1	Post-transcriptional silencing of <i>chalcone synthase</i> is involved in phenotypic lability
2	in petals and leaves of bicolor dahlia (Dahlia variabilis) 'Yuino'
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18	Main conclusion
19	Post-transcriptional gene silencing (PTGS) of a chalcone synthase (DvCHS2) occurred in
20	the white part of bicolor petals and flavonoid-poor leaves, however it did not in red petals
21	and flavonoid-rich leaves.
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### 26 Author contribution

SO and MH conceived the research. SO and MD designed the research. SO, WH and FT
conducted the research. SO wrote the manuscript. All authors read and approved the
manuscript.

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### 32 Abstract

33 Petal color lability is a prominent feature of bicolor dahlia cultivars, and causes plants to 34produce not only original bicolor petals with colored bases and pure white tips, but also frequently single-colored petals without white tips. In this study, we analysed the 3536 molecular mechanisms that are associated with petal color lability using the red-white bicolor cultivar 'Yuino'. Red single-colored petals lose their white tips as a result of 37recover of flavonoid biosynthesis. Among flavonoid biosynthetic genes including four 38chalcone synthase (CHS)-like genes (DvCHS1, DvCHS2, DvCHS3 and DvCHS4), 39 DvCHS1 and DvCHS2 had significantly lower expression levels in the white part of 40 41 bicolor petals than in red petals, while DvCHS3, DvCHS4 and other flavonoid 42biosynthetic genes had almost the same expression levels. Small RNAs from the white part of a bicolor petal were mapped onto DvCHS1 and DvCHS2, while small RNAs from 43a red single-coloured petal were not mapped onto any of the four CHS genes. A 44 relationship between petal color and leaf flavonoid accumulation has previously been 4546 demonstrated, whereby red petal-producing plants accumulate flavonoids in their leaves while bicolor petal-producing plants tend not to. The expression level of DvCHS2 was 47 down-regulated in flavonoid-poor leaves and small RNAs from flavonoid-poor leaves 48were mapped onto DvCHS2, suggesting that the down-regulation of DvCHS2 in 4950flavonoid-poor leaves occurs post-transcriptionally. Genomic analysis also suggested that

51	DvCHS2 is the key gene involved in bicolor formation. Together, these results suggest
52	that post-transcriptional gene silencing of <i>DvCHS2</i> plays a key role in phenotypic lability
53	in this bicolor dahlia.
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55	Key words: CHS, flavonoid, flower color, PTGS, siRNA.
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57	Abbreviation
58	CHS: chalcone synthase
59	HPLC: high-performance liquid chromatography
60	PTGS: post-transcriptional gene silencing
61	SAM: shoot apical meristem
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#### 76 Introduction

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Dahlia (Dahlia variabilis, Asteraceae) is one of popular floriculture crops that are used 7879 as cut flowers and garden plants. Dahlia flowers exhibit huge variations in their traits, 80 including color, shape and size. Flower color is particularly diverse, with purple, red, pink, orange, black, ivory white, yellow, variegated and bicolor cultivars currently available. 81 82 This huge phenotypic variation may be derived from the complicated genetic background 83 and large genome size of this species, as it is believed to be an autoallooctoploid (Gatt et 84 al. 1998) with a genome that is estimated at more than 8.8 Gb per haploid (Temsch et al. 2008). Due to this complicated genetic background and high heterozygosity, the 85 86 propagation of dahlias relies on vegetative propagation, such as through tuberous roots and cuttings (Konishi and Inaba 1964). 87

Bicolor dahlias belong to a group of cultivars that have petals with colored bases 88 and white tips. A prominent feature of these cultivars is the lability, or instability, of petal 89 color, as in addition to the original bicolor petals, they also frequently produce single-90 91 colored petals without white tips, despite being propagated vegetatively (Fig. 1). This 92petal color lability is observed not only among clonal plants but also within a single plant 93 and even within an inflorescence. Bicolor dahlias can bloom inflorescences with only 94 bicolor petals, only single-colored petals, or a mixture of the two. In a mixed inflorescence, 95 the single-colored petals are located in the outer whorls or sectorally in many cases (Ohno 96 et al. 2016), making it unlikely that this phenomenon can be explained by a mutation in particular genes. This petal color lability in bicolor dahlias attracted much research in the 97 1930s (Lawrence 1931; Tammes and Groeneveld-Huisman 1939) and yet the 98 mechanism(s) underlying it remain unknown. 99

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'Yuino' is a red-white bicolor dahlia cultivar that produces petals with red bases

101and white tips, but also often produces red single-colored petals (Ohno et al. 2016, Fig. 1021). Based on the observation that these red petals occur sectorally in an inflorescence 103 (Ohno et al. 2016), it was assumed that whatever phenomenon controls their production 104 occurs meristematically. There were two types of leaves, one was flavonoid-rich leaves 105and the other was flavonoid-poor leaves. A relationship between petal color and leaf 106 flavonoid accumulation has been demonstrated, whereby red petal-producing plants tend 107 to produce flavonoid-rich leaves while solely bicolor petal-producing plants tend to 108 produce flavonoid-poor leaves. This indicates that petal color lability can be considered 109 a phenotypic change at the whole-plant level (Ohno et al. 2016).

It has previously been shown that 'Yuino' accumulates anthocyanins, flavones 110 111 and butein in the red parts of the petals, whereas no flavonoids are accumulated in the 112white tips (Ohno et al. 2011b). The flavonoid biosynthetic genes in dahlia have been largely elucidated, with anthocyanin biosynthesis being regulated by the basic helix-loop-113helix transcription factor DvIVS through regulation of the gene expression of chalcone 114 3-hydroxylase, dihydroflavonol 115synthase (DvCHS1),flavanone 4-reductase. 116anthocyanidin synthase, anthocyanidin 3-O-glucoside-6'-O-malonyltransferase and 117 glutathione S-transferase, but not DvCHS2 and chalcone isomerase (Deguchi et al. 2013; 118 Ohno et al. 2011a, 2013). The flavonoid biosynthetic pathway in dahlia is summarised in Fig. 2. The formation of white tips, i.e. the loss of flavonoids in the white parts of petals, 119 120results from simultaneous post-transcriptional gene silencing (PTGS) of two different 121chalcone synthases, DvCHS1 and DvCHS2 (Ohno et al. 2011b). Small RNA of CHS was detected from the white tips of a yellow-white bicolor line 'OriW1' petals, where 122 DvCHS1 expression was absent in whole petals, indicated that the PTGS of DvCHS2 can 123124be induced without DvCHS1 expression (Ohno et al. 2011b). These results implied that causal gene for PTGS of CHS might be DvCHS2, however, the genomic background has 125

not yet been analysed and so the causal gene for simultaneous PTGS of *CHS* remainsunknown.

This study had two objectives: to elucidate the underlying mechanism for petal color lability in bicolor dahlia and to identify the causal gene for simultaneous PTGS of *CHS*. We examined this by performing a molecular analysis of labile petals and leaves in 'Yuino', and comparing the red petals with bicolor petals and flavonoid-rich leaves with flavonoid-poor leaves. We also analysed the genomic background of *DvCHS2*, which was silenced in bicolor petals and flavonoid-poor leaves, among bicolor and non-bicolor cultivars.

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### 137 Materials and Methods

138 *Plant materials* 

The red-white bicolor dahlia cultivar 'Yuino' (Fig. 1) was used for the experiment. Petals and leaves were collected from field- or greenhouse-grown plants in the experimental field of Kyoto University (Kyoto, Japan) and used in the following analyses. In addition, seven other bicolor cultivars ('Matsuribayashi', 'Kazusa-shiranami', 'Santa Claus', 'OriW1', 'OriW2', 'Shukuhai' and 'Kageboshi') and six single-color cultivars that produce petals without white tips ('Kokucho', 'Ms. Noir', 'Yukino', 'Michael J', 'Fidalgo Blacky' and 'Cupid') were also used in the genomic analysis.

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147 Pigment analysis of petals

Bicolor petals were separated into the outside red area and the inner white area using a razor blade and these parts were collected separately. Red petals were also separated in the same way as bicolor petals and collected. The pigment contents of the different petal

151parts were quantified with high-performance liquid chromatography (HPLC). Fresh petal 152parts were homogenised with a mortar and a pestle under liquid nitrogen, following which 1531 mL of extraction solution (5 % hydrochloric acid in 50 % methanol) was added. The 154mixture was then centrifuged at 4°C at 15,000 rpm for 15 min, and the supernatant was 155collected and diluted 50 times with the same solvent. For hydrolysis, 1.2 mL of the diluted solution was boiled at 95°C for 2 h and 20 µL of the hydrolysed solution was injected 156157into the HPLC apparatus. The analysis was performed using an HPLC system (Hitachi L-7100, L-7200, L-7420, L-7500; Hitachi Systems, Ltd., Tokyo, Japan) with a C18 column 158159(Nihon Waters K.K., Tokyo, Japan) that was maintained at 40°C. The detection 160 wavelength was 350 nm for flavones and chalcones, and 520 nm for anthocyanidins. 161Eluant preparation and HPLC analysis proceeded according to Ohno et al. (2011b). The 162assay was performed with six different petals.

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### 164 Isolation of new CHS homologs

165A partial sequence of DvCHS3 (JN556044) was obtained from NCBI database and a 166 partial sequence of *DvCHS4* was obtained from transcriptome data (unpublished data) composed for 'Yuino', 'Michael J' and 'Kokucho' petals, and SRR797209 (Hodgins et al. 167 2014). The rapid amplification of cDNA 3' and 5' ends was performed using a library 168 constructed with the GeneRacer<sup>™</sup> Kit (Invitrogen, Carlsbad, CA, USA) of 'Michael J' 169 170(Ohno et al. 2011a). Primers for full-length cDNAs, genome sequencing and real-time 171RT-PCR were designed (Table S1) and sequenced using 'Yuino' petal RNA and genomic DNA. Sequencing analysis was performed using a BigDye® Terminator v 3.1 Cycle 172Sequencing Kit and a 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, 173174USA).

### 176 Phylogenetic analysis

A phylogenetic tree was constructed based on the open reading frames or amino acids of
various *CHS* genes and several other polyketide synthase genes using the NeighborJoining method (Saitou and Nei 1987).

180 The accession numbers for the DNA sequences were as follows: DvCHS1-1 (AB576660), DvCHS1-2 (AB576661), DvCHS2-1 (AB591825), DvCHS2-2 (AB591826), 181 182DvCHS3-1 (LC223139), DvCHS3-2 (LC223140) and DvCHS4 (LC223141) in D. 183variabilis; AmCHS1 (X03710) in Antirrhinum majus; AtTT4 (NM\_121396) in 184 Arabidopsis thaliana; GhCHS1 (Z38096), Gh2PS (Z38097), GhCHS3 (Z38098) and GhCHS4 (AM906210) in Gerbera hybrida; GmCHS3 (FJ770471) and GmCHS7 185(AK245977) in Glycine max; InCHSD (AB001818) and InCHSE (AB001819) in Ipomoea 186nil; MsCHS1 (L02901) in Medicago sativa; PhCHSA (AF233638) and PhCHSJ (X14597) 187 in Petunia hybrida; VvCHS1 (EF192464), VvCHS3 (AB066274) and VINST1 188(NM\_001281010) in Vitis vinifera; and ZmWHP (X60204) in Zea mays. 189

The accession numbers for the amino acid sequences were as follows: DvCHS1-190 (BAJ14768), DvCHS1-2 (BAJ14769), DvCHS2-1 (BAJ21531), DvCHS2-2 191 1 (BAJ21532), DvCHS3-1 (BAX02592), DvCHS3-2 (BAX02593) and DvCHS4 192(BAX02594) in Dahlia variabilis; AtTT4 (NP\_196897.1) in Arabidopsis thaliana; 193GhCHS1 (CAA86218), GhCHS3 (CAA86220) and GhCHS4 (CAP20328) in Gerbera 194 195hybrida; GmCHS3 (ACN81822) in Glycine max; InCHSD (BAA21787) and InCHSE 196 (BAA21788) in Ipomoea nil; and PhCHSA (AAF60297) and PhCHSJ (CAA32737) in Petunia hybrida. The amino acid sequence of GmCHS7 was estimated from AK245977. 197 198

199 Real-time RT-PCR

200 To analyse petal flavonoid biosynthetic gene expression, bicolor petals were separated

201into the colored outer part and the pure white inner part, and RNA was extracted from 202each. Red single-colored petals were also separated into the same parts and RNA was 203extracted from each of these. Unfolded bicolor and red petals were collected from the 204same region of the same inflorescence, with pairs of petals being collected from three 205different inflorescences. To analyse leaf flavonoid biosynthetic gene expression, we 206obtained six 3-5 cm leaves with or without flavonoid accumulation, as determined from 207 their ABS<sub>400</sub>/ABS<sub>370</sub> scores measured by a spectrophotometer according to Ohno et al. 208(2016): leaves with a score >0.8 were judged to have accumulated an abundance of 209flavonoids, while leaves with scores <0.5 were judged to have accumulated fewer 210flavonoids. The ABS<sub>400</sub>/ABS<sub>370</sub> scores were 1.005, 0.976, 0.969, 0.955, 0.925 and 0.922 in flavonoid-rich leaves, compared with 0.344, 0.269, 0.264, 0.192, 0.178 and 0.114 in 211212flavonoid-poor leaves.

Total RNA was extracted from petals and leaves using Sepasol RNA I Super G 213(Nacalai Tesque, Kyoto, Japan), purified with a high-salt solution for precipitation 214215(Takara Bio Inc., Ohtsu, Japan) and reverse transcribed with ReverTra Ace® (Toyobo, Osaka, Japan), following which 2 µL of 50-fold diluted RT product was used as a 216template for real-time RT-PCR. Real-time RT-PCR was performed with SYBR® Premix 217Ex Taq<sup>TM</sup> II (Takara Bio Inc.) according to the manufacturer's instructions using the 218LightCycler® 480 system (Roche Diagnostics K.K., Tokyo, Japan). The real-time RT-219220PCR was performed as follows: 95°C for 5 min, followed by 45 cycles at 95°C for 10 s 221and 60°C for 30 s. Single-target product amplification was checked using a melting curve. The primers that were used for real-time RT-PCR are shown in Table S2. 222

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### 224 Protein extraction and western blotting

225 Total protein from petals was extracted using Minute Total Protein Extraction Kit for

226Plant Tissues (Invent Biotechnologies Inc., Eden Prairie, MN). Total protein from leaves 227was extracted according to the method of Wang et al. (2010) with minor modification. 228Briefly, 2 g of leaf was ground to a fine powder under liquid nitrogen and then 6 mL 229extraction solution was added. The extraction solution consisted of 50 mM Tris-HCl (pH 2309.0), 2% SDS, 5 mM ascorbic acid, 14 mM β-mercaptoethanol and 1% proteinase 231inhibitor cocktail for plant cell and tissue extracts, DMSO solution (SIGMA-ALDRICH 232Co., St. Louis, MO). The homogenate was filtered through 4 layers of Miracloth 233(Millipore, Billerica, MA) and centrifuged at 4°C at 15,000 rpm for 20 min. The 234supernatant was used as the total protein. Protein concentration was determined by Protein Quantification Kit-Rapid (Dojindo, Kumamoto, Japan). To determine flavonoid 235accumulation, 100-200 mg of leaf were collected from the same leaf or the same branch 236237of protein extracted leaves. The ABS<sub>400</sub>/ABS<sub>370</sub> scores for flavonoid-rich leaves were 1.000, 0.992 and 0.965, while 0.158, 0.120 and 0.252 for flavonoid-poor leaves. 238

239For western blotting analyses, 10 µg of total proteins in the sodium dodecyl 240sulfate polyacrylamide gel electrophoresis gel (the concentration for running gel was 8% 241and for stacking gel was 3%) were transferred to an Immobilon-P membrane (Millipore). The blots were probed with 1:2000 dilution of the primary anti-GmCHS7 peptide (NH2-242C+FRGPSDTHLDSLVGQ –COOH) IgG (rabbit) and 1 : 20000 dilution of the secondary 243goat Anti-rabbit IgG, HRP-linked Antibody (Cell Signalling Technology, Danvers, MA). 244245The immune complexes were visualized by a peroxidase-catalyzed chemiluminescence 246reaction using an ECL Western blotting kit (GE Healthcare Japan, Tokyo, Japan) following the manufacturer's instructions. The chemiluminescence image was obtained 247using a LAS-3000 Mini (Fujifilm, Tokyo, Japan). For positive control, the DvCHS2-1 248coding sequence was subcloned into pET6xHN-C vector, and the DvCHS2-His fusion 249protein was expressed in E. coli and purified via His60 Ni gravity column using pET 250

Express & Purify Kit-His60 (In-Fusion<sup>®</sup> Ready) (Takara Bio inc.). Primers used for
constructing *DvCHS2*-His fusion protein were pET-6xHN-C-DvCHS2-F: 5'TAAGGCCTCTGTCGAGATGGCATCTTCGGTCGATA-3' (forward) and pET-6xHNC-DvCHS2-R: 5'-CAGAATTCGCAAGCTTGGGCGAAATCGGCATGGTA-3'
(reverse).

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257 Analyses of small RNAs

The detection of *CHS* siRNA by RNA gel blot analysis was performed according to Ohno et al. (2011b). The ABS<sub>400</sub>/ABS<sub>370</sub> scores of leaves used for RNA extraction were 0.907, 0.795 and 0.934 for flavonoid-rich leaves, and 0.200, 0.101 and 0.050 for flavonoid-poor leaves.

262For the mapping analysis of petals, small RNAs were extracted from a 5-cm expanded red petal using the MirVana miRNA Isolation Kit (Applied Biosystems) 263264according to the manufacturer's instructions. Small RNAs were sequenced using an 265Illumina Hiseq (Illumina Inc., San Diego, CA, USA), and 18-32 nt small RNAs were mapped onto DvCHS genes (DvCHS1-1, DvCHS1-2, DvCHS2-1, DvCHS2-2, DvCHS3-2661, DvCHS3-2 and DvCHS4) using the Bowtie software without any mismatch. The 267 number of total reads for a red petal was 13,681,764. For the white part of a bicolor petal, 268previous small RNA data (Ohno et al. 2011b) were used and newly mapped onto 269DvCHS3-1, DvCHS3-2 and DvCHS4. 270

For the mapping analysis of leaves, small RNAs were extracted from a flavonoid-rich leaf and a flavonoid-poor leaf using the MirVana miRNA Isolation Kit (Applied Biosystems) according to the manufacturer's instructions. Each leaf was separated along the midrib using a razor, and one half was used for spectrophotometric measurement and HPLC analysis while the other half was used for RNA extraction. The 276ABS<sub>400</sub>/ABS<sub>370</sub> scores were 0.969 and 0.178 for the flavonoid-rich and flavonoid-poor 277leaves, respectively, and flavonoid accumulation was confirmed by HPLC. A small RNA-278enriched sample was used for the small RNA mapping analysis and the rest of the RNA 279fraction that was depleted of small RNAs was used for the RNA-seq analysis described 280below. Small RNAs were sequenced using an Illumina Hiseq (Illumina Inc.) and 18-30 281nt small RNAs were mapped onto DvCHS genes (DvCHS1-1, DvCHS1-2, DvCHS2-1, 282DvCHS2-2, DvCHS3-1, DvCHS3-2 and DvCHS4) using the Bowtie software without any 283mismatch. The number of total reads was 12,046,092 for a flavonoid-rich leaf and 28411,465,948 for a flavonoid-poor leaf.

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#### 286 RNA-seq analysis

RNA-seq analysis was performed with the HiSeq2000 sequencing system (Illumina) using the RNA fraction that had been depleted of small RNAs from the rest of small RNA mapping analysis. The number of total reads was 4,059,802 for a flavonoid-rich leaf and 4,579,744 for a flavonoid-poor leaf. The differentially expressed genes between these samples were evaluated using edgeR (Robinson *et al.*, 2010) with a border value of logFC = 3 and were confirmed by RT-PCR.

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### 294 DNA gel blot analysis

Genomic DNA was extracted from leaves of bicolor and single-color cultivars using
MagExtractor® Plant Genome (Toyobo). A 20-µg DNA sample was digested with a
restriction enzyme (*Bam*HI or *Hin*dIII) and separated on 0.8 % agarose gel in 0.5 ×TAE
buffer, and subsequently blotted to a Hybond-N+ membrane (GE Healthcare Japan) in 20
×SSC buffer. For the probe, the PCR product was purified from agarose gel slices with
the illustra<sup>TM</sup> GFX<sup>TM</sup> PCR DNA and Gel Band Purification Kit (GE Healthcare Japan),

and labelled with the AlkPhos Direct Labelling and Detection System (GE Healthcare
Japan). The probe was hybridised to the membrane at 55 °C overnight. Detection was
conducted with CDP-Star® (GE Healthcare) and the chemiluminescence image was
obtained using a LAS-3000 Mini (Fujifilm).

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### 306 Inverse and allele-specific PCR

307 To isolate unknown genomic region, inverse PCR was performed. Genomic DNA of 308 'Yuino' and 'Michael J' was digested with a restriction enzyme (BamHI, HindIII or XbaI). 309 A 300-ng digested DNA sample was then self-ligated by T4 DNA ligase (Takara Bio Inc.) 310 in a 200-µL volume. Inverse PCR was performed with Blend Taq (Toyobo) or Takara EX 311Taq (Takara Bio Inc.). To analyse allele composition, allele-specific PCR was performed. 312For allele-specific PCR, genomic DNA extracted using MagExtractor -Plant Genome-(Toyobo) from leaves of six bicolor cultivars ('Yuino', 'Matsuribayashi', 'Kazusa-313314 shiranami', 'Santa Claus', 'OriW1' and 'OriW2') and four single-color cultivars ('Kokucho', 'Ms. Noir', 'Yukino' and 'Michael J') was used. The primers that were used 315316 for inverse PCR, sequencing and allele-specific PCR analysis are shown in Table S3.

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318 *Quantification of the genomic region of* DvCHS1 *and* DvCHS2

Genomic DNA was extracted from leaves of four bicolor cultivars ('Yuino', 'Shukuhai', 'Kazusa-shiranami' and 'Kageboshi') and four single-color cultivars ('Kokucho', 'Fidalgo Blacky', 'Michael J' and 'Cupid'). A 50-ng sample of genomic DNA was used as a template for qPCR, which was performed with SYBR Premix Ex Taq II (Takara Bio Inc.) or THUNDERBIRD® SYBR qPCR Mix (Toyobo) according to the manufacturer's instructions using the Light Cycler 480 system (Roche Diagnostics K.K.). The qPCR was performed as follows: 95°C for 2 min, followed by 40 cycles at 95°C for 10 s, 55°C for 5 s and 72°C for 20 s. Single-target product amplification was checked using a melting curve. The primers that were used for qPCR are shown in Table S4. To amplify all the alleles, we designed these primers on the identical sequence among multiple alleles (*DvCHS1-1* and *DvCHS1-2* for *DvCHS1*, and *DvCHS2-1*, *MJ-1* and *MJ-2* for *DvCHS2*).

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332 Results

333 Comparison of pigments in bicolor and red petals

334'Yuino' produces not only red-white bicolor petals but also red single-colored petals (Ohno et al. 2016; Fig. 1). The analysis of hydrolysed extracts showed that the red single-335 336 colored petals contained anthocyanins (cyanidin and pelargonidin), flavones (apigenin 337 and luteolin) and chalcones (isoliquiritigenin and butein), the composition of which was the same as in the red part of bicolor petals (Fig. 3). The amount of anthocyanins in the 338 339 red parts of bicolor petals (outside) was nearly the same as in the corresponding part of 340 red single-colored petals, whereas only a small amount of anthocyanins, flavones and 341chalcones were detected in the white parts (inside) of bicolor petals (Fig. 3). Analysis of the crude extracts showed that the retention time of the peaks of anthocyanins in red 342single-colored petals and the red part of bicolor petals was the same as in 'Kokucho' petals 343(data not shown), which mainly accumulate pelargonidin 3-(6"-malonylglucoside)-5-344 345glucoside and cyanidin 3-(6"-malonylglucoside)-5-glucoside (Deguchi et al. 2016). 346 Therefore, it was suggested that the formation of red single-colored petals in 'Yuino' resulted from a loss of the white parts of the petals by recovering flavonoid biosynthesis 347348capacity.

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### 350 Four chalcone synthase (CHS)-like genes in dahlia

351Ohno et al. (2011b) previously demonstrated that the formation of white tips, i.e. the loss 352of flavonoids from the white parts of petals, results from simultaneous PTGS of two 353 different chalcone synthases, *DvCHS1* and *DvCHS2*. Before analysing the involvement 354of CHS PTGS in the loss of flavonoids in petals and leaves, we first tried to identify all 355CHS-like genes that are expressed in these organs. Two novel CHS-like genes were 356obtained from a database search and transcriptome data, named DvCHS3 and DvCHS4, 357respectively. For DvCHS3, two cDNA sequences were identified from 'Yuino' petals, 358which were named DvCHS3-1 and DvCHS3-2. The open reading frames of both 359DvCHS3-1 and DvCHS3-2 cDNAs were 1191 bp encoding 397 putative amino acid residues. For DvCHS4, one cDNA was detected and its open reading frame was 1170 bp 360 encoding 390 putative amino acid residues. All four CHS genes are composed of two 361362 exons and one intron, which is typical for the polyketide synthase gene family including CHS, and possess a conserved exon-intron junction (Fig. S1). 363

In the phylogenetic tree based on coding region, *DvCHS1*, *DvCHS2* and *DvCHS4* are close to other *CHS* genes, while *DvCHS3* is close to other polyketide synthases such as stilbene synthase (Fig. 4a). By contrast, in the phylogenetic tree based on putative amino acid sequences, DvCHS1 and DvCHS2 are close to other CHS proteins, while DvCHS3 and DvCHS4 are close to other polyketide synthases (Fig. 4b).

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### 370 Flavonoid biosynthetic gene expression in petals and leaves

Comparison of the expression levels of flavonoid biosynthetic genes, including four chalcone synthase genes (*DvCHS1*, *DvCHS2*, *DvCHS3* and *DvCHS4*), between red and bicolor petals showed that *DvCHS1* and *DvCHS2* had significantly lower expression levels in the white part of bicolor petals than in red petals, while *DvCHS3*, *DvCHS4* and other flavonoid biosynthetic genes had almost the same expression levels (Fig. 5a).

376 Western blot analysis also demonstrated that expression levels of CHS protein was lower 377 in the white part of bicolor petals than in red parts (Fig. 5b). By contrast, comparison of 378 the expression levels of flavonoid biosynthetic genes between flavonoid-rich and 379 flavonoid-poor leaves yielded different results, whereby DvCHS2 but not DvCHS1 was 380 significantly down-regulated in flavonoid-poor leaves (Fig. 6a). The expression levels of 381DvCHS1, DvCHS3 and DvCHS4 were low in both leaf types. This result was confirmed 382by the RNA-seq analysis. Only two genes were significantly up-regulated while five 383 genes were significantly down-regulated in the flavonoid-rich leaf compared with the 384flavonoid-poor leaf (Table S5). However, RT-PCR confirmed that six genes were pseudo positive or pseudo negative due to sample bias (data not shown), and so the only gene 385386 that was actually differentially expressed between flavonoid-rich and flavonoid-poor 387 leaves was DvCHS2. Though the difference of signal intensity between flavonoid-rich leaves and flavonoid-poor leaves was not drastic, Western blot analysis also suggested 388 389 that expression levels of CHS protein was lower in flavonoid-poor leaves than flavonoid 390 rich leaves (Fig. 6b).

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392 Analysis of small RNAs

393 Ohno et al. (2011b) previously mapped small RNAs from the white part of a bicolor petal 394 onto DvCHS1 and DvCHS2 (Fig. 7a). In the current study, we used these data to undertake 395a mapping analysis of small RNAs for newly isolated DvCHS3 and DvCHS4 and 396 recalculated them as matched reads per million reads for comparison. Some small RNAs appeared to be mapped onto DvCHS4 (Fig. S2, Table 1); however, this sequence was 397 identical to DvCHS1 and no small RNAs were detected from the DvCHS4-specific 398 399 sequence, suggesting that these small RNAs were derived from DvCHS1 not DvCHS4. Thus, it was demonstrated that DvCHS3 and DvCHS4 were not silenced in the white part 400

401 of a bicolor petal, presumably due to their low expression. The number of small RNAs402 matched to each gene is shown in Table 1.

Next, we sequenced small RNAs from a red single-colored petal and performed a mapping analysis for all alleles of *DvCHS1*, *DvCHS2*, *DvCHS3* and *DvCHS4*. The number of total reads of 18–32 nt was 13,681,764 for a red petal, and small RNAs from the red petal were rarely mapped onto any of the four *DvCHS* genes (Fig. 7b; Fig. S2; Table 1). This result suggests that simultaneous PTGS of *DvCHS1* and *DvCHS2* was suppressed in red petals. Small RNAs sharing high identity with some microRNAs (miRNAs) were detected in both small RNA libraries (Table S6).

To determine whether this suppression of *DvCHS2* in flavonoid-poor leaves was 410 411 post-transcriptional, we first performed an RNA gel blot analysis. Small RNA of DvCHS2 412was detected in flavonoid-poor leaves but not in flavonoid-rich leaves (Fig. 8), which suggested that down-regulation of DvCHS2 in flavonoid-poor leaves occurs post-413414 transcriptionally. This was confirmed by the small RNA mapping analysis (Fig. 9). While small RNAs from a flavonoid-rich leaf were rarely mapped onto each CHS gene (Fig. 9a, 415416 Fig. S3, Table 1), 10,625 or 10,872 of 11,465,948 reads were mapped onto DvCHS2-1 or 417 DvCHS2-2, respectively, in a flavonoid-poor leaf (Fig 9b, Table 1). Small RNAs from 418 flavonoid-poor leaf were rarely mapped onto the other DvCHS genes (Fig. 9b, Fig. S3, Table 1). Therefore, it was demonstrated that DvCHS2 genes are silenced post-419 420transcriptionally in flavonoid-poor leaves.

421

422 Genomic analysis of DvCHS2

423 Since all bicolor dahlia cultivars inevitably exhibit petal color lability (i.e. the loss of a 424 white area), it is evident that occurrence of PTGS of *CHS* genes is important for this 425 process. It has previously been reported that specific parts of the genomic sequence are important for causing endogenous PTGS (Todd and Vodkin 1996; Stam 1997; Kusaba et
al. 2003; Della Vedova et al. 2005; Kasai et al. 2007; Tuteja and Vodkin 2008; Morita et
al. 2012). Thus, genomic analysis was performed.

429The DNA gel blot analysis using BamHI-digested DNA led to one intense bicolor 430cultivar-specific band being detected when the 3' or 5' coding region was used as a probe 431(Fig. 10a); and when HindIII-digested DNA was used, one intense bicolor cultivar-432specific band was detected when the 3' coding region was used as a probe (Fig. 10a). The 433sequence of this band was determined by inverse PCR and identified as the DvCHS2-1 434allele in 'Yuino' (Fig. 10b). Inverse PCR using 'Michael J' genomic DNA identified two other alleles (MJ-1 and MJ-2, respectively), which coincided with the two bands detected 435436 in the DNA gel blot analysis of *HindIII* using the 3' probe in 'Michael J'. To confirm these 437results, genomic PCR was performed using each allele-specific primer. As with the DNA gel blot analysis, the DvCHS2-1 allele was detected in each of the six tested bicolor 438439cultivars but not in the four tested single-color cultivars (Fig. 10c). The MJ-1 and MJ-2 440 alleles were detected in several cultivars, but there was no correlation between allele retention and flower color (Fig. 10c). 441

The qPCR analysis for quantifying the genome amount showed that bicolor cultivars had almost the same amount of the *DvCHS1* genomic region but twice as much of the *DvCHS2* genomic region as single-color cultivars (Fig. 11). This applied not only to the gene body, but also to the 3' flanking region of *DvCHS2*, suggesting that bicolor cultivars retain a duplicated sequence of *DvCHS2*.

447

448

449 **Discussion** 

450 DvCHS2 is a functional CHS in dahlia

451The CHS multigene family contains multiple paralogous CHS genes that function in 452different organs. CHS-like genes exhibit different evolutionary patterns, with early or late 453divergence, and late-diverged CHS-like genes have experienced a more rapid non-454synonymous substitution rate, yielding new enzyme activities in a relatively short period 455of time (Han et al. 2014). In the present study, four CHS-like genes were isolated from dahlia. A phylogenetic analysis based on the nucleotide sequences suggested that 456457DvCHS1 and DvCHS4, or DvCHS2 and DvCHS3 originated from the same gene (Fig. 4a). By contrast, a phylogenetic analysis based on putative amino acid sequences showed that 458459DvCHS3 and DvCHS4 were more closely related to Gh2PS or VvSTS1 than other CHS genes (Fig. 4b), suggesting that these genes diverged from the other CHS genes more 460 461 recently but now exhibit other enzymatic activities.

462In gerbera (Gerbera hybrida), GhCHS1, which is orthologous to DvCHS2, is associated with flavonoid biosynthesis in petals, while GhCHS4, which is orthologous to 463 464 DvCHS1, is associated with vegetative tissues (Helariutta et al. 1995, Deng et al. 2014). 465However, in dahlia, DvCHS2 is expressed in both petals and leaves, and its expression 466 completely coincides with flavonoid biosynthesis, i.e. the loss of expression of DvCHS2 467 in petals results in the loss of anthocyanin, flavone and butein synthesis (Ohno et al. 468 2011b; Fig. 3), and the loss of expression of DvCHS2 in leaves results in the loss of flavonol and butein synthesis (Ohno et al. 2016, 2017). Therefore, it appears that the 469 470DvCHS2 protein may be the functional CHS enzyme in both petals and leaves, and so is 471associated with all flavonoid biosynthesis in dahlia. DvCHS1 is under the regulation of the basic helix-loop-helix transcription factor DvIVS (Ohno et al. 2011a) and retains the 472 preserved sequences of CHS genes (Ohno et al. 2011b), suggesting that it also has CHS 473474activity in dahlia.

475

476 *Phenotypic lability in 'Yuino' is associated with the occurrence of* DvCHS2 *PTGS* 

'Yuino' produces red single-colored petals in addition to original red-white bicolor petals 477478(Fig. 1; Ohno et al. 2016). Almost no flavonoids were detected in the white parts of petals 479(Fig. 3) due to PTGS of both DvCHS1 and DvCHS2 (Ohno et al. 2011b). However, red-480 colored petals accumulated flavonoids (Fig. 3), suggesting that PTGS of CHS was 481suppressed in these. In fact, expression levels of DvCHS1 and DvCHS2 are higher in the 482inner area of red petal than the corresponding pure white area of bicolor petals (Fig. 5a). 483Western blot analysis also suggested that CHS protein expression was recovered in the 484inner area of red petal (Fig. 5b). In red petals, small RNAs of DvCHS2 were rarely detected by RNA gel blot analysis (Fig. 8) and almost no small RNAs were mapped on 485DvCHS genes (Fig. 7b) suggesting PTGS of CHS was suppressed in red petals. 486

487 Ohno et al. (2016) previously detected a strong relationship between inflorescence color and leaf phenotype, whereby red petal-producing plants accumulated 488 flavonoids in their leaves while plants without flavonoids in their leaves produced only 489 490 bicolor petals. This suggests that the formation of the white part of a petal is related to the 491flavonoid biosynthetic potential of the shoot. The flavonoids in leaves have been 492identified as butein and flavonol derivatives by nuclear magnetic resonance analysis (Ohno et al. 2017). In the present study, only DvCHS2 had lower expression in flavonoid-493poor leaves than in flavonoid-rich leaves, whereas other flavonoid biosynthetic genes had 494 495almost the same expression levels (Fig. 6a). The small RNAs of a flavonoid-poor leaf 496 were detected by RNA gel blot analysis (Fig. 8) and mapped onto the DvCHS2 gene (Fig. 497 9b, Table 1), indicating that this suppression of *DvCHS2* occurs post-transcriptionally. Therefore, PTGS of DvCHS2 is associated with the absence of leaf flavonoid 498499 accumulation, indicating that PTGS of CHS in both petals and leaves is tightly linked to 500petal color lability.

501This then leads to the new question of why PTGS of CHS is not occurred in red 502petals and flavonoid-rich leaves. There are two possible explanations for this: CHS-503specific PTGS has ceased or PTGS of all genes has ceased. The small RNAs sharing high 504identity with miR159 or miR165 or unknown small RNAs were detected in both red-505colored petals and the white part of bicolor petals (Table S6), suggesting that general 506PTGS pathway was functional in both petal types. Therefore, it could be that CHS-507 specific PTGS has ceased in red petals. A similar trend was also observed in leaves. It 508was notable that although seven genes were differentially expressed between a flavonoid-509rich and a flavonoid-poor leaf by RNA-seq analysis (Table S5), only DvCHS2 was 510substantially different (Table S5), whereas the other genes were pseudo positive. This 511suggests that the suppression of PTGS in flavonoid-rich leaves is also DvCHS2-specific, 512but not the PTGS component itself.

513In the case of soybean (*Glycine max*), it has been shown that the silencing of CHS gene family members occurs only in the seed coats and not in other organs (Tuteja 514et al. 2004, 2009), and that both primary and secondary CHS siRNAs are not significantly 515516produced in the germinated cotyledon, immature cotyledon, leaf, root, shoot tip or stem 517tissues (Cho et al. 2013). However, in bicolor dahlia 'Yuino', the production of CHS 518siRNAs was not limited to the petal tips but also observed in the leaves. This CHS siRNA-519producing state was assumed to be maintained after vegetative propagation because plants 520that are propagated from flavonoid-poor plants, in which CHS PTGS occurs, produce 521only bicolor petals (Ohno et al. 2016).

522

523 Genomic background for PTGS of CHS genes

In dahlia, two different *CHS* genes were silenced in the white part of bicolor petals: *DvCHS1* and *DvCHS2*. *CHS* siRNA was also detected in the pure white part of the yellow-

white bicolor cultivar 'OriW1', which does not express DvCHS1 in the petals, suggesting that the PTGS of DvCHS2 can be induced without DvCHS1 expression (Ohno et al. 2011b). Thus, it had been presumed that DvCHS2 was a potential causal gene of CHSPTGS in bicolor dahlia and that the silencing of DvCHS1 was incidental due to high homology with some secondary DvCHS2 siRNAs. This was supported by our analysis of the leaves, in which only DvCHS2 was silenced because DvCHS1 is expressed at a relatively low level in both flavonoid-rich and flavonoid-poor leaves (Figs. 6a, 9).

533It has previously been reported that specific parts of the genomic sequence, such 534as inverted repeats, cause endogenous PTGS (Todd and Vodkin 1996; Stam 1997; Kusaba et al. 2003; Della Vedova et al. 2005; Kasai et al. 2007; Tuteja and Vodkin 2008; Morita 535et al. 2012). With respect to PTGS of CHS, inverted repeats of CHS are a potent cause in 536maize (Zea mays; Della Vedova et al. 2005) and soybean (Todd and Vodkin 1996; Kasai 537et al. 2007; Tuteja and Vodkin 2008), while a tandem repeat is a potent cause in petunia 538(Petunia hybrida; Stam 1997; Morita et al. 2012). DNA gel blot analysis suggested the 539existence of a bicolor cultivar-specific DvCHS2 allele, which is identical to 'Yuino' 540541DvCHS2-1 (Fig. 10a, b). This allele was only detected in bicolor cultivars (Fig. 10c), 542suggesting that it is linked to the silencing causal gene.

543Quantification of the genome amount showed that bicolor cultivars contain twice as much of the DvCHS2 region but almost the same amount of the DvCHS1 region as 544 545single-color cultivars (Fig. 11). This suggests that bicolor cultivars have a duplicated 546DvCHS2 region. The 3' proximal DvCHS2 region also doubled in bicolor cultivars (Fig. 11), whereas no repeat sequence of CHS was detected from 3-kb downstream of DvCHS2-5471 (Fig. 10b). The DNA gel blot analysis digested with BamHI detected intense bands of 548approximately 8 kb with the 5' probe and 10 kb with the 3' probe (Fig. 10a), suggesting 549that at least an 18-kb region that includes DvCHS2 is duplicated in bicolor cultivars. This 550

region is a strong candidate for the *CHS* PTGS causal gene, but further analysis is required
to confirm this.

553

554 Candidate mechanism for petal color lability in bicolor dahlias

555The causal gene of PTGS of DvCHS is presumed to be DvCHS2 (see above), and so petal color lability in bicolor dahlias could be interpreted as resulting from the switching on 556557and off of DvCHS2 PTGS. Since all shoot organs including petals and leaves are 558differentiated from the shoot apical meristem (SAM), this switching could occur in the 559SAM. Consequently, here we propose a candidate model of SAM state for each flower color pattern. This model is based on the observation that when both red petals and bicolor 560561petals occur in the same inflorescence, red petals were located inevitably in the outer 562whorls rather than bicolor petals, and the inverse pattern was never observed (Ohno et al., 2016). This indicates an obvious direction for the SAM state from red petal to bicolor 563petal. Considering the strong link between petal color and leaf flavonoid accumulation 564(Ohno et al. 2016), the SAM state of single-colored petal formation and flavonoid-rich 565566leaves could be defined as a 'switch OFF' of DvCHS2 PTGS, while the SAM state of 567 bicolor petal formation and flavonoid-poor leaves is a 'switch ON' of *DvCHS2* PTGS. Therefore, when the SAM maintains a 'switch OFF' state, the inflorescence will be a 568single color due to the absence of DvCHS2 PTGS, whereas when SAM maintains a 569570'switch ON' state, the inflorescence will be bicolor as a result of DvCHS2 PTGS. 571Furthermore, if the SAM maintains a 'switch ON' state sectorally, the inflorescence will be mixed with sectorial single-colored petals; and if the switch of CHS PTGS turns 'ON' 572after the formation of outer whorl petals, the inflorescence will be mixed with single-573colored petals in the outer whorls. Original bicolor cultivars may change to 'switch ON' 574before formation of the inflorescence. The timing of switching depends on the plant, 575

because plants that produce flavonoid-rich leaves could bloom both bicolor and red
inflorescences (Ohno et al. 2016). In this case, it is assumed that if switching occurs
before flowering, the inflorescence will be bicolor, whereas if switching does not occur,
the inflorescence will be red.

580The above model can explain the phenotypic relationship between petal color and leaf flavonoid accumulation. However, another factor is required to explain the 581582lability itself. The phenomenon of color lability is not limited to dahlias, but is also found 583in other plants, such as *Rhododendron* spp. and transgenic petunia, suggesting that there 584may be a common mechanism across species. Transgenic petunia plants carrying a sense CHS transgene (Jorgensen 1995) or antisense CHS transgene (van der Krol et al. 1990) 585586also exhibit flower color lability. In co-suppressed petunia carrying a sense CHS transgene, 587 the expression of CHS is suppressed in the leaf tissues of white branches and reverted in the leaf tissues of the revertant violet branches (Napoli et al. 1990). In this phenotypic 588reversion, it was demonstrated that epigenetic changes that interfere with the initiation of 589transgene transcription lead to a reversion of the PTGS phenotype (Kanazawa et al. 2007). 590591Therefore, an endogenous factor is inferred to be involved in this switch control, perhaps 592an epigenetic factor such as DNA methylation. When the gene or locus that triggers DvCHS PTGS is activated in the SAM, an endogenous factor might fully or sectorally 593suppress its activation in the SAM, resulting in petal color lability. Recently, it was 594595reported that soybean Argonaute5 (AGO5) affects the distribution of siRNAs targeting 596the CHS genes in the seed coat (Cho et al. 2017). In nonfunctional AGO5 background (k1), production of CHS siRNAs was suppressed even in the presence of the normally 597 dominant I allele. Thus, not only CHS itself but also AGO and other components of RNA-598induced silencing complex is a candidate trigger gene of CHS PTGS. Isolation of the gene 599or locus that triggers DvCHS PTGS would be of great value in further elucidating the 600

601	mechanism behind color lability.
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620	Supplemental materials
621	<b>Table S1.</b> Primers used for the isolation of <i>DvCHS3</i> and <i>DvCHS4</i> .
622	Table S2. Primers used for real-time RT-PCR.
623	Table S3. Primers used for identification of the DvCHS2 flanking region.
624	Table S4. Primers used for qPCR for genomic quantification.
625	Table S5. Genes that showed significantly different expression levels between a

- 626 flavonoid-rich leaf and a flavonoid-poor leaf in *Dahlia variabilis* 'Yuino'.
- **Table S6.** Top 80 most highly abundant small RNAs for which the number of total reads
- 628 per million (RPM) was over 1000 in a red petal and the white part of a bicolor petal of
- 629 Dahlia variabilis 'Yuino'.
- **Fig. S1.** Genome structure of *DvCHS1–DvCHS4*.
- Fig. S2. Mapping of chalcone synthase (*CHS*) small RNAs from petals onto *DvCHS3* and *DvCHS4*.
- 633 Fig. S3. Mapping of chalcone synthase (CHS) small RNAs from leaves onto DvCHS3
- 634 and *DvCHS4*.

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### 651 **Figure legends**

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Fig. 1. Petal color lability in field-grown *Dahlia variabilis* 'Yuino'. Both bicolor petalsand red petals were produced in the inflorescence on the right.

655

Fig. 2. Flavonoid biosynthetic pathway in dahlia (*Dahlia variabilis*). This figure was
based on Ohno et al. (2011a) with some modifications. ANS, anthocyanidin synthase;
CH3H, chalcone 3-hydroxylase; CHI, chalcone isomerase; CHR, chalcone reductase;
CHS, chalcone synthase; DFR, dihydroflavonol 4-reductase; F3H, flavanone 3hydroxylase; F3'H, flavonoid 3'-hydroxylase; FLS, flavonol synthase; FNS, flavone
synthase; GST, glutathione S-transferase; GT, anthocyanidin glucosyltransferase; 3MaT,
anthocyanidin 3-*O*-glucoside-6"-*O*-malonyltransferase.

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**Fig. 3.** Amount of flavonoid per 100 mg fresh weight (FW) of red and bicolor petals of *Dahlia variabilis* 'Yuino'. Bicolor petals were separated into the outside red area (Bicolor out) and the inner white area (Bicolor in) and collected separately. Red petals were also separated in the same way as bicolor petals, namely separated into the outside area (Red out) and the inner area (Red in). All data represent the mean  $\pm$  SE (n = 6). Cy, cyanidin; Pg, pelargonidin; Lt, luteolin; Ap, apigenin; Iso, isoliquiritigenin; Bt, butein.

670

**Fig. 4.** Phylogenetic tree for chalcone synthase (CHS). a, Phylogenetic tree based on the coding regions; b, phylogenetic tree based on putative amino acid sequences. The bootstrap values of 1,000 retrials are indicated on each branch and the bar indicates a genetic distance of 0.1. The abbreviation in front of each coding region/protein indicates the plant species: Am, *Antirrhinum majus*; At, *Arabidopsis thaliana*; Dv, *Dahlia*  676 variabilis; Gh, Gerbera hybrida; Gm, Glycine max; In, Ipomoea nil; Ms, Medicago
677 sativa; Ph, Petunia hybrida; Vv, Vitis vinifera; Zm, Zea mays. Proteins are abbreviated as
678 follows: STS, stilbene synthase; 2PS, 2-pyrone synthase.

679

680 Fig. 5. Expression analysis of flavonoid biosynthetic pathway in petals. Bicolor petals 681 were separated into the outside red area (Bicolor out) and the inner white area (Bicolor 682in) and collected separately. Red petals were also separated in the same way as bicolor 683 petals, namely separated into the outside area (Red out) and the inner area (Red in). a, 684 Relative expression levels of flavonoid biosynthetic genes and transcription factors in red 685 and bicolor petals of Dahlia variabilis 'Yuino' by real-time RT-PCR. All data represent 686 the mean  $\pm$  SE (n = 3). Each value represents the expression level relative to that of inside 687 part of red petal (Red in). DvActin was used as the internal standard. n.d., not detected. b, Western blot analysis of CHS in petals. P. C. (positive control): recombinant DvCHS2-688 689 6xHN-C. Two biological replications were performed for each part.

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**Fig. 6.** Expression analysis of flavonoid biosynthetic pathway in flavonoid-rich and flavonoid-poor leaves. a, Relative expression levels of flavonoid biosynthetic gene in leaves of *Dahlia variabilis* 'Yuino' by real-time RT-PCR. Scores were calculated relative to flavonoid-poor leaves (= 1). The vertical bars indicate the SE (n = 6). b, Western blot analysis of CHS protein in leaves. P. C. (positive control): recombinant DvCHS2-6xHN-C. Three biological replications were performed for each flavonoid-rich leaf and flavonoid-poor leaf.

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Fig. 7. Mapping of chalcone synthase (*CHS*) small RNAs from petals onto *DvCHS1* and *DvCHS2*. a, white part of a bicolor petal. b, a red petal; Small RNAs of 18–32 nt with a

100% match to *DvCHS1-1*, *DvCHS1-2*, *DvCHS2-1* or *DvCHS2-2* were mapped onto
either the sense (above the *x*-axis) or antisense (below the *x*-axis) strand. The number of
total reads of 18–32 nt was 17,455,041 for the white part of a bicolor petal and 13,681,764
for a red petal. To compare the white area of a bicolor petal with a red petal, values for *DvCHS1-1*, *DvCHS1-2*, *DvCHS2-1* and *DvCHS2-2* in the white part of a bicolor petal
were obtained from Ohno *et al.* (2011b) and recalculated as the number of matched reads
per million reads.

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Fig. 8. Detection of chalcone synthase (*CHS*) small RNAs in *Dahlia variabilis* 'Yuino' leaves by RNA gel blot analysis. Three biological replications were performed for each flavonoid-rich leaf and flavonoid-poor leaf. Whole part of bicolor petals were used as a positive control.

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**Fig. 9.** Mapping of chalcone synthase (*CHS*) small RNAs from leaves onto *DvCHS1* and *DvCHS2*. a, a flavonoid-rich leaf; b, a flavonoid-poor leaf. Small RNAs of 18–30 nt with a 100% match to *DvCHS1-1*, *DvCHS1-2*, *DvCHS2-1* or *DvCHS2-2* were mapped onto either the sense (above the *x*-axis) or antisense (below the *x*-axis) strand. The number of total reads of 18–30 nt was 12,046,092 for a flavonoid-rich leaf and 11,465,948 for a flavonoid-poor leaf.

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Fig. 10. Genomic structure of *DvCHS2*. a, DNA gel blot analysis of *DvCHS2*; the blot
was hybridised with a 5' probe (left two panels) and a 3' probe (right two panels). b,
Location of primers, probes and restriction enzyme recognition sites of *DvCHS2* alleles;
B and H indicate restriction enzyme recognition sites of *Bam*HI and *Hin*dIII; arrows
indicate the position of the primers used in c. c, genomic polymerase chain reaction (PCR)

726	analysis of allele-specific sequences in the 3' flanking region of DvCHS2. Numbers above
727	the lanes indicate each cultivar: 1, 'Yuino'; 2, 'Matsuribayashi'; 3, 'Kazusa-shiranami';
728	4, 'Santa Claus'; 5, 'OriW1'; 6, 'OriW2'; 7, 'Kokucho'; 8, 'Ms. Noir'; 9, 'Yukino' and
729	10, 'Michael J'.
730	
731	Fig. 11. Quantification of the <i>DvCHS1</i> and <i>DvCHS2</i> genomic regions. ①-⑤ indicates
732	position of designed primers. Primers were designed on the identical sequence among
733	multiple alleles (DvCHS1-1 and DvCHS1-2 for DvCHS1, and DvCHS2-1, MJ-1 and MJ-
734	2 for <i>DvCHS2</i> ). qPCR analyses to quantify the amount of the genomic regions of <i>DvCHS1</i>
735	and <i>DvCHS2</i> . Data represent the mean $\pm$ SD scores of three technical replicates of three
736	biological replicates relative to 'Yuino' (= 1).
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## **Table**

## 

		Pet	tal	Lea	af
		Red petal	White part of bicolor petal	Flavonoid-rich leaf	Flavonoid-poor leaf
Tota	l Read	13,681,764	17,455,041	12,046,092	11,465,948
	CHS1-1	665	63,361	1	7
	CHS1-2	656	61,594	2	2
	CHS2-1	483	371,899	83	10,625
Gene	CHS2-2	337	296,406	36	10,872
	CHS3-1	1	2	0	1
	CHS3-2	1	0	0	0
	CHS4	11	148	0	0

- **Figures**
- **Fig. 1**













b





42kDa



b

# Red petal







939 Fig. 9

b



# Flavonoid-poor leaf

300

200

100

100

0 T Exon1





Exon2

100bp

UTR

- 941 Fig. 10





### 956 Supplemental materials

- 957 **Table S1.** Primers used for the isolation of *DvCHS3* and *DvCHS4*.
- 958 **Table S2.** Primers used for real-time RT-PCR.
- 959 **Table S3.** Primers used for identification of the *DvCHS2* flanking region.
- 960 **Table S4.** Primers used for qPCR for genomic quantification.
- 961 Table S5. Genes that showed significantly different expression levels between a
- 962 flavonoid-rich leaf and a flavonoid-poor leaf in *Dahlia variabilis* 'Yuino'.
- 963 **Table S6.** Top 80 most highly abundant small RNAs for which the number of total reads
- 964 per million (RPM) was over 1000 in a red petal and the white part of a bicolor petal of
- 965 Dahlia variabilis 'Yuino'.
- 966 **Fig. S1.** Genome structure of *DvCHS1–DvCHS4*.
- Fig. S2. Mapping of chalcone synthase (CHS) small RNAs from petals onto DvCHS3 and
  DvCHS4.
- 969 Fig. S3. Mapping of chalcone synthase (CHS) small RNAs from leaves onto DvCHS3
- 970 and *DvCHS4*.

### 972 Supplemental Figure legends

973



Fig. S2. Mapping of chalcone synthase (*CHS*) small RNAs from petals onto *DvCHS3* and *DvCHS4*. a, white part of a bicolor petal. b, a red petal; Small RNAs of 18–32 nt with a
100% match to *DvCHS3-1*, *DvCHS3-2* or *DvCHS4* were mapped onto either the sense

979 (above the *x*-axis) or antisense (below the *x*-axis) strand. The number of total reads of 18–

- 980 32 nt was 17,455,041 for the white part of a bicolor petal and 13,681,764 for a red petal.
- 981 Fig. S3. Mapping of chalcone synthase (CHS) small RNAs from leaves onto DvCHS3

and DvCHS4. a, a flavonoid-rich leaf; b, a flavonoid-poor leaf. Small RNAs of 18–30 nt

983 with a 100% match to DvCHS3-1, DvCHS3-2 or DvCHS4 were mapped onto either the

984 sense (above the *x*-axis) or antisense (below the *x*-axis) strand. The number of total reads

985 of 18–30 nt was 12,046,092 for a flavonoid-rich leaf and 11,465,948 for a flavonoid-poor

986 leaf.

989

990

Table S1 Primers used for the isolation of DvCHS3 and DvCHS4 Gene Purpose sequence (5'-3') Name Orientation GTGCTCCGCAGGAGGCATGGTCCTT Race CHS3 3'Race sense CHS3 3'Race nested GGCTCGCGTGTACTTGTTGTCTGCT Race sense CHS3 5'Race GCACGCGCCTACTCGCCCTCAACTT Race antisense CAGCTCCACCTCGTCGAGTATGCGT CHS3 5'Race nested Race antisense CHS3 3'Race-Re Race CACGTTGATTCTTTGGTCGGCCAAG sense DvCHS3 Race CHS3 3'Race-1 sense GGTCTACGTGACTGGAATTCGATGT Sequence CHS3 Full-F sense ACACATTCTTCAATAGATCAAGTTA Sequence CHS3 Full-R antisense TTACTTATTAATTTACTACCAATCA Sequence CHS3 Full-R2 antisense CATTATTACATCAACCGTTACTTAT CHS3-178F CAACGACCATATGATTGATCTTAAA Sequence sense

	Oequence	0103-170	361136	CARCOACCATATOATTOATTOATTOAT
	Sequence	CHS3-446R	antisense	TTCCGGATGTGGTGCAAAAGATGAG
	Race	CHS4 3'Race	sense	TGGCTCGGTCCTTCGTTTGGCCAAA
	Race	CHS4 3'Race nested	sense	TTTTGGGTTGCGCATCCGGGTGGCC
	Race	CHS4 5'Race	antisense	ACCGAAGGATTAAGATCCAGGAGTT
	Race	CHS4 5'Race nested	antisense	CAATATCTCTTCGGTTAAGTACATG
DUCHSA	Sequence	CHS4 Full-F	sense	CACCACATACAAATTGTAACTTCAC
DV01134	Sequence	CHS4 Full-R	antisense	GTATGAAATTTCATATTATGTAATA
	Sequence	CHS4 163F	sense	AAAAAGTGAGCATATGAAGGAGCTC
	Sequence	CHS4 Int 1R	antisense	TCATTAAGAAACTAAAAAAGCCCAA
	Sequence	CHS4 Int 1F	sense	CACGGTTCAATGATAACTTACAAAG

antisense

CTTATTCCAATTTTGAACTTTTG

CHS4 Int 2R

Sequence

991

Table S2. Prir	ners used for real-time RT-PCR.	
Genes	Forward primers	Reverse primers
DvCHS1	CATGTGCTAAGCGAATACGG	CCTCTCCGGTGGTATTGAAC
DvCHS2	TGTCCCAACTACCATGCCGATTTC	TTACACATTAAAATGACACAGTGA
DvCHS3	CACCGGTGAAGGTTTAGATTGGGGT	CATTATTACATCAACCGTTACTTAT
DvCHS4	TTGGTATGCCCTATTTTCATCATGC	AAATTACATGAACAAAACATGTTT
DvCHI	AGAAGCTGGGAATGCAGTGT	GAGATCTGAGAGCCTTGATGC
DvF3H	TTGGAGGGAGATTGTGACCT	GGCCCATTAACTCCTTGCTA
DvDFR	CAACTTCCGGTCTATGACGAG	TTTCGGCCAATGTTTTTGAC
DvANS	GCTCCAACTCTTCTACAACG	GAAATCCTGACCTTCTCCTT
DvGT	AAACATCACCCTTCTTACTCT	TTGAAAAGCGCGATGGATGTT
Dv3MaT	AAATACGAAGTTGTTTCAATC	TTGCACTTTCTAATCCATCAT
DvGST	ATGTGGTGGGATGATATTTCAAACA	AACATTTATTTGTGAGTCACATACA
DvF3'H	GTAGTTATACGCAATATGCTC	CATAACTGCCTTACTATTGTAC
DvFNS	GTGTGTTTCCCTTTGCTTCGTAAAA	GCGAAGGGAAACACACTAGATTCGT
DvMYB1	GTTCACTACTTTAGCAAACG	GACTTTGATATCAACCGGAT
DvDEL	ATCTAAGTTAAAGAGTTGTACAGC	TGAAACTTGGAAAATTGGACTCAA
DvIVS	CATAACCCAAGTAAAGAAAGCCATT	CATCCATTTTTAAATTGTTTGTGGT
DvWDR1	AGGCGTTGTGGAAACTCAAT	TTATCGCGAAGGTCGAAAAC
DvActin	TGCTTATGTTGGTGATGAAG	CCCTGTTAGCCTTAGGATTT

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	Table S3. P	rimers used for identificatio	n of the DvCHS2 flanking region.		
	Cultivar	Target region	Purpose	Name	Sequence (5'-3')
	Yuino	CHS2 5' flanking region	Inverse PCR-F	CHS2-3'Race-nested	GCTCGGTAAAGATGCGGCTGTCAAA
			Inverse PCR-R	CHS2 Right Walk-2	CCACGTCCTGACGGGCGTCCAAAGA
		CHS2 3' flanking region	Inverse PCR-F	CHS2 1146F	AAGGTTTGGATTGGGGTGTTCTGTT
			Inverse PCR-R	CHS2 924R	GGCGAAAACGCTTGCACCAACGCCT
			Inverse PCR-2nd-F, Sequencing	CHS2 3'Genome 6F	AATGTAATTGTCTATTTTGTTCTCT
			Inverse PCR-2nd-R	CHS2 762R	CGCTCAGTTGTCAAGTCC
			Sequencing	CHS2 3'Genome 1F	GTTGTGTTGTATGGTTCTGCTTTCT
			Sequencing	CHS2 3'Genome 2F	CCGGAGGTGCCACGCGGGGCCTTTT
			Sequencing	CHS2 3'Genome 3F	CGTGGCAATTAAACACGTTCACCT
			Sequencing	CHS2 3'Genome 4F	TCGCAAGATAATTCACGCGTGGCAA
			Sequencing	CHS2 3'Genome 5F	AGATCATAAGCGGGACG
			Sequencing	CHS2 3'Genome 7F	TCCGGCGTGAGAATCAGTATACAA
			Sequencing	CHS2 3'Genome 8F	TTGTTTTTGCGGGATGACGTCATGT
			Sequencing	CHS2 3'Genome 9F	TCACGCAAGTTCATCCGTTGGGGAT
			Sequencing	CHS2 3'Genome 1R	GGAGAAGAAAGAACAAGAAGAAAAC
			Sequencing	CHS2 3'Genome 2R	AACAGTTGACCCGACACATTCATAT
			Sequencing, allele-specific PCR	CHS2 3'Genome 3R	GGAAACACAAAATGTGTCAAAAACT
			Sequencing	CHS2 3'Genome 4R	CCTCACGCGCACCATCCCCAACGG
			Sequencing	CHS2 3'Genome 5R	AATACCAAACCCCATCATACGGTAA
			Sequencing	CHS2 3'Genome 6R	ACTAAGGACGGACGCAGCGGAACC
			Sequencing	CHS2 3'Genome 7R	GGAGCACCAGCCATTCTTGAAAAAT
			Sequencing	CHS2 3'Genome 8R	ATGGAGGAAGCATTGTTAATTAGT
	Michael J	CHS2 3' flanking region	Inverse PCR-F, allele-specific PCR	CHS2-1216F	TGTCCCAACTACCATGCCGATTTC
			Inverse PCR-R	CHS2 762R	CGCTCAGTTGTCAAGTCC
			Sequencing	CHS2-Inv-MJR-Bam-Clone1-1R	GAGGGATATAAGTTTGATAAACTT
			Sequencing	CHS2-Inv-MJR-Bam-Clone2-1R	TATATGCTGTATGACTATAGTAGA
			Sequencing, allele-specific PCR	CHS2-Inv-MJR-Bam-Clone1-2R	TAACCAAGGGGGAGTGTTATAAA
1002			Sequencing, allele-specific PCR	CHS2-Inv-MJR-Bam-Clone2-2R	AGATTGATTAACAATCGAAGTTA

Table S4. Pr	Table S4. Primers used for qPCR for genomic quantification.		
Genes	Forward primers	Reverse primers	
DvCHS1	GGTGACGGTGCAGCCGCGATCAT	CGCGCCTCCACTATCCGGTAGAATA	
DvCHS1 ②	CATGTGCTAAGCGAATACGG	CCTCTCCGGTGGTATTGAAC	
DvCHS2③	TCTTATTACTGCTCGCAATATCTT	GGCGGGGTTGCAGTGCCGATGGCAA	
DvCHS2@	AAGGTTTGGATTGGGGTGTTCTGTT	AGTTAGGGCGAAATCGGCATGGTA	
DvCHS25	TCGCAAGATAATTCACGCGTGGCAA	ATGGAGGAAGCATTGTTAATTAGT	

	logFC	logCPM	FDR	ID	Description by Blastx
Up-regulated in	4.10	10.08	0.001	c15993_g1	Yuino CHS2-1 (Dahlia pinnata) (AB591825)
flavonoid-rich leaf	3.65	6.19	0.031	c13892_g2	PREDICTED: 23 kDa jasmonate-induced protein-like (Malus domestica)
Lin regulated in	-5.23	4.25	0.031	c15904_g1	BTB and TAZ domain protein 1 (Arabidopsis thaliana) (NP201121)
	-4.33	4.77	0.031	c15650_g1	transcription factor bHLH35 (Arabidopsis thaliana) (NP568850)
op-regulated in flovonoid poor loof	-4.23	7.01	0.002	c15638_g1	RESPONSE TO LOW SULFUR 2 (Arabidopsis thaliana) (NP197854)
navonoiu-poor ieai	-4.01	5.17	0.031	c15986_g1	sugar transporter ERD6-like 16 (Arabidopsis thaliana) (NP568367)
	-3.65	8.09	0.012	c19320_g2	5'-adenylylsulfate reductase 1 (Arabidopsis thaliana) (NP192370)

Table S6. Top 80 most highly abundant small RNAs for which the number of total reads per million (RPM) was over 1000 in a red petal and the white part of a bicolor petal of Dahlia variabilis 'Yuino'.

Sequence	Length	Description	RPM Sum	Red p	etal	White part of a b	icolor petal
TTCCACGGCTTTCTTGAACTG	21		34936	Read count	RPM 22611	Read count	RPM 2225
TCGGACCAGGCTTCATTCCCC	21	miR165	28151	364709	26657	26077	1494
GGTTTCATTGTCACTTGACTG	21		10472	142327	10403	1204	69
CTGGATTATGACTGAACGCCT	21		4850	66081	4830	349	20
ATTCAGACGCTATGAACCATTAAG	24		3671	244	18	63768	3653
TTTGGATTGAAGGGAGCTCT	20	miR159a	3522	7653	559	51719	2963
GTTGAGGGCACGTCTGCCTGGGCGTCACG	30		3204	15345	1122	36352	2083
TCATACACCTITICCAAATCCGC	22		3029	27701	51	51982	2978
	21		2925	3//91	2/02	Z041 45084	2634
TTGAGGGCACGTCTGCCTGGGCGTCACGC	29		2596	4091	299	40100	2004
CAGAATCCGGGCTAGAAGCGA	21		2471	31129	2275	3424	196
TTCAACATCTGATCATACACC	21		2354	27948	2043	5438	312
GCGACCCCAGGTCAGGCGGGACT	23		2288	31133	2276	218	12
TAAAACGACTCTCGGCAACGGATATCTC	28		2132	8987	657	25745	1475
TICGTIGICIGITCGACCTIG	21		2114	5320	389	30108	1725
AAGAAGAACTITGATGACGG	21		2005	22575	1650	6937	2000
TGGATTGTGCTGCATCGTCTG	21		2005	371	27	34519	1978
TCGGACCAGGCTTCATTCCTC	21		1991	17630	1289	12267	703
GCCCGCGACGTCGCGAGAATTCCACTGAAC	32		1952	26559	1941	188	11
TGGAACAATGTAGGCAAGGGAAGTCGGCA	29		1893	16	1	33018	1892
AGATTCAGCCCTGCGTCGCTCAGATTCGT	29		1890	25510	1865	451	26
GTICGAGAAAACTGTGGGAAA	21	miD167	1853	5965	436	24730	1417
TAAAACGACTCTCGGCAACGGATATC	21	miR 167	1793	1230	90	28722	1703
AACGACTCTCGGCAACGGATATCTCGGCT	20		1710	21607	1579	2408	138
TAAAACGACTCTCGGCAACGGA	22		1688	18100	1323	6374	365
CATAGTAGCCAAGGAAGACGA	21		1640	57	4	28555	1636
TGGACCGACGGATACTCATCT	21	DvCHS2	1624	0	0	28351	1624
CCAGGGATCAGCGGATGTTGC	21		1605	21847	1597	139	8
CAAGTGACAATGAAACCGCCT	21		1594	21718	1587	119	7
TTGTGACGAAGATGGCGAGAA	21		1561	4498	329	21507	1232
TAAAACGACTCTCGGCAACGGATATCTCGG	31		1528	3520	258	22175	1270
AGTTAAAAAAGCTCGTAGTTGGACC	20		1502	19781	1446	987	427
TTTGGATTGAAGGGAGCTCTA	21	miR159a	1453	4474	327	19656	1126
TTTAAGAAGATGGAACGTCAG	21	DvCHS2	1451	0	0	25320	1451
GCAGTTAAAAAGCTCGTAGTTGGACC	26		1441	18803	1374	1169	67
TGGAACAATGTAGGCAAGGGAAGTCGGC	28		1407	18	1	24533	1405
ATGGAACAATGTAGGCAAGGGAAGTCGGC/	30		1399	1399	102	22637	1297
	21	miR166n	1366	18584	1358	143	1222
CAACCTITIGTTIGGGGGCGGT	21	DVCHS2	1307	325	24	23204	1333
TGGAACAATGTAGGCAAGGGAAGTCGG	27	0101102	1342	7	1	23424	1342
ATCCGGTTAGGATCGATCTAAACCAGCC	28		1329	13294	972	6235	357
AGAAACGATTGTCCTTACTTT	21		1327	17798	1301	459	26
TGACAGAAGAGAGTGAGCAC	20	miR156a	1313	2525	185	19689	1128
TTCCACAGCTTTCTTGAACTT	21		1283	16903	1235	837	48
TTTGCACTITTGATATCACGG	21		1278	3589	262	17721	1015
TTAAGAGGGGCTTGAACACGG	21		1209	30	2	22107	1207
CACGAACTITGATATCTAGGC	21		1260	382	28	21502	1232
AACGACTCTCGGCAACGGATATC	23		1241	14076	1029	3700	212
TCTGAACAGAAGATGGACCACC	22		1233	305	22	21127	1210
TAAAACGACTCTCGGCAACGGATATCTCGG	32		1226	8756	640	10226	586
GAGAGACCTATTAACGAGCCT	21		1222	16693	1220	37	2
TGGAACAATGTAGGCAAGGGAAGTCGGCA	30		1215	5	0	21200	1215
CTCTAAGTCAGAATCCGGGGCT	21		1176	16047	1173	49	3
TTECAETTAAAAAGCTCGTAETTEGACC	21		11/1	15021	1098	19431	1113
GATGGAACAATGTAGGCAAGGGAAGTCGG	30		1126	97	7	19539	1119
TCCATTGTCGTCCAGCGGTTAGGATA	26		1108	14793	1081	472	27
CAGAAGACTCAGCTCGCTCCT	21		1095	8263	604	8578	491
TGGAACAATGTAGGCAAGGGAAGTCGGCA	31		1087	9	1	18968	1087
CAATGTAGGCAAGGGAAGTCGG	22		1059	20	1	18468	1058
CGTGGACCGACGGATACTCAT	21	DvCHS2	1059	0	0	18482	1059
ATCCGGTTAGGATCGATCTAAACCAGCCC	29		1052	12339	902	2626	150
CGGATACTCATCTTGATTGC	21	DVCHes	1020	13	1	18120	1049
TAAAACGACTCTCGGCAACGG	21	0101102	1035	6294	460	10043	575
TAAAACGACTCTCGGCAACGGATATCT	27		1033	3520	257	13547	776
TTCCGAGCCTACCGATTCCAGC	22		1033	3514	257	13543	776
GAGATTCAGCCCTGCGTCGCTCAGATTCGT	30		1033	13918	1017	269	15
TGGAACAATGTAGGCAAGGGAAGTCGGCA	32		1028	8	1	17930	1027
ATGGAACAATGTAGGCAAGGGAAGTCGGC	29		1021	231	17	17532	1004
TTAAAGAAGAACTTTGATGACG	20		1015	12/8	93	16083	921
CAACAAGGCTGTAGCGAGGGG	22		1014	08 1323	4	15921	912
CAGTTAAAAAGCTCGTAGTTGGACC	25		1008	13077	956	919	53

1015 Total read of red petal and the white part of the bicolor petal was 13,681,764 and 17,455,041, respectively.

Fig. S1



1037 Fig. S2

а

# White part of bicolor petal



b

# Red petal





1039 Fig. S3





# Flavonoid-poor leaf



60			
50	CHS3-2		
40			
30			
20			100hp
10			
0		Evon?	
0		EXONZ	
10			

1040