2004, Genomic
- 557 identification of direct target genes of LEAFY, Proc. Nat. Acad. Sci., 101, 1775–1780.
- 558
- 559 32. Okamuro, J.K., Den Boer, B.G., and Jofuku, K.D. 1993, Regulation of Arabidopsis flower
- 560 development, *Plant Cell*, 5, 1183-1193.
- 561
- 33. Weigel, D., Alvarez, J., Smyth, D.R., Yanofsky, M.F., and Meyerowitz, E.M. 1992, LEAFY
 controls floral meristem identity in Arabidopsis, *Cell*, 69, 843-859.
- 564
- 565 34. Carpenter, R. and Coen, E.S. 1990, Floral homeotic mutations produced by transposon-566 mutagenesis in *Antirrhinum majus*, *Gene. Dev.*, 4, 1483–1493.
- 567
- 35. Hofer, J., Turner, L., Hellens, R. et al. 1997, UNIFOLIATA regulates leaf and flower
 morphogenesis in pea, *Curr. Biol.*, 7, 581–587.
- 570
- 571 36. Ahearn, K.P., Johnson, H.A., Weigel, D., and Wagner, D.R. 2001, NFL1, a Nicotiana tabacum
- 572 LEAFY-like gene, controls meristem initiation and floral structure, *Plant Cell Physiol.*, 42, 1130–
 573 1139.
- 574
- 575 37. Bowman, J.L., Smyth, D.R., and Meyerowitz, E.M. 1989, Genes directing flower development in

576 Arabidopsis, *Plant Cell*, 1, 37-52.

578	38. Kitamura,	Y., Hosokawa,	M., U	Jemachi, T.,	and Yazawa,	S. 2009,	Selection	of ABC g	genes for
-----	---------------	---------------	-------	--------------	-------------	----------	-----------	----------	-----------

579 candidate genes of morphological changes in hydrangea floral organs induced by phytoplasma

580 infection, Sci. Hort., 122, 603-609.

582 39. Kramer, E.M. and Irish, V.F. 2000. Evolution of the petal and stamen development programs:

583 Evidence from comparative studies of the lower eudicots and basal angiosperms, Int. J. Plant Sci., 161,

- 584 s29-s40

	D iii i	Sequenc	e variant	TT-1	Frequency of double flower phenotype (double flower/all)					
Scaffold	Position at Phase 0	Posy Bouquet Grace	. (70)		Homozygous of 'Posy Bouquet Grace'	Heterozygous	Homozygous of 'Blue Picotee Manasulu			
0008F-2	3250598	А	G	100	37/37	0/61	0/47			
0008F-2	3250523	А	С	100	37/37	0/61	0/47			
0008F-2	780104	С	А	100	37/37	0/60	0/48			
0259F	404610	Т	А	100	37/37	0/60	0/48			
1207F	365533	С	Т	100	38/38	0/61	0/48			
1207F	372121	С	А	100	38/38	0/61	0/47			
0012F	1318350	Т	С	97.9	37/39	1/59	0/48			
0437F	170787	G	А	97.9	36/37	1/60	1/49			
0437F	180821	А	G	97.9	36/37	1/60	1/49			
0994F	216439	С	Т	97.9	36/37	1/60	1/49			

Table 1. SNPs correlated (fitting rate more than 95%) with double flower phenotype in 12GM1 population

	Position at	Sequence variant		Fitting rate	Frequency of double flower phenotype (double flower/all)					
Scaffold	Phase 0	Kirakiraboshi	Frau Yoshimi	(%)	Homozygous of 'Kirakiraboshi'	Heterozygous	Homozygous of 'Frau Yoshimi'			
0577F	1204837	AG	AAACATG	98.9	22/22	0/51	1/20			
3145F	55089	TA	TAA	98.9	22/22	0/51	1/20			
3145F	55109	G	А	98.9	22/22	0/51	1/20			
3145F	55446	G	А	98.9	22/22	0/51	1/20			
0109F	868569	С	G	95.7	22/25	0/44	1/24			

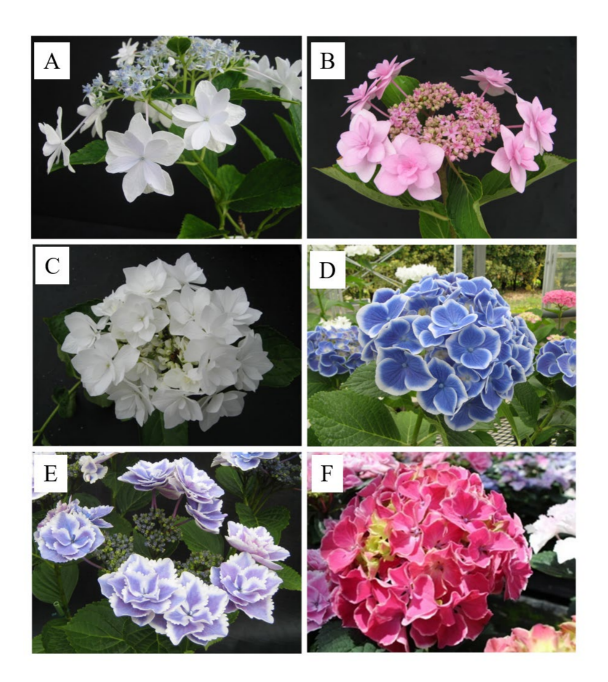
Table 2. SNPs correlated (fitting rate more than 95%) with double flower phenotype in KF population

-

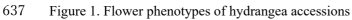
A	Dh an a fam a	Genotype				
Accession name	Phenotype	J01	S01			
Jogasaki	Double	117_50/117_50	250/280			
Posy Bouquet Grace	Double	117_50/117_50	280/280			
Izunohana	Double	117_50/117_50	250/280			
Chikushinokaze	Double	117_50/117_50	250/280			
Chikushinomai	Double	117_50/117_50	280/280			
Chikushiruby	Double	117_50/117_50	280/280			
Corsage	Double	117_50/117_50	280/280			
Dance Party	Double	117_50/117_50	280/280			
Fairy Eye	Double	117_50/117_50	250/280			
Posy Bouquet Casey	Double	117_50/117_50	250/280			
Sumidanohanabi	Double	167/167	236/236			
Kirakiraboshi	Double	167/167	236/236			
HK01	Double	167/167	236/236			
HK02	Double	167/167	236/236			
03JP1	Single	117_50/167	280/280			
Amethyst	Single	167/167	250/280			
Blue Picotee Manaslu	Single	167/167	280/280			
Blue Sky	Single	167/167	280/280			
Bodensee	Single	167/167	250/250			
Chibori	Single	167/167	280/280			
Furau Mariko	Single	167/167	250/250			
Furau Yoshiko	Single	167/167	280/280			
Furau Yoshimi	Single	167/167	250/280			
Green Shadow	Single	167/167	280/280			
Kanuma Blue	Single	167/167	250/280			
Mrs. Kumiko	Single	167/167	280/280			
Paris	Single	167/167	280/280			
Peach Hime	Single	167/167	280/280			
Picotee	Single	167/167	282/282			
Ruby Red	Single	167/167	280/280			
Shinkai	Single	167/167	280/280			
Tokimeki	Single	167/167	280/282			
Uzuajisai	Single	167/167	250/280			

Table 3. Genotypes of DNA marker J01 and S01 in H. macrophylla varieties

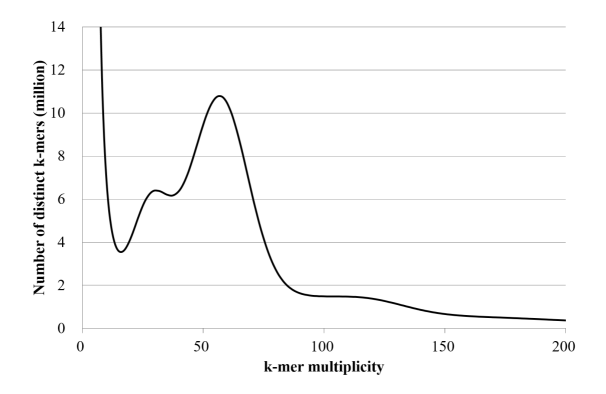
635 Genotypes shown as gray indicate homozygous of double flower allele.



636

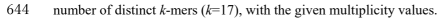


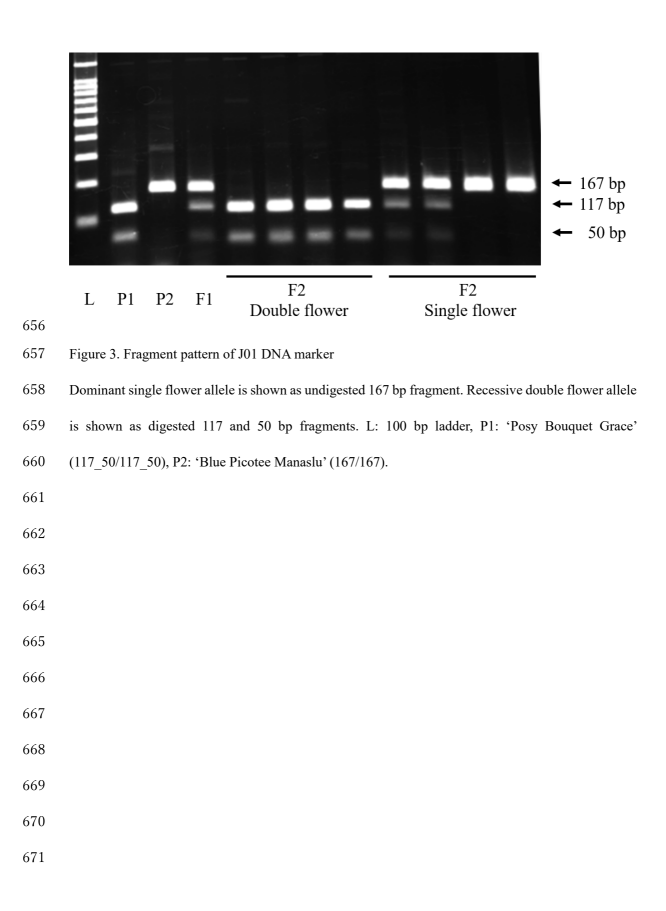
- 638 A: 'Sumidanohanabi' (double flower). B: 'Jogasaki' (double flower). C: 'Posy Bouquet Grace' (double
- 639 flower). D: 'Blue Picotee Manaslu' (single flower). E: 'Kirakiraboshi' (double flower). F: 'Frau
- 640 Yoshimi' (single flower).

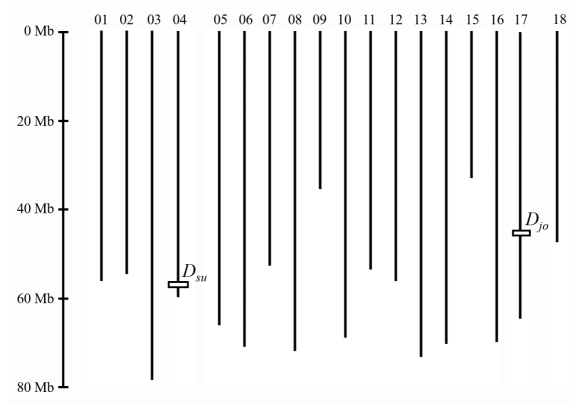




643 Figure 2. Genome size estimation for the hydrangea line 'Aogashima-1' with the distribution of the







672

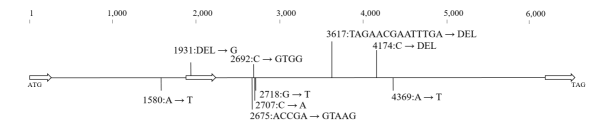
673 Figure 4. Schematic model of pseudomolecules

674 Double flower phenotype controlling loci D_{su} and D_{jo} are shown. D_{jo} is shown as J01 marker position

46,326,384 in CHR17. D_{su} is shown as tightly linked SNP at 0109F_868569, since the S01 marker

676 sequence was not on the pseudomolecule.

- 678
- 679
- 680
- 681
- 682
- 683
- 684
- 685



- 687 Figure 5. DNA polymorphisms in *LFY* genomic sequence
- 688 LFY sequence polymorphisms observed specifically in 'Kirakiraboshi' genomic sequence
- 689 The sequence is started from the initiation codon (ATG) at 678,200 to the termination signal (TAG) at
- 690 684,639 in phase 1 sequence of 0577F of HMA_r1.2. White arrows indicate coding sequences, CDS1:
- 691 1 to 454 bp, CDS2: 1,888 to 2,255 bp, CDS3: 6,078 to 6,440 bp. Genetic variants are shown as from
- 692 Hma1.2 sequence to 'Kirakiraboshi'.
- 693
- 694
- 695
- 696
- 697
- 698
- 699
- 700
- 701
- 702
- 703

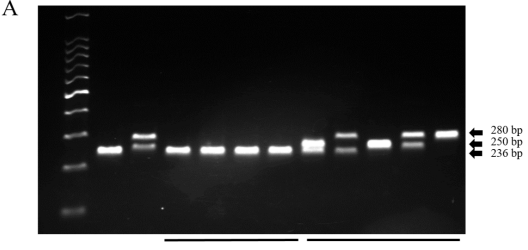
		20		40		60		80		100
Arabidopsis thaliana	MDPEGFTSGL	FRWNPTRALV	QAPP	PVPPPLQQQP	VTPQTAAFGM	R LGGLEG	LFGPYGIRFY	TAAKIAELGF	TASTLVGMKD	EELEEMMNSL 91
Antirrhinum majus	MDPDAF L	FKWDHRTALP	QPNRLLDAVA	PPPPPPQAP	S Y SM	RPRELGGLEE	LFQAYGIRYY	TAAKIAELGF	TVNTLLDMRD	EELDEMMNSL 91
HMA_r1.2p_g182220 Frau Yoshimi	MDSDAFSASL	FKWDPRGALP	PPNRLLDPMA	PLPPPP-PPP	SSTAAAAYPV		LFQAYGIRYY	TGAKIAELGF	TVNTLLNMKD	DELDDMMTSL 99
Kirakiraboshi splicing 1	MDSDAFSASL MDSDAFSASL	FKWDPRGALP	PPNRLLDPMA PPNRLLDPMA	PLPPPPLPPP	SSTAAAAYPV SSTAAAAYPV	RPRELGGLED RPRELGGLED	L FQAYGIRYY L FQAYGIRYY	TGAKIAELGF	TVNTLLNMKD	DELDDMMTSL 100 DELDDMMTSL 100
Kirakiraboshi_splicing 2	MDSDAFSASL	FKWDPRGALP	PPNRLLDPMA	PLPPPPLPPP	SSTAAAAYPV	RPRELGGLED	LFQAYGIRYY	TGAKIAELGF	TVNTLLNMKD	DELDDMMTSL 100
Conservation					Nonno 100nn					
•		120		140		160		180		200
Arabidopsis thaliana	SHIFRWELLV	GERYGIKAAV	RAERRRLQEE	EEEESSRRRH	LLLSAAGDSG	THHALDALSQ	EGLSEEPVQQ	QDQTDAAGNN	GGGGSG - YWD	AGQGKMKKQQ 190
Antirrhinum majus	CQIFRWDLLV	GERYGIKAAV	RAERRRIDEE	E VRRRH	LLLG D	TTHALDALSQ	EGLSEEPVQQ	EKEAMGS	GGGGVGGVWE	MMG-AGGRKA 178
HMA_r1.2p_g182220	SQIFRWDLLV	GERYGIKAAV	RAERRRLDEE	E ARRRH	LL SVD	TANALDALSQ	EGLSEEPVQQ	EKEAGGS	GGGG GTWE	VVVAAGSKKK 185
Frau Yoshimi	SQIFRWDLLV	GERYGIKAAV	RAERRRLDEE	E ARRRH		TANALDALSQ	EGLSEEPVQQ			VVVAAGSKKK 186
Kirakiraboshi_splicing 1 Kirakiraboshi_splicing 2	SQIFRWDLLV	GERYGIKAAV	RAERRRLDEE	E ARRRH	LLSVD	TANALDALSQ	EGLSEEPVQQ	EKEAGGEWWR	WRNVGGGGCG	
100	SQIFRWDLLV	GERYGIKAAV	RAERRRLDEE	E ARRRH	LLSVD	TANALDALSQ	EG			153
Conservation		220		240						annannanna
		T		T		260 I		280 I		300 I
Arabidopsis thaliana	QQRRRKK	- PMLTSVET -	DEDVNEG			QREHPFIVTE		GLDYLFHLYE	QCREFLLQVQ	TIAKDRGEKC 281
Antirrhinum majus	PQRRRKNYKG	RSRMASMEED	DDDDDDETEG	AEDDENIVS -		QREHPFIVTE	PGEVARGKKN	GLDYLFHLYE	QCRDFLIQVQ	TIAKERGEKC 269
HMA_r1.2p_g182220 Frau Yoshimi	- QRRRK	DODMACVE	DEDETDG				PGEVARGKKN			NIAKERGEKC 261
Kirakiraboshi splicing 1	- QRRRK TG DRAVEDGIGG		* * KWRRRWWR		GGER GAG*GGTWQK	ERP*LSIPSL	*AVP*FLDPS			NIAKERGEKC 272 QVCQEGRCEL 291
Kirakiraboshi splicing 2		GRORDGWER						SENC GERRE		QVCQEGRCEL 168
Conservation										
0		320		340		360		380		400
Arabidopsis thaliana		1				1		100000000		
Arabiaopsis inaliana Antirrhinum majus	PTKVTNQVFR	YAKKSGASYI YAKKAGANYI	NKPKMRHYVH	CYALHCLDEE	ASNALRRAFK	ERGENVGSWR		ACRHGWDIDA	VFNAHPRLSI	WYVPTKLRQL 381 WYVPTKLRQL 369
HMA r1.2p g182220	PTKVTNQVFR		NKPKMRHYVH	CYALHCLDED	SSNALRRIFK		QACYKPLVAI	AARQGWDIDA	IFNAHPRLAI	WYVPTKLRQL 361
Frau Yoshimi										WYVPTKLRQL 372
Kirakiraboshi_splicing 1	HKQAQNAPLR	PLLCPTLSRR	GLIKRTKADF	QGEGRERRGV	EASVLQAPRG	HSGPTGLGYR	CHFQRPSTTC	HLVRPHQTPS	ALSRRAE*RC	HGAGGAEHLC 391
Kirakiraboshi_splicing 2	HKQAQNAPLR	PLLCPTLSRR	GLIKRTKADF	QGEGRERRGV	EASVLQAPRG	HSGPTGLGYR	CHFQRPSTTC	HLVRPHQTPS	ALSRRAE*RC	HGAGGAEHLC 268
Conservation	nnnn nnn l									
		420								
Arabidopsis thaliana	CHLERNNAVA	AAAALVGGIS	CIGSSISGRG	GCGGDDLRE 4	20					
Antirrhinum majus	CHAERSSAAV	AATSSITGGG	PADHLPF		96					
HMA_r1.2p_g182220	CHAERSSA				69					
Frau Yoshimi	CHAERSSAAM	VPVGPSTSAF	RGC		95					
Kirakiraboshi_splicing 1 Kirakiraboshi_aplicing 2	F*RVL				96					
Kirakiraboshi_splicing 2	F*RVL			2	73					
Conservation	Mallanha.									

705 Figure 6. Alignment of LFY protein sequences

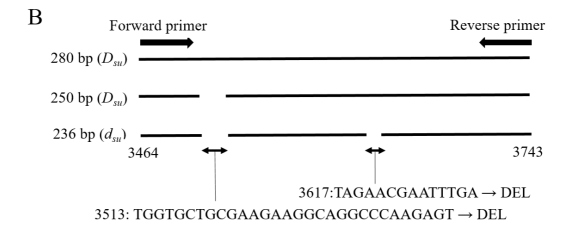
Amino acids with gray background show frameshifted regions. Splicing variant was observed, and

507 both sequences showed frameshift in 'Kirakiraboshi'. Arabidopsis thaliana: ABE66271.1 Antirrhium

majus: AAA62574.1.



L P1 P2 Double flower Single flower



- 719 Figure 7. Fragment pattern of S01 DNA marker
- A. Fragment pattern of S01 DNA marker. Dominant single flower alleles are shown as 250 bp and 280
- bp fragments. Recessive double flower allele is shown as 236 bp fragments. L: 100 bp ladder, P1:
- 722 'Kirakiraboshi' (236/236), P2: 'Frau Yoshimi' (250/280).
- 723 B. INDEL polymorphisms in alleles of DNA marker S01 amplified sequences. Position on schematic
- models were the same as in Figure 5.