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Transcript profiling of structural genes involved in cyanidin-based anthocyanin biosynthesis between purple and non-purple carrot (*Daucus carota* L.) cultivars reveals distinct patterns

Zhi-Sheng Xu, Ying Huang, Feng Wang, Xiong Song, Guang-Long Wang and Ai-Sheng Xiong*

**Abstract**

**Background:** Carrots (*Daucus carota* L.) are among the 10 most economically important vegetable crops grown worldwide. Purple carrot cultivars accumulate rich cyanidin-based anthocyanins in a light-independent manner in their taproots whereas other carrot color types do not. Anthocyanins are important secondary metabolites in plants, protecting them from damage caused by strong light, heavy metals, and pathogens. Furthermore, they are important nutrients for human health. Molecular mechanisms underlying anthocyanin accumulation in purple carrot cultivars and loss of anthocyanin production in non-purple carrot cultivars remain unknown.

**Results:** The taproots of the three purple carrot cultivars were rich in anthocyanin, and levels increased during development. Conversely, the six non-purple carrot cultivars failed to accumulate anthocyanins in the underground part of taproots. Six novel structural genes, *CA4H1*, *CA4H2*, *4CL1*, *4CL2*, *CHI1*, and *F3′H1*, were isolated from purple carrots. The expression profiles of these genes, together with other structural genes known to be involved in anthocyanin biosynthesis, were analyzed in three purple and six non-purple carrot cultivars at the 60-day-old stage. *PAL3/PAL4*, *CA4H1*, and *4CL1* expression levels were higher in purple than in non-purple carrot cultivars. Expression of *CHS1*, *CHI1*, *F3H1*, *F3′H1*, *DFR1*, and *LDOX1/LDOX2* was highly correlated with the presence of anthocyanin as these genes were highly expressed in purple carrot taproots but not or scarcely expressed in non-purple carrot taproots.

**Conclusions:** This study isolated six novel structural genes involved in anthocyanin biosynthesis in carrots. Among the 13 analyzed structural genes, *PAL3/PAL4*, *CA4H1*, *4CL1*, *CHS1*, *CHI1*, *F3H1*, *F3′H1*, *DFR1*, and *LDOX1/LDOX2* may participate in anthocyanin biosynthesis in the taproots of purple carrot cultivars. *CHS1*, *CHI1*, *F3H1*, *F3′H1*, *DFR1*, and *LDOX1/LDOX2* may lead to loss of light-independent anthocyanin production in orange and yellow carrots. These results suggest that numerous structural genes are involved in anthocyanin production in the taproots of purple carrot cultivars and in the loss of anthocyanin production in non-purple carrots. Unexpressed or scarcely expressed genes in the taproots of non-purple carrot cultivars may be caused by the inactivation of regulator genes. Our results provide new insights into anthocyanin biosynthesis at the molecular level in carrots and for other root vegetables.

**Keywords:** Anthocyanin pathway, Root development, Purple carrot, Cyanidin, *Daucus carota* L, Gene expression
Background

Anthocyanins are widely distributed water-soluble pigments belonging to the flavanoid group of phytochemicals. Over 635 types of anthocyanins have been identified [1]. These mainly possess six common aglycones (cyanidin, pelargonidin, delphinidin, peonidin, petunidin, and malvidin) and various types of glycosylated and acylated compounds [2]. Anthocyanins protect plants from strong light, heavy metals, and pathogens, and play an important role in flowers [3,4]. As they have low toxicity and vary in color they are often used as healthier alternatives to synthetic colorants [5]. Previous studies have confirmed that anthocyanins provide antioxidants for human health that protect against a broad range of diseases, including a high blood cholesterol level, cardiovascular disease, and ultraviolet radiation damage [2,6-8].

Carrots (Daucus carota L.) are among the 10 most economically important vegetable crops grown worldwide [9]. Carrot cultivars appear in five taproot color types: purple, orange, yellow, red, and white. Although orange carrot cultivars (D. carota ssp. sativus var. sativus) account for the majority of production, purple carrots (D. carota ssp. sativus var. atrorubens Alef.) are enjoying increased popularity, largely because they contain high amounts of anthocyanin in their flesh taproots. Purple carrot cultivars have existed for over 3000 years, and are much older than orange carrot cultivars [10]. Anthocyanins from purple carrots are commonly used as natural food colorants in candies, ice cream, and beverages, this is because they remain stable when exposed to heat and light, and have increased pH values [11,12]. Purple carrots mainly contain cyanidin-based anthocyanins; some cultivars also contain trace amounts of peonidin- or pelargonidin-based anthocyanins in their taproots [13].

The anthocyanin biosynthesis path has been extensively studied in numerous plant species, including bilberry (Vaccinium myrtillus L.), grape (Vitis vinifera L.), apple (Malus × domestica), Arabidopsis (Arabidopsis thaliana), Mitchell petunia [Petunia axillaris × (Petunia axillaris × Petunia hybrida cv. 'Rose of Heaven')], and sweet potato (Ipomoea batatas L. Lam.) [14-20]. Two classes of genes participate in the anthocyanin biosynthesis pathway: structural genes and regulatory genes. Structural genes encode enzymes that directly catalyze reaction steps leading to the formation of anthocyanins; the transcription of these genes is controlled by regulatory genes, such as MYB, bHLH, and WD40 genes [16,17,21]. Structural genes that participate in anthocyanin biosynthesis have been identified in numerous plant species [14-18,20-23]. Some functional genes participating in this pathway have also been identified in carrots; these include the phenylalanine ammonia-lyase (PAL), chalcone synthase (CHS), flavanone 3-hydroxylase (F3H), dihydroflavonol 4-reductase (DFR), and leucoanthocyanidin dioxygenase (LDOX) genes [24]. We described the presence of the UDP-galactose: cyanidin 3-O-galactosyltransferase (UCGT) in the purple carrot cultivars in our previous study [25]. However, cinnamate 4-hydroxylase (CA4H), 4-coumaroyl-coenzyme A ligase (4CL), chalcone-flavonone isomerase (CHI), and flavonoid 3′-hydroxylase (F3′H) genes have not been identified in carrots.

To obtain insights into differences in anthocyanin biosynthesis between purple and non-purple carrot cultivars, we cloned six novel structural genes involved in anthocyanin biosyntheses from taproot-derived cDNA. The expression patterns of 13 structural genes in the taproots of three purple and six non-purple carrot cultivars were analyzed at the transcriptional level. The accumulation of anthocyanins was determined in parallel. The aim of this work was to determine the stage at which the anthocyanin biosynthesis pathway switches off, thus leading to loss of anthocyanins in non-purple carrot cultivars.

Results

Taproot color of nine carrot cultivars at different development stages

Anthocyanins accumulate in different parts of carrot taproots at different development stages; this leads to taproots displaying a purple or dark color. In 60-day-old carrots, anthocyanins accumulated in the cortex and xylem of ‘Deep purple’ and ‘Purple 68’ cultivar taproots, but only in the cortex of ‘Tianzi2hao’ (Figure 1). In the six other carrot cultivars, no purple or dark coloring was detected in the taproot. In 90- and 120-day-old carrots, anthocyanins accumulated in the cortex, phloem, and xylem of ‘Deep purple’, ‘Purple 68’, and ‘Tianzi2hao’ taproots. In the six other carrot cultivars, no purple or dark coloring was detected in these taproot parts, with the exception of the hypocotyl-derived root part of 90- and 120-day-old ‘Kuroda’, 90-day-old ‘Sanhongliucun’, and 120-day-old ‘Junchuanhong’; these displayed dark color in the epidermis when exposed to light (Figure 2). As hypocotyl-derived parts of the taproots of 120-day-old ‘Sanhongliucun’ and 90-day-old ‘Junchuanhong’ cultivars were not exposed to light, purple or dark coloring was not detectable in their epidermis.

Anthocyanin content in the taproots of nine carrot cultivars at different developmental stages

Total anthocyanin content in the taproots of the three purple carrot cultivars (‘Deep purple’, ‘Purple 68’, and ‘Tianzi2hao’) increased significantly during development (Figure 3). These cultivars accumulated anthocyanins more efficiently in their taproots between the 60- and 90-day-old stages than between the 90- and 120-day-old stages. Of these three purple carrot cultivars, ‘Purple 68’ showed the highest anthocyanin content in
the taproots at all three stages. In the taproots of the three orange carrot cultivars (‘Kuroda’, ‘Sanhongliucun’, and ‘Junchuanhong’), anthocyanin was not detected at the 60-day-old stage. At the 90-day-old stage, the anthocyanin content in ‘Kuroda’ and ‘Sanhongliucun’ taproots was 6.47 mg/100 g fresh weight (fw) and 1.35 mg/100 g fw, respectively; anthocyanin was not detected in ‘Junchuanhong’ taproots at this stage. The taproots of 120-day-old ‘Kuroda’, ‘Sanhongliucun’, and ‘Junchuanhong’ contained 0.56, 0.21, and 0.23 mg/100 g fw anthocyanins, respectively. Anthocyanins did not accumulate in the taproots of the three yellow carrot cultivars in any of the three stages.

Analysis of carrot structural genes for cyanidin-based anthocyanin biosynthesis

As cyanidin-based anthocyanins represent almost all anthocyanin content in carrots, we analyzed structural genes for cyanidin-based anthocyanin biosynthesis. We propose the following cyanidin-based anthocyanin biosynthesis pathway in purple carrots (Figure 4). PAL, CA4H, and 4CL code enzymes implicated in the general phenylpropanoid pathway of anthocyanin biosynthesis in carrots. CHS, CHI, F3H, F3’H, DFR, LDOX, and UCGT code enzymes involved in the anthocyanin pathway of anthocyanin biosynthesis in carrots. The full names of these genes and their corresponding accession numbers in GenBank and CarrotDB are listed in Table 1 [25].

PAL1 (GenBank ID:D85850.1), PAL3 (GenBank ID: AB089813.1), PAL4 (GenBank ID:AB435640.1), CHS1 (GenBank ID:AJ006779.1), CHS2 (GenBank ID: D16255.1), CHS9 (GenBank ID: D16256.1), F3H1 (GenBank ID: AF184270.1), DFR1 (GenBank ID:AF184271.1), LDOX1 (GenBank ID:AF184273.1), and LDOX2 genes (GenBank ID:AF184274.1) were present in GenBank. The CA4H1, CA4H2, 4CL1, 4CL2, CHI1, and F3’H1 genes were identified in CarrotDB using a BLAST-based search tool; these genes were further identified by cloning and sequencing genes from the ‘Deep purple’ cultivar. The nucleotide and deduced amino acid sequences of these genes were deposited at the National Center for Biotechnology Information (NCBI). The NCBI accession numbers of CA4H1, CA4H2, 4CL1, 4CL2, CHI1, and F3’H1 are listed in Table 1.

The PAL3 and PAL4, CHS2 and CHS9, and LDOX1 and LDOX2 genes were considered as allelic genes because they share very high identity in their nucleotide acid
sequences (>95%); furthermore, only one gene corresponding to each pair of genes was found in the CarrotDB. Therefore, only one pair of primers, specific to each pair of genes, was used for quantitative real-time polymerase chain reaction (qRT-PCR).

Expression of cyanidin-based anthocyanin biosynthetic genes in the taproot at the 60-day-old stage

Purple carrot cultivars had accumulated anthocyanins in taproots by the 60-day-old stage. Nucleotide sequences of primer pairs used for qRT-PCR specific to each anthocyanin biosynthetic genes are given in Table 2. PAL3/PAL4, CA4H1, 4CL1, CHS1, CHI1, F3H1, F3’H1, DFR1, and LDOX1/LDOX2 genes showed significantly higher transcript abundance in the taproots of ‘Deep purple’, ‘Purple 68’, and ‘Tianzi2hao’ than in the taproots of ‘Kuroda’, ‘Sanhongliucun’, ‘Junchuanhong’, ‘Bejo1719’, ‘Qitouhuang’, and ‘Baiyu’ (Figure 5). Correlation analysis results revealed that CHS1, CHI1, F3H1, F3’H1, DFR1, and LDOX1/LDOX2 were highly correlated with anthocyanin presence among the genes encoding enzymes implicated in the anthocyanin pathway in anthocyanin biosynthesis (Additional file 1: Table S1). Among the cultivars, ‘Deep purple’ showed the highest taproot mRNA levels of PAL3/PAL4, CA4H1, 4CL1, CHS1, F3H1, F3’H1, and LDOX1/LDOX2. ‘Purple 68’ had the highest taproot mRNA levels of PAL1, 4CL2, and DFR1. ‘Tianzi2hao’ had the highest taproot mRNA levels of CA4H2 and CHI1, and ‘Kuroda’ showed the highest mRNA levels of CHS2/CHS9 in the taproots. Transcript levels of PAL1 and CA4H2 were lower than PAL3/PAL4 and CA4H1, respectively, in the taproots of all carrot cultivars. Transcript levels of 4CL2 in the taproots of ‘Kuroda’, ‘Junchuanhong’, ‘Bejo1719’, ‘Qitouhuang’, and ‘Baiyu’ were higher than those of 4CL1, but lower than observed in the taproots of ‘Deep purple’, ‘Purple 68’, ‘Tianzi2hao’, and ‘Sanhongliucun’. In the three purple carrot cultivars, mRNA abundance of CHS2/CHS9 in the taproot was significantly lower than that of CHS1. In the six other carrot cultivars, CHS2/CHS9 showed similar mRNA expression levels as CHS1 in the taproots (Figure 5).

Discussion

In many plants, anthocyanin biosynthesis can be light independent or light induced. The light-independent anthocyanin biosynthesis pathway has been investigated in other plant species, including apple and sweet potato [16,22]. Light-induced anthocyanin biosynthesis has been observed in several plant species, including apple, grape, and Mitchell petunia [18,20,23]. In this work, purple carrot cultivars could produce rich anthocyanins in taproots light independently. In contrast, yellow carrot cultivars failed to produce anthocyanins in the taproots, while orange carrots could only produce small amounts of anthocyanins in the epidermis.
of hypocotyl-derived root parts light dependently. To determine genes involved in light-independent anthocyanin biosynthesis in purple carrots and those responsible for losses of light-independent anthocyanin production in non-purple carrots, we analyzed anthocyanin pathway structural genes in three purple and six non-purple cultivars.

Some structural genes (PAL1, PAL3/PAL4, CHS1, CHS2/CHS9, F3H1, DFR1, and LDOX1/LDOX2) involved in anthocyanin biosynthesis have been previously cloned, and their expression profiles analyzed under ultraviolet light [24,26]. Our previous study identified that UCGT1 expressed in purple carrot cultivars [25]. In this study, six structural genes (CA4H1, CA4H2, 4CL1, 4CL2, CHI1, and F3′H1) present in purple ('Deep purple') and non-purple ('Kuroda') carrot cultivars were cloned and sequenced. PAL3/PAL4, CA4H1, 4CL1, CHS1, F3H1, F3′H1, DFR1, and LDOX1/LDOX2 genes may be involved in the anthocyanin biosynthesis in purple carrots as these genes showed higher expression levels in purple carrots than non-purple carrots. CHS1, CHI1, F3H1, F3′H1, DFR1, and LDOX1/LDOX2 genes were strongly correlated with the presence of anthocyanins, as indicated by high gene expression levels in the taproots of purple carrot cultivars but no or scarce expression in the taproots of non-purple carrot cultivars. This suggests that these genes predominantly lead to loss of anthocyanin production in non-purple carrot cultivars. A similar result was observed in sweet potatoes [22]. Loss of anthocyanins in the taproots of non-purple carrots is possibly caused by inactivation of regulator genes such as MYB, bHLH, and WD40 genes. Future investigation will focus on transcription factors controlling expression of hypocotyl-derived root parts light dependently.

![Figure 4 Schematic of the proposed cyanidin-based anthocyanin biosynthetic pathway.](image)

| Gene family | Annotation | GenBank ID | Scaffolds ID in CarrotDB | Predict genes ID in CarrotDB |
|-------------|------------|------------|--------------------------|-----------------------------|
| PAL         | PAL1       | D85850.1   | scaffold_229, 168181     | g443,75570                  |
|             | PAL3/PAL4  | AB089813.1 | scaffold_16648, 16649, 169243 | g17456, 17457, 76158        |
| CA4H        | CA4H1      | KM359961   | scaffold_25749           | g24733                      |
|             | CA4H2      | KM359962   | scaffold_45729           | g36870                      |
| 4CL         | 4CL1       | KM359963   | scaffold_1380            | g1988                       |
|             | 4CL2       | KM359966   | scaffold_2619            | g3566                       |
| CHS         | CHS1       | AJ006779.1 | scaffold_27333, 21408    | g25908, 21339               |
|             | CHS2/CHS9  | D16255.1/D16256.1 | scaffold_139899, 4932 | g61298, 6227               |
| CHI         | CHI1       | KM359964   | scaffold_8876            | g10389                      |
| F3H         | F3H1       | AF184270.1 | scaffold_17226           | g17950                      |
|             | F3H1       | KM359965   | scaffold_36531           | g31974                      |
| DFR         | DFR1       | AF184271.1 | scaffold_1955, 1956      | g31520, 2745                |
| LDOX        | LDOX1/LDOX2| AF184273.1 | scaffold_20348, 20349, 20350 | g20467, 78025, 20472       |

Table 1 Cyanidin-based anthocyanin biosynthetic genes annotation and accession numbers in GenBank or CarrotDB

Gene abbreviations: PAL, Phenylalanine ammonia-lyase; CA4H, Cinnamate 4-hydroxylase; 4CL, 4-coumaroyl-coenzyme A ligase; CHS, Chalcone synthase; CHI, Chalcone-flavonone isomerase; F3H, Flavanone 3-hydroxylase; F3′H, Flavanoid 3′-hydroxylase; DFR, Dihydroflavonol 4-reductase; LDOX, Leucoanthocyanidin dioxygenase.
these structural genes to identify the key gene(s) involved in anthocyanin production in purple carrot cultivars and responsible for anthocyanin loss in non-purple carrot cultivars.

The taproots of purple carrots are rich in anthocyanins, reaching a maximum of 175 mg/100 g fw in some cultivars [27]. In this study, anthocyanin content varied significantly in the taproots of the three purple carrot cultivars at the three different stages, as visually indicated by the degree of root coloring. Anthocyanin content in 60-day-old stage taproots of the three purple carrot cultivars was comparable to previously reported for several genotypes of purple carrots [13]. In 90- and 120-day-old purple carrots, anthocyanin content was higher than previously reported, this may be because of the different growth conditions and harvest time of the carrots [13]; anthocyanin accumulation in carrots is sensitive to variations in growth conditions, such as temperature, light, and nutrients [28-30]. Anthocyanin contents in taproots of the three purple carrot cultivars at the 120-day-old stage were higher than those in strawberries, red onion, and red grapes but lower than that observed in blueberries [31]. Anthocyanins accumulated in the epidermis of the hypocotyl-derived root part of the three orange carrot cultivars after they were exposed to light. This suggested that a light-induced anthocyanin biosynthesis pathway is found in these orange carrot cultivars.

**Conclusions**

Purple carrot cultivars produced rich amounts of anthocyanins in the taproots light independently, whereas non-purple cultivars did not. The anthocyanin content in purple carrot cultivars increased as root growth occurred. Six novel candidate structural genes that existed in both purple and non-purple carrots were successfully cloned and sequenced. *PAL3/PAL4*, *CA4H1*, *4CL1*, *CHS1*, *F3H1*, *DFR1*, and *LDOX1/LDOX2* may participate in anthocyanin biosynthesis in the taproots of purple carrot cultivars. *CHS1*, *CHI1*, *F3H1*, *F3′H1*, *DFR1*, and *LDOX1/LDOX2* were unexpressed or scarcely expressed in non-purple carrots, thus may lead to the loss of light-independent anthocyanins production in orange and yellow carrots. Our results provide new insights into anthocyanin biosynthesis in carrots at the molecular level and are of importance for other root vegetables.

**Methods**

**Plant materials and growth conditions**

Three purple carrot cultivars (‘Deep purple’, ‘Purple 68’, and ‘Tianzi2hao’), three orange carrot cultivars (‘Kuroda’, ‘Sanhongliucun’, and ‘Junchuanhong’), and three yellow carrot cultivars (‘Bejo1719’, ‘Qitouhuang’, and ‘Baiyu’) were chosen for this work (Figure 2). Seeds were sown in pots containing a soil/vermiculite mixture (1:1) in a controlled artificial climatic chamber, with a photoperiod of 12 h light (2000–3000 lux) and 12 h dark at day/night temperatures of 22°C/18°C. Carrot plants were grown under the same conditions. Taproots of carrots at 60-, 90-, and 120-day-old stages were harvested, immediately frozen in liquid nitrogen, and stored at −70°C for future analysis. Three taproots were sampled for each carrot cultivar at each stage.

| Gene      | Forward primer 5′-3′ | Reverse primer 5′-3′ |
|-----------|----------------------|----------------------|
| PAL1      | ACATTACCCCTTGCTTGCCACT | TCAAAAGATTCACCACTCAACCTC |
| PAL3/PAL4 | CTGGCATGGCCTCTATGGTACT | TGGCTCTGGATGTCCTCAACCTC |
| CA4H1     | GTGGAGGCCAACGGAAATG   | CGTCCGATTGATGCACGGAAAT |
| CA4H2     | CCGGCAAAGAGCAAAAGTTGT | CAGCCCCGAATGTAGGAAAT |
| 4CL1      | AAACACCTGGCGTACTCTGCG | CCGAAAGCAAGATCATCAGGTATC |
| 4CL2      | AGACCGCAAGTTCTCTTAGC  | TCCGGTGTAGCTGTTCAGT |
| CHS1      | TTCCACCTCCTCAAAAGATGTC | GCTCAACTCTGTTCACCTCAACCTC |
| CHS2/CHS9 | ATCCAGAAAGGGAGCAAGGGGC | ATCCGCTGTAGACCAACAT |
| CHI1      | TCCGTCACAGGTCAAACCT   | AAGACGGAGCGAGGAAGAT |
| F3H1      | GATGACATGGAAAGGCGATGTC | GTGGAGGGCAGTGAGGAAAT |
| F3′H1     | TTAGAGTGCTGGAGGTTGAGGGA | CTTTTGCGGCGGCAAGAC |
| DFR1      | GTATCAGCCTACGTCACAGG   | AGTTCAGCACGAGGAGTGAAT |
| LDOX1/LDOX2 | AGGTGCCACAGTGCCACATAGC | CCGCGTTCACGCCACCTAA |
| DcActin1  | CGGTATTTGTGGGACTCTGTGAGT | CAGCAAGTCAGACGAGTATAGG |

Table 2 Nucleotide sequences of primers specific to cyanidin-based anthocyanin biosynthetic genes and *Actin1* gene used for qRT-PCR

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Determination of anthocyanin content
Carrot taproots were ground to a fine powder in the presence of liquid nitrogen before anthocyanins were extracted. The total anthocyanin content of carrot taproots was determined in accordance with a previously described method [20]. Total anthocyanin quantities were reported in mg cyanidin 3-O-galactoside equivalents per 100 g fw (mg/100 g fw). Values were means of three independent experiments.

RNA isolation and cDNA synthesis
Total RNA was extracted from the taproots of carrots using an RNA Simple Total RNA Kit (Tiangen, Beijing, China) according to manufacturer’s instructions. First-strand cDNA was synthesized from 1 μg of total RNA using a PrimeScript™ RT reagent Kit with a gDNA Eraser (Perfect Real Time) kit (Takara, Dalian, China) following the manufacturer’s protocol. cDNA was diluted 20-fold for gene cloning and qRT-PCR analysis.

Figure 5 Expression of cyanidin-based anthocyanin biosynthetic pathway genes in 60-day-old stage carrot taproots. The mRNA level of actin1 was defined as 1. Data represents means of biological triplicate qRT-PCRs ± SD. Statistical analysis of differences was performed using Duncan’s multiple range test. Significant differences are indicated by different letters at the $P < 0.05$ level.
Gene identification, cloning, and sequencing

Nucleotide sequences of genes were searched from the GenBank of the NCBI. Genes not accessed from the NCBI were searched using the genome and transcriptome databases of carrots [25]. Genes were further identified by cloning and sequencing from ‘Deep purple’ in accordance with a previously described method [32].

qRT-PCR analysis

Primer pairs for qRT-PCR were designed using Primer 5 with a temperature of 59–62°C, length of 19–24 bp, and GC content of 45–55% (Table 2). qRT-PCR was performed on a MyiQ Real-Time PCR Detection System (Bio-Rad) with a SYBR Premix Ex-Taq (Takara) in accordance with the manufacturer’s protocol. A total of 20 μL of each reaction contained 10 μL of SYBR Premix Ex-Taq, 2 μL of diluted cDNA, 0.2 μM of each primer, and 7.2 μL of ddH2O. The qRT-PCR conditions were as follows: denaturation at 95°C for 30 s; 40 cycles of 95°C for 10 s; and 60°C for 30 s. To confirm amplification purity, melt-curve analysis was performed over a temperature range of 60–95°C at the end of the qRT-PCR. The DcActin1 gene was used as an internal standard. Experiments were conducted in biological triplicate using three biological RNA samples for each carrot cultivar.

Statistical analysis

Differences in structural gene expression levels between different carrot genotypes were statistically analyzed using Duncan’s multiple-range test at a 0.05 significance level. Correlation analysis was performed to determine relationships between expression levels of genes encoding enzymes implicated in the anthocyanin pathway of anthocyanin biosynthesis (CHS1, CHS2/CHS9, CHI1, F3H1, F3’H1, DFR1, and LDOX1/LDOX2) and anthocyanin presence by logistic regression analysis at a 0.05 significance level.

Availability of supporting data

The data supporting the results of this article are included within the article.

Additional file

**Additional file 1: Table S1.** Correlation between the expression levels of CHS1, CHS2/CHS9, CHI1, F3H1, F3’H1, DFR1, and LDOX1/LDOX2 and anthocyanin presence by logistic regression analysis.

Abbreviations

PAL: Phenylalanine ammonia-lyase; CA4H: Cinnamate 4-hydroxylase; 4CL: 4-coumaroyl-coenzyme A ligase; CHS: Chalcone synthase; CHI: Chalcone-flavonone isomerase; F3H: Flavanone 3-hydroxylase; F3’H: Flavonoid 3’-hydroxylase; DFR: Dihydroflavonol 4-reductase; LDOX: Leucoanthocyanidin dioxygenase; UCGT: UDP-galactose:cytidine 3-O-galactosyltransferase.

Competing interests

The authors declare that there are no competing interests.

Author contributions

Conceived and designed the experiments: ASX ZSX. Performed the experiments: ZSX YH FW SX GLW ASX. Analyzed the data: ZSX. Contributed reagents/materials/analysis tools: ASX. Wrote the paper: ZSX. Revised the paper: ZSX ASX. All authors read and approved the final manuscript.

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