RESEARCH ARTICLE

Identification and Characterization of DcUSAGT1, a UDP-Glucose: Sinapic Acid Glucosyltransferase from Purple Carrot Taproots

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Abstract

Purple carrots accumulate abundant cyanidin-based anthocyanins in taproots. UDP-glucose: sinapic acid glucosyltransferase (USAGT) can transfer the glucose moiety to the carboxyl group of sinapic acid thereby forming the ester bond between the carboxyl-C and the C1 of glucose (1-O-sinapoylglucose). 1-O-sinapoylglucose can serve as an acyl donor in acylation of anthocyanins and generate cyanidin 3-xylosyl (sinapoylglucosyl) galactoside in purple carrots. This final product helps stabilize the accumulation of anthocyanins. In this study, a gene named DcUSAGT1 encoding USAGT was cloned from ‘Deep purple’ carrot taproots. Enzymatic activity was determined using high performance liquid chromatography (HPLC). The optimal temperature and pH value were 30°C and 7.0, respectively. Kinetic analysis suggested a $K_m$ (sinapic acid) of 0.59 mM. Expression profiles of DcUSAGT1 showed high expression levels in the taproots of all the three purple carrot cultivars but low expression levels in those of non-purple carrot cultivars. The USAGT activity of different carrots in vitro indicated that crude enzyme extracted from the purple carrot taproots rather than non-purple carrot taproots exhibited USAGT activity. These results indicated that DcUSAGT1 may influence anthocyanin biosynthesis of purple carrot taproots.

Introduction

As described in detail previously, carrots (*Daucus carota* L.) located in the second place in the aspect of economic value among all the vegetables in the European Union. They are also important vegetable crops worldwide [1, 2]. Carrots are biannual diploids (2n = 2x = 18) with a comparatively small genome about 470 Mb [3]. In the large family of Apiaceae, carrot is the most typical species at the genetic and molecular levels [4, 5]. Carrots are rich in a variety of nutrition, such as carotenoids, hydroxycinnamic acids, anthocyanins, potassium, vitamin B6, vitamin C, folic acid, and magnesium [6]. Purple carrots (*D. carota* ssp. *sativus* var. *atorrubens* Alef.) particularly accumulate rich anthocyanins in taproots. Anthocyanins are antioxidants
that can enhance memory and vision [7]. Under natural conditions, free anthocyanins are rare and are usually acylated with sinapic, ferulic, and coumaric acids in the last step of anthocyanin biosynthesis [8].

The involvement of UDP-glucose is well identified in the formation of phenylpropanoid glycosides. Sinapic acid is a major phenylpropanoid participating in two different pathways, which operates at different stages and in different tissues during the development of plants [9]. Glucosyltransferases play important roles in the formation of the intermediates during these pathways. One pathway produces sinapoylmalate and sinapoylcholine through the synthesis of 1-O-sinapoylg glucose [10]. The other pathway goes through the offspring of sinapyl alcohol-4-O-glucoside, generating syringyl units found in lignin [11]. UDP-glucose: sinapic acid glucosyltransferase (USAGT) participates in the first pathway (Fig 1). It is an essential enzyme during the transfer of sinapic acid to the product of 1-O-sinapoylglucose.

UDP-glucose glucosyltransferase activity has been detected in many plants, such as UDP-glucose: flavonol 3-O-glucosyltransferase from soybean cell cultures [12], cyanidin 3-O-glucosyltransferase from flower cell cultures [13], UDP-glucose: thiohydroximate glucosyltransferase from seedlings of Brassica napus [14], and USAGT from cultured cell extracts of D. carota [15]. Although the enzymatic activity of these glucosyltransferases has been recognized for many years, most of their corresponding genes have not been identified [16]. The gene encoding for USAGT was cloned from B. napus [17]. However, the USAGT-gene in carrot has not been identified in detail. Furthermore, the potential function of USAGT in anthocyanin biosynthesis of purple carrot taproots has not been studied yet. Our work aimed to investigate the role of USAGT in anthocyanin biosynthesis of purple carrot taproots. Here, a gene named DcUSAGT1 that encodes for USAGT was identified from purple carrot taproots. We also reported the purification of rDcUSAGT1 (recombinant DcUSAGT1) protein and detected rDcUSAGT1 activity.

DcUSAGT1 may play important roles in the stability of anthocyanin accumulation, the expression levels of DcUSAGT1 gene in the purple carrot taproots may be higher than those in non-purple carrot taproots. Three purple carrot cultivars ‘Deep purple’, ‘Purple68’, ‘Tianzi2-hao’, and three non-purple carrot cultivars ‘Kurodagosun’, ‘Sanhonglucun’, ‘Junchuanhong’ were selected for expression profiles performance to verify this assumption. Expression patterns of DcUSAGT1 and crude enzyme of DcUSAGT1 in purple and non-purple carrots were also detected and analyzed. Those results will help us to further determine the effect of DcUSAGT1 on cyanidin-based anthocyanin biosynthesis.

Materials and Methods
Sequence database construction

The template sequence for database search is glycosyltransferase (AT3G21560) from Arabidopsis thaliana. Then the sequences of DcUSAGT1 gene were retrieved from CarrotDB, a genomic
and transcriptomic database for carrots built by our group (http://apiaceae.njau.edu.cn/carrotdb/) [18]. The nucleotide sequence was submitted to GenBank and assigned a GenBank accession number of KT595241. All primers used in this study are listed in Table 1.

Structure prediction and structure-based sequence alignment

Protein 3D structures were predicted on the Phyre2 website (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index) [19]. The domain of the proteins was analyzed using the Conserved Domain Database of the NCBI website. DNAMAN 6.0 software was used to complete the multiple sequence alignments of UDP-glucose: glucosyltransferase domains.

Plant material and treatment

This experiment was conducted in the State Key Laboratory of Crop Genetics and Germplasm Enhancement of Nanjing Agricultural University. The carrot seeds: purple carrot cultivars 'Deep purple', 'Purple68', and 'TianziZhaor', non-purple carrot cultivars 'Kurodagosun', 'Sanhongliucun', and 'Junchuanhong' used in this study were kept in our Lab (Lab of Apiaceae Plant Genetics and Germplasm Enhancement, Nanjing Agricultural University). After seven-day germination, carrots were transferred into plates which mixed with organic soil and vermiculite (3:1). The artificial climate chamber programmed for 12 h/12 h day/night at 25°C/18°C with a relative humidity of 60%–70%. The intensity of illumination is 300 μmol·m⁻²·s⁻¹.

For gene cloning, the whole taproots of purple carrot cultivar 'Deep purple' were harvested after two months. For expression profiles experiment, whole taproots of three purple carrot cultivars and three non-purple carrot cultivars were harvested after two, three, and four months, respectively. For the USAGT activity test in vitro, two-month taproots of two carrot cultivars 'Deep purple' and 'Kurodagosun' were harvested. All samples were immediately frozen in liquid nitrogen and stored at −80°C in a refrigerator before RNA extraction. The chemical reagents used for HPLC analysis in this study, such as sinapic acid and UDP-glucose, were purchased from Sigma (San Francisco, USA).

Constructing the expression vector to host the DcUSAGT1 gene

Total RNA from the taproot tissues of carrot cultivar 'Deep purple' were extracted using a total RNA kit (RNA Simply, Tiangen, Beijing, China) according to the manufacturer’s protocol. Reverse transcription into cDNA was conducted using PrimeScript RT reagent kit (TaKaRa, Dalian, China). cDNA was synthesized using 5 μg of total RNA as the template of the reverse transcription system in a 20 μL reaction volume. The DcUSAGT1 gene was amplified using diluted cDNA as the template with DcUSAGT1-Forward primer and DcUSAGT1-Reverse primer, respectively (Table 1). The DcUSAGT1 gene was cloned by polymerase chain reaction (PCR) using a forward primer 5’-ATGAGCCCTGTATATCAAGACTCCA-3’ and a reverse

| Name                | Oligonucleotides sequences        |
|---------------------|-----------------------------------|
| DcUSAGT1-Forward primer | ATGAGCCCTGTATATCAAGACTCCA         |
| DcUSAGT1-Reverse primer   | TCAGTTCACTCATACGCTTATCCACA        |
| Dactin-Forward primer    | CGGTATTGTGTTGGACTCTGGTGAT         |
| Dactin-Reverse primer    | CACCAAGGTCAGCAGACCGAGATGG         |
| DcUSAGT1-QForward primer | TTTGATTACGGTGTTGGTGTTGGT         |
| DcUSAGT1-QReverse primer | GCTGGATAAGGAGGTTGGAAGGTTG        |

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primer 5’-TCAGTTCATCATCAGCTTATCCACA-3’, and was then sequenced by Applied Biosystems 3730 DNA Analyzer (Genscript, Nanjing, China). The PCR conditions were as follows: 95°C for 5 min; followed by 35 cycles of 94°C for 30 s, 54°C for 15 s, and 72°C for 60 s; and 72°C for 10 min. The PCR product was recovered and inserted in a plasmid pMD-19 vector (TaKaRa, Dalian, China), and the ligation mixture was applied to transform the *Escherichia coli* strain (DH5α). The ampicillin resistant bacteria in the LB medium were identified, and plasmid DNA was sequenced afterward (Genscript, Nanjing, China). The *DcUSAGT1* gene was then inserted into a pET-30a (+) expression vector (Novagen, Darmstadt, Germany) with the Sac I/Hind III sites and transformed into *E. coli* DH5α. The authenticity of the result was confirmed by PCR analysis and sequencing.

**Purification of DcUSAGT1 protein**

To produce DcUSAGT1 protein, a single colony of *E. coli* BL21DE3 cells (TransGen, Beijing, China) carrying recombinant plasmid pET-30a (+)-DcUSAGT1 was grown in 50 mL of fresh LB medium containing 50 mg/mL kanamycin until OD600 value reached 0.4 to 0.6 at 37°C. The recombinant protein was then induced with the addition of 1 mM IPTG at 18°C, whereas the control culture lacked IPTG induction. After over 12 h of induction, the cells were harvested by centrifugation for 8 min (4°C, 4,000 × g) and re-suspended in 2 mL lysis buffer (pH = 7.5) containing 50 mM NaH2PO4, 300 mM NaCl, 10% glycerol, 10 mM β-mercaptoethanol, and 10 mM imidazole. The suspension was then sonicated on ice for 20 min. The mixture was centrifuged for 10 min (4°C, 12,000 × g) and filtered with 0.22 μm microfiltration membranes.

The protein was purified in a column containing Ni-NTA-agarose resin (1.5 mL bed volume) according to the instructions from the QIA-expression system (Qiagen, Hilden, Germany). The recombinant protein fraction was desalted in buffer (pH 7.5, 50 mM NaH2PO4-Na2HPO4, 0.1 mM EDTA, and 1 mM DTT) using HiTrap Desalting Column (GE, Fairfield, USA) according to the manufacturer’s instructions. The purified protein assay was quantified using Bradford assay kit with BSA as standard.

**Enzyme activity assay**

The assay mix (100 μL) was consisted of 5 μL recombinant protein, 5 mM sinapic acid, 50 mM UDP-glucose. The activity was performed in 50 mM buffer (NaH2PO4-Na2HPO4, pH 7.0) at 30°C for 30 min. The enzyme which was boiled at 99°C for 5 min was taken as control. After incubation, the reaction was terminated by adding 100 μL absolute ethyl alcohol. A total of 20 μL of the supernatant was used for analysis by HPLC.

**Optimum pH and temperature of the enzyme activity**

The determination of optimum pH for enzyme activity was performed in 50 mM NaH2PO4-Na2HPO4 at different pH values from 6.0–8.0 and in 50 mM glycine-NaOH (pH 8.5–9.5) at 35°C for 3 h. The optimum temperature for enzyme activity was determined at pH 7.0 in 50 mM NaH2PO4-Na2HPO4 at different temperatures (20–50°C) for 3 h.

**Optimal kinetic analysis of the enzyme activity**

The reactions contained fixed UDP-glucose concentration of 50 mM and sinapic acid concentration varying from 0.05 mM to 1.6 mM in 50 mM NaH2PO4-Na2HPO4. The reaction products were carried out in pH 7.0 at 30°C for 3 h according to the pH optima and linearity of the
two preceding activities. Afterward, the reaction was stopped by the addition of 100 μL of quick-frozen ethanol and stored at –20°C before the HPLC analysis.

**HPLC analysis**

Enzymatic activity was determined by HPLC in absorbance at 306 nm. Precision instrument (Agilent Technologies1200 series) with a Columbus C18 column (4.6×150 mm) was used for HPLC analysis. Acetonitrile in H2O (containing 1% glacial acetic acid) was used as mobile phase. Separations were accomplished for 20 min in a linear gradient from 10% to 40% acetonitrile in H2O with the flow rate of 1 mL/min.

Expression profile analysis of *DcUSAGT1* gene in purple and non-purple carrot cultivars

Quantitative real-time polymerase chain reaction (qRT-PCR) systems were performed on ABI7500 (Applied Biosystems,USA) with SYBR Premix Ex Taq (TaKaRa, Dalian, China). Each reaction was repeated three times with independent RNA samples for relative accuracy. Expression levels of mRNA for *DcUSAGT1* gene were measured using the primer pairs of *Dcactin* gene for qRT-PCR expression analysis [20]. All primer pairs used in qRT-PCR reaction are listed in Table 1. Reaction volume (20 μL) was composed of 10 μL of SYBR Premix Ex Taq, 7.2 μL of ddH2O, 0.4 μL each of primer, and 2 μL of diluted cDNA. qRT-PCR conditions were 95°C for 30 s, followed by 40 cycles of 95°C for 10 s and 60°C for 30 s, and finally 65°C for 15 s.

**Determination of the USAGT activity of carrot in vitro**

The purple carrot ‘Deep purple’ and non-purple carrot ‘Kurodagosun’ were selected to extract crude enzyme for *DcUSAGT1* activity determination. The reaction system was the same to the glucosyltransferase activity assay described previously. A total of 20 μL of the supernatant was used for analysis by HPLC.

**Results**

**Sequence and structure analysis of DcUSAGT1**

The result of cloning of the cDNA showed that the gene *DcUSAGT1* has an open reading frame of 1,473 bp encoding 491 amino acids. The calculated relative molecular mass (Mr) of *DcUSAGT1* is 54.55 kDa. Based on NCBI search, the protein contains conserved domains of the glycosyltransferase family, which belongs to the glycosyltransferase-B type superfamily. Structure-based sequence alignment was performed on the *DcUSAGT1* of other plant UDP-glucose: glucosyltransferases (Fig 2), such as UDP-glucose: flavonoid 3-O-glucosyltransferase from *Medicago truncatula* (Accession No. XP_003610163.1), UDP-glucose: anthocyanidin 3-O-glucosyltransferase from *Clitoria ternatea* (Accession No. 3WC4_A), UDP-glucose: flavonoid 3-O-glucosyltransferase from *Vitis vinifera* (Accession No. AAB81682.1), cytokinin-O-glucosyltransferase from *M. truncatula* (Accession No. XP_003618665.1), and UDP-glycosyltransferase 72B1 (UGT72B1) from *A. thaliana* (Accession No. NP_192016.1). The consensus of domain sequence was 42.19%.

3D structure was constructed based on UGT72B1 as the closest model. *DcUSAGT1* displays 25% identity with UGT72B1. *DcUSAGT1* and UGT72B1 both belong to glycosyltransferase-B type superfamily, as expected of the glycosyltransferase-1 enzyme family. The glycosyltransferase-B structure exists in the domain of *DcUSAGT1* and UGT72B1. *DcUSAGT1* features of N- and C-terminal domains, which is similar to those of UGT72B1. In *DcUSAGT1*, the N-terminal domain contains eight alpha helixes and five beta sheets, whereas the C-terminal covers
seven alpha helixes and six beta sheets. However, eight alpha helixes and six beta sheets were observed in the N-terminal of UGT72B1, whereas nine alpha helixes and six beta sheets occurred in the C-terminal. These two domains were separated by a deep cleft for substrate binding (Fig 3).
Screening for rDcUSAGT1 activity

Initial screening for rDcUSAGT1 activity was carried out at pH 7.0 (50 mM NaH2PO4-Na2HPO4) under 30°C for 30 min. The retention time of product (1-O-sinapoylglucose) and substrate (sinapic acid) was about 5.6 and 9.1 min, respectively (Fig 4). Comparing the control (using deactivated rDcUSAGT1) and the blank control, no product activity was detected.

Fig 3. Predicted 3D structure of UGT72B1 and DcUSAGT1. (A) 3D structure of UGT72B1. (B) 3D structure of DcUSAGT1. N represents the N-terminal, C represents the C-terminal. Red-helix represents α-helix, blue-sheet represents β-sheet. The long deep cleft in the middle of the structure is used for substrate binding.

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Fig 4. Chromatograms of product formation by the purified enzyme of UDP-glucose: sinapic acid glucosyltransferase in carrots by HPLC. Peak area A represents the reaction mix without enzyme as the blank control. Peak area B represents the reaction volume with deactivated enzyme. Peak area C represents the reaction system with active enzyme. The retention time of the product: 1-O-sinapoyl-glucose was about 5.6 min, and the peaks for sinapic acid were 9.1 min in average.

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Effect of temperature and optimum pH

rDcUSAGT1 activity was determined at temperatures from 20°C to 50°C and the optimal temperature for rDcUSAGT1 activity was 30°C (Fig 5A). Whether the temperature increased or decreased, the activities always gradually declined. When the temperature reached 50°C, rDcUSAGT1 activity was not detectable. The optimum pH for the reaction was 7.0 in a pH range of 6.5–7.5 (Fig 5B), and the activity of rDcUSAGT1 stopped when the pH exceeded 9.0.

Kinetic parameters of rDcUSAGT1

The pattern of dependence of the rDcUSAGT1 reaction rate was consistent with Michaelis-Menten kinetic behavior. \( K_m \) and \( V_{\text{max}} \) values were estimated based on a Hanes-Woolf plot. \( K_{\text{cat}} \) value was calculated using the derived rDcUSAGT1 Mr of 54,556 g.mol\(^{-1}\). For the purified enzyme, the calculated \( V_{\text{max}} \), \( K_m \), \( K_{\text{cat}} \), and \( K_{\text{cat}}/K_m \) values of sinapic acid as substrate are listed in Table 2. The calculated \( V_{\text{max}} \) was 0.74 nkcat mg\(^{-1}\), \( K_m \) was 0.59 mM, \( K_{\text{cat}} \) was 0.04 s\(^{-1}\), and \( K_{\text{cat}}/K_m \) was 67.79 M\(^{-1}\)s\(^{-1}\).

Expression profiles of DcUSAGT1 in the taproots of six carrot cultivars at different stages

A total of 18 samples were chosen to perform qRT-PCR analysis (Fig 6). The expression profiles indicated that the DcUSAGT1 gene was up-regulated more significantly in purple carrots ('Deep purple', 'Purple68', and 'Tianzi2hao') than in non-purple carrots ('Kurodagosun', 'Sanhongliucun', and 'Junchuanhong') (Fig 7). The trend of gene expression levels in the root development stage of the non-purple carrots (90 d > 60 d > 120 d) was similar to that of the purple carrots.

Table 2. Kinetic analysis data of rDcUSAGT1.

| Substrate   | \( V_{\text{max}} \) (nkcat mg\(^{-1}\)) | \( K_m \) (mM) | \( K_{\text{cat}} \) (s\(^{-1}\)) | \( K_{\text{cat}}/K_m \) (M\(^{-1}\)s\(^{-1}\)) |
|-------------|-----------------------------------|--------------|----------------------------|---------------------------------|
| Sinapic acid| 0.74                              | 0.59         | 0.04                       | 67.79                           |

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carrot ‘Purple68’. The trend of DcUSAGT1 expression levels in the taproots of ‘Deep purple’ and ‘Tianzi2hao’ cultivars was 120 d > 90 d > 60 d. Among the purple carrots, the expression level of DcUSAGT1 gene from purple carrot was highest in ‘Purple68’ (at the stage of 90 d), followed by ‘Deep purple’ and ‘Tianzi2hao’.

The USAGT activity of different carrots in vitro

As illustrated in the chromatogram, the retention time of the produced 1-O-sinapoylglucose and sinapic acid was 9.9 and 15.1 min, respectively (Fig 8A). It indicated that the crude protein extracted from the taproots of the purple carrots exhibited USAGT activity. However, no product was detected in Fig 8B, which indicated that the crude protein from the non-purple carrots did not exhibit USAGT activity. In total, USAGT activity was only detectable in the taproots of the purple carrots.

Discussion

During the secondary metabolism of plants, numerous glycosides have already been isolated as biologically active compounds, and some of them have been widely used as important medicines [21]. Glycosyltransferases usually act in the final stages of plant secondary metabolism. They are used to stabilize and dissolve various low molecular mass compounds, such as flower pigments [22], and to regulate the action of functional compounds, such as plant hormones [23].

Purple carrot cultivars contain abundant anthocyanins in the taproots. In Glässgen’s report, the main activity for UDP-glucose glucosyltransferase in anthocyanin production used sinapic acid as substrate [24]. Sinapic acid plays an important role in maintaining the stability of cyanidin-based anthocyanins [8]. USAGT can transfer sinapic acid from UDP-glucose to the

Fig 6. Photos of purple and non-purple carrot cultivars used in this study. D represents ‘Deep purple’, T represents ‘Tianzi2hao’, and P represents ‘purple68’, S represents ‘Sanhongliucun’, K represents ‘Kurodagosun’, and J represents ‘Junchuanhong’.

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product of 1-O-sinapoylglucose [25]. 1-O-sinapoylglucose can serve as an acyl donor in acylation of anthocyanins and generate cyanidin 3-xylosyl (sinapoylglucosyl) galactoside in purple carrots. This final product helps stabilize the accumulation of anthocyanins [8].

Structure prediction of DcUSAGT1 is based on the model UGT72B1. UGT72B1 is the first UDP-gluc-dependent NGT ever reported with glycosyltransferase-B topology [26]. The glycosyltransferase fold contains two β/α/β Rossmann domains, each corresponding to N- and C-terminals [27]. The structural model of DcUSAGT1 from purple carrot is similar to that of UGT72B1 from A. thaliana, which contains a similar motif. In purple carrot, sinapic acid may combine with the N-terminal domain, whereas UDP-glucose mainly binds to the C-terminal domain. The similar structural model between DcUSAGT1 and UGT72B1 indicated that DcUSAGT1 may also perform the function of UDP-glucose glycosyltransferase.

The Mr of glycosyltransferases is generally between 45 kDa and 60 kDa [28]. In this study, the expression vector hosting DcUSAGT1 gene was constructed, then rDcUSAGT1 protein was purified to detect enzyme activity. Here, we focused on the use of sinapic acid as substrate for
the determination of rDcUSAGT1 activity. Other studies also used other phenylpropanoids as substrates to detect enzymatic activity, such as cinnamic acid, caffeic acid, ferulic acid, and p-coumaric acid [11, 24]. The Mr of DcUSAGT1 was calculated to be 54.55 kDa. In a previous study, the gene encoding UDP-glucose: sinapic acid glucosyltransferase cloned from B. napus was predicted to have a Mr of 55.97 kDa [17]. Halaweish and Dougall found that crude extracts from wild carrots grown as cell cultures can catalyze the formation of 1-O-sinapoylglucose from sinapic acid and UDP-glucose. The activity of the purified enzyme was measured to identify its optimal temperature and pH value [15]. This report showed no significant difference in the amount of product formed at 30°C or 37°C, and optimum pH was between 6.5 and 7.0 [15]. In the present study, the peak of rDcUSAGT1 activity was at 30°C at pH value of 7.0. The $K_m$ value of rDcUSAGT1 for 5 mM of sinapic acid was 0.59 mM with 50 mM of UDP-glucose under optimal conditions. The $K_m$ value of rDcUSAGT1 was higher than that of UGT84A1 and UGT84A2 but lower than that of UGT84A3 for 1 mM of sinapic acid with 5 mM of UDP-glucose under optimal conditions in A. thaliana [10]. It indicated that the affinity of rDcUSAGT1 toward the substrate was high.

A previous study reported that genes PAL3/PAL4, CA4H1, 4CL1, CHS1, CHI1, F3H1, F3′H1, DFR1, and LDOX1/LDOX2 are powerful indicators for the production of anthocyanins in the taproots of purple carrot cultivars [29]. The transcripts for genes CHS1, DFR1, F3H, LDOX2, and PAL3 accumulated at high levels in purple carrots, less in purple-orange carrots, and low or no transcripts were detected in orange carrots [30]. In the present study, significant differences in DcUSAGT1 gene expression levels in purple carrots and non-purple carrots were detected. Significantly higher expression levels of the DcUSAGT1 gene in the taproots of all three purple carrots than non-purple carrot taproots were detected, which is similar to previous expression patterns [29, 30]. It indicated that the low expression of DcUSAGT1 gene in non-purple carrot taproots may be the determining step lost in anthocyanin production. To further prove the function of USAGT in carrot taproots, the USAGT activity of different carrot cultivars ‘Deep purple’ and ‘Kurodagosun’ was tested. The result showed that USAGT activity was detectable in the taproots of purple carrot in vitro instead of in non-purple carrot taproots.

In conclusion, the results presented here suggested that DcUSAGT1 is an efficient enzyme in the transformation of sinapic acid as substrate to product 1-O-sinapoylglucose and UDP. DcUSAGT1 maybe plays important roles in anthocyanins biosynthesis of purple carrot.
taproots. The study will provide a foundation for future understanding and manipulation of sinapic acid metabolism and anthocyanins synthesis in purple carrot taproots.

Supporting Information

S1 File. Relevant data underlying the findings described in the manuscript. Figure A. Nucleotide acid and deduced amino acid sequences of DcUSAGT1 from purple carrot.

(DOC)

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Author Contributions

Conceived and designed the experiments: ASX YYC. Performed the experiments: YYC ZSX. Analyzed the data: YYC ZSX. Contributed reagents/materials/analysis tools: ASX. Wrote the paper: CYY. Revised the paper: ASX ZSX.

References

1. Budahn H, Baranski R, Grzebelus D, Kielkowska A, Straka P, et al. (2014) Mapping genes governing flower architecture and pollen development in a double mutant population of carrot. Frontiers in Plant Science 5: 504. doi:10.3389/fpls.2014.00504 PMID: 25339960

2. Chen YY, Li MY, Wu XJ, Huang Y, Ma J, et al. (2015) Genome-wide analysis of basic helix-loop-helix family transcription factors and their role in responses to abiotic stress in carrot. Molecular Breeding 35: 125.

3. Schrader O, Ahne R, Fuchs J (2014) Karyotype analysis of Daucus carota L. using Giemsa C-Banding and FISH of 5S and 18S/25S rRNA specific genes. Caryologia 56: 149–154.

4. Alessandro M, Galmarini C, Iorizzo M, Simon P (2013) Molecular mapping of vernalization requirement and fertility restoration genes in carrot. Theoretic Applied Genetics 126: 415–423.

5. Wang GL, Xiong F, Que F, Xu ZS, Wang F, et al. (2015) Morphological characteristics, anatomical structure, and gene expression: novel insights into gibberellin biosynthesis and perception during carrot growth and development. Horticulture Research 2: 15028. doi:10.1038/hortres.2015.28 PMID: 26504574

6. Luby CH, Maeda HA, Goldman IL (2014) Genetic and phenological variation of tocochromanol (vitamin E) content in wild (Daucus carota L. var. carota) and domesticated carrot (D. carota L. var. sativa). Horticulture Research 1: 14015. doi: 10.1038/hortres.2014.15 PMID: 26504534

7. Böhm HG (1994) Mazza und E. Miniati: Anthocyanins in Fruits, Vegetables and Grains. 362 Seiten, zahlr. Abb. und Tab. CRC Press, Boca Raton, Ann Arbor, London, Tokyo 1993. Food / Nahrung 38: 343.

8. Montilla EC, Arzaba MR, Hillebrand S, Winterhalter P (2011) Anthocyanin composition of black carrot (Daucus carota ssp. sativus var. atrorubens Alef.) cultivars Antonina, Beta Sweet, Deep Purple, and Purple Haze. Journal of Agriculture and Food Chemistry 59: 3385–3390.

9. Milkowski C, Strack D (2010) Sinapate esters in brassicaceous plants: biochemistry, molecular biology, evolution and metabolic engineering. Planta 232: 19–35. doi:10.1007/s00425-010-1168-z PMID: 20428885

10. Lim EK (2001) Identification of glucosyltransferase genes involved in sinapate metabolism and lignin synthesis in Arabidopsis. Journal of Biological Chemistry 276: 4344–4349. PMID: 11042211

11. Whetten RW, MacKay JJ, Sederoff RR (1998) Recent advances in understanding lignin biosynthesis. Annual Review of Plant Physiology and Plant Molecular Biology 49: 585–609. PMID: 15012247

12. Poulton JE, Kauer M (1977) Identification of an UDP-glucose: Flavonol 3-O-glucosyl-transferase from cell suspension cultures of soybean (Glycine max L.). Planta 136: 53–59. doi: 10.1007/BF00387925 PMID: 24420227
13. Saleh NA, Fritsch H, Witkop P, Grisebach H (1976) UDP-Glucose: Cyanidin 3-O-glucosyltransferase from cell cultures of *Haplopappus gracilis*. Planta 133: 41–45. doi: 10.1007/BF00386004 PMID: 24425177
14. Reed DW, Davin L, Jain JC, Deluca V, Nelson L, et al. (1993) Purification and properties of UDP-glucose: thiohydroximate glucosyltransferase from *Brassica napus* L. seedlings. Archives of Biochemistry and Biophysics 305: 526–532. PMID: 8373190
15. Halaweish FT, Dougall DK (1990) Sinapoyl glucose synthesis by extracts of wild carrot cell cultures. Plant Science 71: 179–184.
16. Kovinich N, Saleem A, Arnason JT, Miki B (2010) Functional characterization of a UDP-glucose: flavonoid 3-O-glucosyltransferase from the seed coat of black soybean (*Glycine max* (L.) Merr.). Phytochemistry 71: 1253–1263. doi: 10.1016/j.phytochem.2010.05.009 PMID: 20621794
17. Milkowski C, Baumert A, Strack D (2000) Cloning and heterologous expression of a rape cDNA encoding UDP-glucose: sinapate glucosyltransferase. Planta 211: 883–886. PMID: 11144274
18. Xu ZS, Tan HW, Wang F, Hou XL, Xiong AS (2014) CarrotDB: a genomic and transcriptomic database for carrot. Database (Oxford). 2014, pii: bau096.
19. Kelley LA, Mezulis S, Yates CM, Wass MN, Sternberg MJ (2015) The Phyre2 web portal for protein modeling, prediction and analysis. Nature Protocols 10: 845–858. doi: 10.1038/nprot.2015.053 PMID: 25950237
20. Tian C, Jiang Q, Wang F, Wang GL, Xu ZS, et al. (2015) Selection of suitable reference genes for qPCR normalization under abiotic stresses and hormone stimuli in carrot leaves. PLoS One 10(2): e0117569. doi: 10.1371/journal.pone.0117569 PMID: 25658122
21. Jones P, Vogt T (2001) Glycosyltransferases in secondary plant metabolism: tranquilizers and stimulant controllers. Planta 213: 164–174. PMID: 11469580
22. Nishihara M, Nakatsuoka T (2011) Genetic engineering of flavonoid pigments to modify flower color in floricultural plants. Biotechnology Letters 33: 433–441. doi: 10.1007/s10529-010-0461-z PMID: 21053046
23. Bari R, Jones JD (2009) Role of plant hormones in plant defence responses. Plant Molecular Biology 69: 473–488. doi: 10.1007/s11103-008-9435-0 PMID: 19083153
24. Gläßgen WE, Rose A, Madlung J, Koch W, Gleitz J, et al. (1998) Regulation of enzymes involved in anthocyanin biosynthesis in carrot cell cultures in response to treatment with ultraviolet light and fungal elicitors. Planta 204: 490–498. PMID: 9684371
25. Strack D (1980) Enzymatic synthesis of 1-sinapoylglucose from free sinapic acid and UDP-glucose by a cell free system from *Raphanus sativus* seedlings. Zeitschrift füür Naturforschung C: Journal of Biosciences 35: 204–208.
26. Brazier-Hicks M, Offen WA, Gershater MC, Revett TJ, Lim EK, et al. (2007) Characterization and engineering of the bifunctional N- and O-glucosyltransferase involved in xenobiotic metabolism in plants. Proceedings of the National Academy of Sciences 104: 20238–20243.
27. Mulichak AM, Losey HC, Walsh CT, Garavito RM (2001) Structure of the UDP-Glucosyltransferase Gtb that modifies the heptapeptide aglycone in the biosynthesis of vancomycin group antibiotics. Structure 9: 547–557. PMID: 11470430
28. Kubo A, Arai Y, Nagashima S, Yoshikawa T (2004) Alteration of sugar donor specificities of plant glycosyltransferases by a single point mutation. Archives of Biochemistry and Biophysics 429: 198–203. PMID: 15313223
29. Xu ZS, Huang Y, Wang F, Song X, Wang GL, et al. (2014) Transcript profiling of structural genes involved in cyanidin-based anthocyanin biosynthesis between purple and non-purple carrot (*Daucus carota* L.) cultivars reveals distinct patterns. BMC Plant Biology 14: 262. doi: 10.1186/s12870-014-0262-y PMID: 25269413
30. Yildiz M, Willis DK, Cavagnaro PF, Iorizzo M, Abak K, Simon PW, et al. (2013) Expression and mapping of anthocyanin biosynthesis genes in carrot. Theoretic Applied Genetics 126: 1689–1702.