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Gene regulation of anthocyanin biosynthesis in two blood-flesh peach (*Prunus persica* (L.) Batsch) cultivars during fruit development^{*#}

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Abstract: The blood-flesh peach has become popular in China due to its attractive anthocyanin-induced pigmentation and antioxidant properties. In this study, we investigated the molecular mechanisms underlying anthocyanin accumulation by examining the expression of nine genes of the anthocyanin biosynthesis pathway found in the peach mesocarp. Expression was measured at six developmental stages in fruit of two blood-flesh and one white-flesh peach cultivars, using quantitative reverse transcription polymerase chain reaction (qRT-PCR). Results show that the expression of the chalcone synthase (*CHS*) gene was closely related to anthocyanin accumulation in both of the blood-flesh peaches. In the white-flesh peach, we found that the transcription level of phenylalanine ammonia-lyase (*PAL*) during fruit development was much lower than that in the blood-flesh peach, even though all other genes of the anthocyanin biosynthesis pathway were highly expressed, suggesting that the *PAL* gene may be limiting in anthocyanin production in the white-flesh peach. Moreover, the transcription levels of the *CHS* and UDP-glucose-flavonoid 3-O-glucosyltransferase (*UFGT*) genes were markedly up-regulated at three days after bag removal (DABR) in the blood-flesh peach, suggesting that *CHS* and *UFGT* are the key genes in the process of anthocyanin biosynthesis for both of the blood-flesh peaches. The present study will be of great help in improving understanding of the molecular mechanisms involved in anthocyanin accumulation in blood-flesh peaches.

Key words:Blood-flesh peach, Anthocyanins, Bagging, Gene expressiondoi:10.1631/jzus.B1400086Document code: ACLC number: S662.1

1 Introduction

Anthocyanins synthesized by the phenylpropanoid pathway have been well characterized in different plant species (Pandey *et al.*, 2014). The major anthocyanins include pelargonidin (orange or red), cyanidin (pink or red), and delphinidin (purple or blue) and their formation depends on their hydroxylation pattern (Niu *et al.*, 2010). Anthocyanins, as well as other health-promoting phytonutrients, are present in relatively large amounts in fresh fruit, and their bright colors help to make them more attractive to the consumer. Moreover, anthocyanins have been found to have strong antioxidant properties and are thought to reduce the incidence of certain cancers and heart disease (Dragsted *et al.*, 2006; Shin *et al.*, 2006; Butelli *et al.*, 2008; Williams *et al.*, 2008). Thus, the investigation of anthocyanin biosynthesis in fruit has become a very popular area of research.

The blood-flesh peach [*Prunus persica* (L.) Batsch] belongs to the Rosaceae family. Some peach

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cultivars exhibit fruits with red pigmentation of the mesocarp around the stone that generally appears during the later stages of fruit ripening (Hsia et al., 1965; van Blaricom and Senn, 1967; Shen et al., 2013; Zhou et al., 2013). The Canadian peach cultivar 'Harrow Blood' was the first to exhibit a fully blood-flesh peach, producing fruits with purple pigmentation of the mesocarp and skin, associated with a red midrib color on the abaxial side of the leaf (Chaparro et al., 1995). Anthocyanin accumulation was found to be initiated early, at about 50 d after full bloom (DAFB) (Shen et al., 2013). A similar profile for anthocyanin accumulation was observed in the skin and flesh of 'Stark Red Gold' nectarines (Ravaglia et al., 2013). Taken together, these cultivars provide interesting models for dissecting the mechanisms involved in anthocyanin accumulation.

Plant anthocyanin biosynthetic genes have been identified or cloned in many plant species, e.g., apples (Takos et al., 2006; Chagné et al., 2007; Espley et al., 2007), pear (Zhang et al., 2011; Yu et al., 2012), and Chinese bayberry (Niu et al., 2010). Of the genes discovered, phenylalanine ammonia-lyase (PAL), chalcone synthase (CHS), chalcone isomerase (CHI), flavanone 3-hydroxylase (F3H), dihydroflavonol 4-reductase (DFR), anthocyanin synthase (ANS), and UDP-glucose-flavonoid 3-O-glucosyltransferase (UFGT) have all been found to participate in the biosynthesis of anthocyanin. In addition, transcription factors (TFs) that regulate anthocyanin biosynthesis have been identified and characterized in several plant species (Allan et al., 2008). For instance, the TFs of anthocyanin biosynthesis in Arabidopsis were found to contain an R2R3-MYB domain, a basic helix-loop-helix (bHLH) domain, and WD40 repeats (WDRs) (Morita et al., 2006) that work together to control the expression of the gene involved in the catalysis of the biosynthesis of anthocyanin (Baudry et al., 2004; 2006; Lepiniec et al., 2006). Similarly, the induction of anthocyanin biosynthesis in transient assays using MYB10 is dependent on the co-expression of the bHLH3 domain in peach (Ravaglia et al., 2013) and apple (Espley et al., 2007).

In this paper, we report on a study of the molecular mechanisms of anthocyanin synthesis examining seven enzyme-encoding genes and two TFs involved in the anthocyanin biosynthetic pathway of two blood-flesh peach cultivars during fruit development.

2 Materials and methods

2.1 Plant materials

We collected fruit of two blood-flesh peach cultivars, 'Beijingyixianhong' and 'Heiyoutao', and one white-fleshed peach, 'Xuebaitao'. All peaches were collected during the growing season (from March to August, 2013) from an orchard located in the germplasm resource center of peach (Nanjing, China). The sampling times of 'Beijingyixianhong' were 50 (S1), 64 (S2), 72 (S3), 80 (S4), 82 (S5), and 87 (S6) DAFB, while the sampling times of both 'Heiyoutao' and 'Xuebaitao' were 60 (S1), 74 (S2), 88 (S3), 97 (S4), 109 (S5), and 123 (S6) DAFB. For fruit bagging experiments, similarly-sized trees were randomly divided into three groups (replicates), with three trees in each group. The fruit in each group was covered with double layer (yellow outside and black inside) paper bags (Yunong (Qingdao) Co., Ltd., China) at 55 DAFB. Bags were removed at 72 DAFB for 'Beijingyixianhong' and 88 DAFB for 'Heiyoutao' and 'Xuebaitao', and the fruit was harvested at 0, 3, 5, 10, and 15 d after bag removal (DABR). Fruit skin (about 1 mm of the cortical tissue) and flesh samples (divided into two parts) were collected separately from at least 12 replicate fruit. In a separate experiment, fruit skin (FS), young leaves (YL), petals (P), and young stems (YS) were sampled at 60 DAFB to test the tissue-specific expression of the genes. For all sampling times, tissues were immediately frozen in liquid nitrogen and stored at -80 °C for further use.

2.2 Extraction and measurement of anthocyanin

Frozen fruit (3 g) was homogenized with 7.0 ml of a solvent containing methyl alcohol/water/formic acid/trifluoroacetic acid (70:27:2:1, v/v) for 24 h in the dark. The separations were centrifuged (12000 r/min, 20 min, 4 °C), filtered (0.22 μ m), and analyzed via an Agilent 1100 series high performance liquid chromatography (HPLC) system (Agilent, USA). Samples (10.0 μ l of extract) were further analyzed using a ZORBAX SB-C₁₈ column (4.6 mm×250 mm, 5 μ m) coupled with an ultraviolet (UV) detector at 515 nm with a flow rate of 0.8 ml/min at 35 °C. The mobile phase was a linear gradient of 10% aqueous formic acid (solvent A) and formic acid/acetonitrile (85:15, v/v; solvent B). Conditions: initial 95% A, 5% B; in 30 min to 88% A, 12% B; in 40 min to 75% A, 25% B;

in 50 min to 70% A, 30% B; in 60 min to 95% A, 5% B. Finally, the compounds were quantified by comparing the peak areas and presented as mg of cyanidin-3-glucoside and cyanidin-3-rutinoside (Sigma-Aldrich, USA) per 100 g fresh tissue.

2.3 RNA extraction and quantitative reverse transcription polymerase chain reaction (qRT-PCR)

Total RNA was extracted from each sample using the EASY spin plant RNA extract kit (Biomed, Beijing, China). RNA was digested with DNase I and subsequently reverse-transcribed using a Supermo III RT kit (BioTeKe, Beijing, China). Samples of cDNA were diluted to 1:10 with dH₂O for use in qRT-PCR. All gene-specific primers were designed using Primer3 (http://frodo.wi.mit.edu/cgibin/primer3/primer3 www. cgi), according to the available databases (http://www. ncbi.nlm.nih.gov/Genbank/index.html) and published papers (Espley et al., 2007; Lin-Wang et al., 2010; Zhou et al., 2013) (Table 1). The specificity of each primer pair was analyzed by a melting curve at the end of the PCR run, with one single peak and no dimer formation. The specificity of these primers was confirmed by cloning and sequencing of qRT-PCR products. We conducted qRT-PCR using an ABI 7500 System (Applied Biosystems, Foster, CA, USA) with the SYBR Premix Ex Taq[™] (TaKaRa, Dalian, China). The qRT-PCR reaction solution consisted of 10 µl of SYBR Green PCR Master Mix, 0.4 µl forward primer (10 μ mol/L), 0.4 μ l reverse primer (10 μ mol/L), 0.4 µl ROX Reference dye (50×; all from TaKaRa), 6.8 µl dH₂O, and 2.0 µl template cDNA in a total volume of 20 µl. The gRT-PCR amplification procedure was initiated with a preliminary step of 95 °C

for 30 s, followed by 40 cycles of 95 °C for 5 s and 60 °C for 34 s. All qRT-PCR reactions were normalized using the housekeeping genes *UBQ10* and *TEF2* (Tong *et al.*, 2009). Relative gene expression levels were calculated using the $2^{-\Delta\Delta C_T}$ method (Livak and Schmittgen, 2001) with three measurements for each biological replicate sample.

2.4 Statistical analysis

All statistical analyses were performed using the Data Processing System (DPS, Version 14.10; Zhejiang University, Hangzhou, China). All results were summarized as mean±standard deviation (SD); least significant differences (LSDs; α =0.05) were considered statistically significant. The graphs were drawn with Origin software version 8.1 (Microcal Software Inc., Northampton, MA, USA).

3 Results

3.1 Anthocyanin content in blood-flesh peach during fruit development

The total anthocyanin content, comprising cyanidin-3-glucoside and cyanidin-3-rutinoside, varied greatly among different tissues of 'Heiyoutao' and 'Beijingyixianhong' during fruit development (Figs. 1 and 2). The anthocyanin content of 'Heiyoutao' increased significantly from 60 to 88 DAFB and declined thereafter. Anthocyanin was not detected until 64 DAFB in 'Beijingyixianhong' and then increased continuously from 64 to 87 DAFB before reaching its maximum. The highest content of anthocyanin was found in the fruit skin, followed by the outer layer of

Fable 1 Primers used for RT-PCR analy	vsi	S
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Gene	GenBank ID	Forward primer (5' to 3')	Reverse primer (5' to 3')	Product size (bp)
PpPAL	HM543574.1	AAGCTGCTGAAAAGGTGCAT	TCATTTTGGTTGCTGCTCTG	126
PpCHS	HM543568.1	CCGTGAAGTTGGGCTTACAT	TGGGTGTGCAATCCAGAATA	139
РрСНІ	HM543569.1	AATATGGGGTGCAGCTTGAG	TGCTGGAAAATGAAACGTGA	155
PpF3H	HM543570.1	CAAAACCCAGCTCAAGAAGC	CTCCGAGTCCTGCAATTGTT	174
PpDFR	HM543571.1	TGAATGTCGAAGAGCACCAG	AAATGGGCCAATCACAAGAG	197
PpUFGT	JX149550.1	TGGCCTCTCAAGGACAGTTT	CCGCCTGCTATAGTCTCCAG	173
PpANS	HM543572.1	AGGAGTTGAAGAAGGCAGCA	GCTCAATGGGAAGATCGAAA	123
PpMYB10	EU155160.1	GGCAGGACTGCGAATAATGT	CCCACCAATCACGTTGAGTA	196
PpTT8	KF585000.1	CCGACCACCCACATTTCC	TGACCGTTGCGGCCCGTCTC	163
TEF2	TC3544	GGTGTGACGATGAAGAGTGATG	TGAAGGAGAGGGAAGGTGAAAG	129
UBQ10	TC2782	AAGGCTAAGATCCAAGACAAAGAG	CCACGAAGACGAAGCACTAAG	146



Fig. 1 Changes of color in the mesocarp of three peach cultivars during fruit development (a) Beijingyixianhong; (b) Heiyoutao; (c) Xuebaitao. Samples were assayed at 50 (S1), 64 (S2), 72 (S3), 80 (S4), 82 (S5), and 87 (S6) d after full bloom (DAFB) from 'Beijingyixianhong', and at 60 (S1), 74 (S2), 88 (S3), 97 (S4), 109 (S5), and 123 (S6) DAFB from both 'Heiyoutao' and 'Xuebaitao'



Fig. 2 Anthocyanin concentration in the fruit of 'Heiyoutao' (a, c, e) and 'Beijingyixianhong' (b, d, f) during fruit development, as measured by HPLC

(a, b) Fruit skin; (c, d) Outer layer of mesocarp; (e, f) Inner layer of mesocarp. Fruit skin and flesh samples were collected separately from at least 12 replicate fruit of each cultivar. The vertical bars represent the standard error (SE) of three replicates

the mesocarp. We found that cyanidin-3-glucoside was the major anthocyanin in all layers of the fruit. Cyanidin-3-rutinoside was not detected in the mesocarp of 'Beijingyixianhong' and only small amounts were found in the fruit skin. Interestingly, the leaf midrib of 'Heiyoutao' contained both cyanidin-3glucoside and cyanidin-3-rutinoside, as did its fruit (Fig. S1). 'Xuebaitao', which produces fruits with a red color surrounding the stone during the mature stage, expressed only cyanidin-3-glucoside (Fig. S2).

3.2 Expression of anthocyanin biosynthetic genes in two blood-flesh peach cultivars

Nine genes were differentially expressed in the peach cultivars (Fig. 3). Firstly, the transcription of *PpPAL*, *PpUFGT*, and *PpMYB10* reached peak levels during the initial stages of fruit development at S2 or S3. During this time, the expression levels of *PpUFGT* and *PpMYB10* were higher in 'Beijingyixianhong' than in 'Heiyoutao'. Post-peak levels, expression of these genes continuously declined until maturity.



Secondly, the transcription of PpCHI, PpF3H, PpDFR, PpANS, and PpTT8 reached peak levels during the initial stages of fruit development at S2 or S3 in 'Beijingyixianhong' and 'Xuebaitao'. However, the overall expression of these genes in 'Heiyoutao' was very low and gradually declined throughout development. Finally, the expression of the gene PpCHS was closely related to the anthocyanin content in 'Beijingyixianhong', but had a tendency to continuously decline to maturity in 'Heiyoutao'. The expression of the gene PpCHS reached its peak level

Fig. 3 Expression analysis of anthocyanin biosynthetic genes in the mesocarp of 'Heiyoutao', 'Beijingyixianhong', and 'Xuebaitao' during fruit development

Expression levels of the *UBQ10* and *TEF2* genes were used to normalize the expression of all other genes under identical conditions. The vertical bars represent the standard error (SE) of three replicates

at S2 in the white-fleshed cultivar 'Xuebaitao'. This expression pattern was similar to that of 'Heiyoutao'.

3.3 Expression of anthocyanin biosynthetic genes after the bag removal

Following bag removal, anthocyanin accumulated in the mesocarp of the blood-flesh peach (Fig. 4) in a similar fashion to the natural state (as described in Fig. 2). Small amounts of anthocyanins were detected in the mesocarp of the 'Xuebaitao' fruit at 15 DABR. The transcription levels of seven genes (PpPAL, PpCHI, PpF3H, PpDFR, PpMYB10, PpANS, and PpTT8) were markedly up-regulated in both bloodflesh peaches at 3 or 5 DABR, followed by a sharp drop (Fig. 5). The transcription levels of PpPAL and PpTT8 were lower in the white-flesh of 'Xuebaitao', suggesting that these genes were non-sensitive to the bag removal treatment. Moreover, the transcription levels of PpCHS and PpUFGT still had relatively high expression in 'Beijingyixianhong' from 3 to 15 DABR, suggesting a common transcription mechanism that uses multiple pathways all contributing to the biosynthesis of anthocyanin.



Fig. 4 Anthocyanin concentrations following bag removal in the mesocarp

Samples were assayed at 0, 3, 5, 10, and 15 d after bag removal (DABR). The vertical bars represent the standard error (SE) of three replicates





Expression levels of the *UBQ10* and *TEF2* genes were used to normalize the expression of all other genes under identical conditions. The vertical bars represent the standard error (SE) of three replicates

3.4 Tissue-specific expression analysis

Anthocyanin content varied greatly among the different organs and tissues of 'Heiyoutao', with the highest levels found in the petals and the lowest in the young stems (Fig. 6). The anthocyanin content in the petals was significantly higher in 'Heiyoutao' than in both 'Beijingyixianhong' and 'Xuebaitao'. These results are in accordance with visual observations (Fig. S3). Tissue-specific expression analysis revealed that the genes were differentially expressed in various organs and tissues (Fig. 7). Firstly, *PpCHS*, *PpUFGT*, and *PpTT8* were all highly expressed in the fruit skin of 'Heiyoutao' and 'Beijingyixianhong'.



Fig. 6 Anthocyanin concentrations in various organs and tissues of the peach

P: petal; FS: fruit skin; YL: young leaf; YS: young stem. The vertical bars represent the standard error (SE) of three replicates



Fig. 7 Expression analysis of the anthocyanin biosynthetic genes in various organs and tissues The expression levels of the *UBQ10* and *TEF2* genes were used to normalize the expression of all other genes under identical conditions. P: petal; FS: fruit skin; YL: young leaf; YS: young stem. The vertical bars represent the standard error (SE) of three replicates

The transcription levels of the *PpUFGT* and *PpTT8* genes were lower in 'Xuebaitao'. Secondly, *PpPAL*, *PpCHI*, *PpF3H*, *PpDFR*, and *PpANS* were all expressed at the highest level in the fruit skin and the young stems of 'Heiyoutao' and 'Beijingyixianhong'. These genes were also expressed at the highest level in the leaves of 'Xuebaitao', suggesting that the expression of the flavonoid "early" biosynthetic genes (EBGs) (including *PpPAL*, *PpF3H*, and *PpDFR*) is less associated with anthocyanin levels. Finally, *PpMYB10* was expressed at the highest level in the petals and was highly correlated with anthocyanin levels in different tissues.

4 Discussion

Anthocyanin concentrations have previously been measured by ultraviolet spectrometry in the peach (Leng et al., 2000; Zhou et al., 2013). However, these studies showed the difficulty in distinguishing different anthocyanin components, as their concentrations across different organs and/or tissues were not identifiable. In the current study, we identified that cyaniding-3-glucoside is the major anthocyanin in the leaves, petals, and stems of the peach, consistent with previous studies (Cheng and Crisosto, 1997; Cevallos-Casals et al., 2006; Orazem et al., 2011). On the other hand, cyaniding-3-rutinoside was found in the fruit skin of 'Heiyoutao' and was nondetectable in the mesocarp of 'Beijingyixianhong', suggesting that production of this anthocyanin component differs among cultivars. Furthermore, we estimate that there are six types of anthocyanin pigments in the petals of these peach cultivars, making them much more complex than other organs or tissues (data not shown). However, the molecular mechanisms of anthocyanin biosynthesis are still not fully understood.

In this study, the expression of *PpCHS* was found to correlate with the accumulation of anthocyanin, suggesting that it is a key gene involved in the process of anthocyanin biosynthesis. These findings also echo previous research (Tsuda *et al.*, 2004), and thus this pattern may be common to other red fruit of the Rosaceae family. We found that the transcription levels of *PpPAL* and *PpUFGT* were much lower in the white-flesh cultivar 'Xuebaitao' than in the blood-flesh peaches. Taken together, this pattern of mRNA expression suggests that *PpUFGT* is under a different regulatory regime, compared with the other flavonoid pathway genes in the peach, and that anthocyanin synthesis is controlled at a later stage than seen in previously studies (Boss et al., 1996; Kobayashi et al., 2001). In addition, these results suggest that anthocyanin biosynthesis is not active in the mesocarp of the white-flesh peach due to the absence of *PpPAL*. In contrast, the early-expressing genes of the flavonoid pathway (PpCHS, PpF3H, and *PpDFR*) were all expressed in the white-flesh peach 'Xuebaitao' and are therefore likely to be involved in the synthesis of other secondary metabolites, such as free-auxins (Schwalm et al., 2003; Shen et al., 2013). Similar results have been found in grape (Boss et al., 1996), apple (Kim et al., 2003), Chinese bayberry (Niu et al., 2010), kiwifruit (Montefiori et al., 2011), and litchi (Wei et al., 2011). Furthermore, Schwalm et al. (2003) reported that CHS-dependent flavonoid aglycones have the potential to be endogenous regulators of the basipetal auxin flux, thereby leading to free-auxin accumulation. 'Heiyoutao' has the genotype bf, and the blood-flesh phenotype has been associated with reduced tree height in progeny derived from this cultivar (Shen et al., 2013). This suggests that the phenotype may be closely related to auxin flux, but the detailed mechanisms remain to be elucidated.

Results show that bagging has a significant effect on anthocyanin synthesis (Dussi et al., 1995). As expected, anthocyanin concentrations were generally higher in the natural state than following bagging. After bag removal, the transcription levels of most of the anthocyanin biosynthetic genes were markedly enhanced at 3 DABR, except for *PpPAL* and *PpTT8*. These results indicate that a single gene is not responsible for anthocyanin accumulation and that the coordinated action of many genes is likely involved in anthocyanin biosynthesis. This hypothesis is supported by previous findings in apple (Honda et al., 2002; Takos et al., 2006), pear (Yu et al., 2012), and grape (Walker et al., 2007). In particular, PpUFGT was clearly expressed in the white-flesh peach 'Xuebaitao', but anthocyanin accumulation was not detected (Figs. 4 and 5). These findings suggest that *PpUFGT* is likely more highly regulated than the other flavonoid genes. Furthermore, Ravaglia *et al.* (2013) reported that *PpMYB10* positively regulates the promoters of *PpUFGT* and *PpDFR*. Our qPCR results show that the expression pattern of the *PpMYB10* gene is highly correlated with *PpUFGT* and *PpDFR* in 'Heiyoutao', but not in 'Beijingyixianhong'. These results suggest that *PpMYB10* plays a different role in the anthocyanin regulation of the two blood-flesh peaches. Moreover, some members of *MYB10* are thought to be involved in regulating anthocyanin biosynthesis (Rahim *et al.*, 2014; Uematsu *et al.*, 2014). Further studies should focus on their responsiveness to both developmental and environmental stimuli in blood-flesh peaches.

Finally, in the two blood-flesh peach cultivars that we investigated, the petals were pink, owing to a greater accumulation of anthocyanins compared with other tissues (Fig. S3). Tissue-specific expression analysis revealed that *PpMYB10* expression seemed more pronounced in petals than in other tissues (Fig. 7), strongly indicating that *PpMYB10* is a key regulatory gene involved in anthocyanin biosynthesis within the petals of these peach cultivars. In contrast, the other anthocyanin biosynthesis genes (PpPAL, PpCHI, pPpF3H, PpDFR, and PpANS) were all highly expressed in the stems and leaves of the white-flesh peach 'Xuebaitao', even though anthocyanin was not detected. This result indicates that these genes are likely involved in processes other than anthocyanin production.

In conclusion, this study demonstrated large differences in anthocyanin accumulation among the two blood-flesh peaches studied. The transcription level of *PpCHS* was closely related to anthocyanin accumulation, due to its diverse roles in the biosynthetic pathway. Further studies are required to elucidate the molecular mechanisms of action of *PpCHS* in the anthocyanin biosynthesis pathway.

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Compliance with ethics guidelines

Yun JIAO, Rui-juan MA, Zhi-jun SHEN, Juan YAN, and Ming-liang YU declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by any of the authors.

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List of electronic supplementary materials

- Fig. S1 HPLC chromatogram of the anthocyanin components in the leaf midrib of 'Heiyoutao'
- Fig. S2 HPLC chromatogram of the cultivar 'Xuebaitao' from the flesh surrounding the stone at maturity
- Fig. S3 Petals during full bloom of 'Beijingyixianhong' (a), 'Heiyoutao' (b), and 'Xuebaitao' (c)

本文题目:	两个红肉桃品种果实发育期花色素苷合成与分子调控研究
	Gene regulation of anthocyanin biosynthesis in two blood-flesh peach (<i>Prunus persica</i> (L.
研究目的:	深入分析两个红肉桃品种果实发育期花色素苷合成差异的分子机理,为红肉桃品种的遗传改
	良与利用提供理论依据。
创新要点:	阐明两个红肉桃品种果实发育期花色素苷生物的合成与相关基因的时空表达规律,探讨造成
	两个品种花色素苷积累差异的内在分子机制。
研究方法:	应用高效液相色谱(HPLC)和逆转录聚合酶链式反应(RT-PCR)分别鉴定两个红肉桃品和
	花色素苷的积累规律与合成相关基因的表达水平,进一步探讨两者花色素苷合成的内在分子调控机理。
重要结论:	两个红肉桃品种果实发育期花色素苷合成与 PpCHS 表达水平密切相关,同时 PpPAL 在白肉
	桃品种果实发育期的表达水平低于红肉桃品种,推测这两种基因可能是限制白肉桃品种果实
	中花色素苷合成的因素之一。
关键词组:	红肉桃;花色素苷;套袋;基因表达