Transcriptomic dynamics changes in development of carmine radish (Raphanus sativus L.) fleshy roots using RNA-seq method

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Jian Gao
Yangtze Normal University

Mao Luo
Southwest Medical University

Yi Liu
Sichuan Agricultural University

Fabo Chen
Yangtze Normal University

Hua Peng
Sichuan Tourism College

Wenbo Li  liverb@163.com
Yangtze Normal University

Corresponding Author

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Abstract

Radish (Raphanus sativus L.), belonging to biennial root vegetable crop of Brassicaceae family, is an economically important vegetable crop with an edible taproot. Recently, most of differential expressed genes associating with anthocyanin biosynthesis have been identified in most of important fruit crops. However, transcriptome analysis of anthocyanin biosynthesis and expression of anthocyanin biosynthesis related genes in ‘Hongxin’ radish have not been fully investigated. Here, based on results from HPLC analysis, young fleshy roots obtained from the dynamics development stage of fleshy roots in carmine radish ‘Hongxin 1’ was used for RNA-Seq, including fleshy roots from seedling stage (SS), initial expansion (IE), full-expansion (FE), bolting stage (BS), initial flowering stage (IFS); full-bloom stage (FBS) and podding stage (PS). Subsequently, the putative candidate genes involved in the dynamics development stage of fleshy roots in carmine radish were identified. After that, DGE (differential gene expression) profile analysis was used to identify the pupative transcripts, compared with fleshy roots from seedling stage (SS). In addition, co-modulated DEGs (Common DEGs in the dynamic growing stages of fleshy root in carmine radish) were also identified, from which most DGEs were more likely to participate in anthocyanin biosynthesis, including two transcription factors RsMYB and Rs RZFP. In addition, some related proteins e.g. RsCHS, RsDFR, RsANS, RsF’3H, RsF3GGT1, Rs3AT1, glutathione S-transferase F12, RsUFGT78D2-like and RsUDGT-75C1-like were significantly contributed to the regulatory mechanism during anthocyanin synthesis in the development stage of fleshy roots. Furthermore, GO terms comprised of “anthocyanin-containing compound biosynthetic process” and “anthocyanin-containing compound metabolic process” were commonly overrepresented in the other dynamics growing stages of fleshy roots after initial expansion of fleshy roots. Moreover, these results indicated that five significantly enrichment pathways of DEG were identified for the dynamics
growing stages of fleshy roots in carmine radish, including Flavonoid biosynthesis, Flavone and flavonol biosynthesis, Diterpenoid biosynthesis, Anthocyanin biosynthesis, as well as Benzoxazinoid biosynthesis. These results will expand our understanding of complex molecular mechanism of the putative candidate genes involved in the dynamics development stage of fleshyroot in carmine radish.

Background

Anthocyanins, recognized as regulator for red to purple colors in nature, thereby producing water-soluble pigments belonging to the flavonoid group [1]. Most research has demonstrated that anthocyanins, as the benefit food additive worldwide, could release major public health threat (comprised of cardiovascular disease, inflammatory, obesity, and diabetes) caused by chemical-synthesis food additive[2, 3]. In addition, most of regulatory genes have been extensively found involved in the anthocyanin biosynthetic pathway, which were largely conserved among flowering plants[4]. Previous studies have demonstrated that anthocyanins are firstly formed from phenylalanine by series enzymes through phenylpropanoid metabolism, such as phenylalanine ammonia-layse (PAL), cinnamic 4-hydroxylase (C4H) and 4-coumarate-CoA ligase (4CL); subsequently followed by chalcone synthase (CHS), and then the product 4, 2′ 4′ 6′-tetrahydrocychalcone are further catalyzed successively by four enzymes [Chalcone Isomerase (CHI), flavanone 3-Hydroxylase (F3H), dihydroflavonol 4-Reductase (DFR), as well as anthocyanidin synthase (ANS/LDOX)][5, 6]. However, the molecular mechanism of anthocyanin biosynthesis regulation is still not fully understood in the dynamic development of fleshy roots in radish.

Recently, based on the global transcriptome technology (such as RNA-seq technology), most of differential expressed genes associating with anthocyanin biosynthesis and expression of anthocyanin biosynthesis have been identified in most of important fruit
crops. However, transcriptome analysis of anthocyanin biosynthesis and expression of anthocyanin biosynthesis related genes in ‘Hongxin’ radish (That is famous for containing natural red pigment (red radish pigment), as produced in Fuling district of Chongqing city, China) have not been fully investigated.

Results

**Dynamics Anthocyanidin profiles of fleshy root in development of carmine radish**

To demonstrate the dynamics Anthocyanidin profiles of fleshy root in development of carmine radish, Anthocyanidin profiles of fleshy root from seedling stage, initial expansion, full-expansion, Bolting stage, initial flowering stage, full-bloom stage and podding stage in five local *cultivars of carmine radish* (‘Hongxin 1’, ‘Guanguan’, ‘Longquan 1’, ‘Yanzhi 1’ and ‘Yanzhi 2’) were investigated by HPLC analysis. The results showed that Anthocyanidin was significantly increased in ‘Hongxin 1’ differential fleshy roots types with the development of dynamic growing stages of fleshy roots than other local cultivars of carmine radish, from seedling stage to full-bloom stage, but decreased in podding stage (Fig. S1).

**Illumina sequencing and De novo assembly**

Based on results from HPLC analysis, young fleshy roots obtained from the development stage of carmine radish ‘Hongxin 1’ was used for RNA-Seq in this study. The cDNAs obtained from fleshy roots of seven growth phases (seedling stage, initial expansion, full-expansion, Bolting stage, initial flowering stage, full-bloom stage and podding stage) were sequenced using Illumina sequencing technology. After filtering out adaptor-only reads, trimming reads and low-quality reads (base quality ≤10), high-quality Reads were aligned to the SSU and LSU rRNA sequences to remove rRNA reads by a home-made perl script. And the percentage of clean reads counts almost 70% after removing rRNA sequences on average among raw tags in each library, respectively *(Table 1)*. Moreover, 198, 342
assembled transcripts from the raw sequence reads were constructed with an average length of 411 bp, and 34,927 Unigenes were generated using paired-end reads with an average length of 768 bp through de novo assembly technology (Fig. S2).

**DEGs related to the dynamics growing stages of fleshy roots in carmine radish**

To identify the putative candidate genes with great changes involved in the dynamics growing stages of fleshy roots in carmine radish, normalized expression levels for all global expressed genes were analyzed, indicating that high distinct gene expression profiles exists in the dynamics growing stages of fleshy roots (Fig. 1, Table S2). More interestingly, we found that the putative candidate genes belong to Cluster 8 was consistently with dynamics anthocyanidin profiles of fleshy root in development of carmine radish, but the putative candidate genes categorized into Cluster 9 were found oppositely.

Furthermore, DEGs were identified the dynamics growing stages of fleshy roots among other different development periods (IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’) between ‘SS_root’ group, the results indicated that 1,629, 1,037, 1,385, 1,521, 1,574 and 917 DEGs were generated in IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’, compared with ‘SS_root’, including up-regulated (878, 755, 718, 838, 852 and 555 transcripts) and down-regulated genes (751, 282, 667, 683, 722 and 362 transcripts) (Fig. 2A). 126 Co-modulated DEGs (Common DEGs in the dynamic growing stages of fleshy roots in carmine radish) were identified based on venny graph (Fig. 2B), and expression changes pattern of co-modulated DEGs were displayed with different colors using heatmap (Fig. 2C). More importantly, we found some of co-modulated DEGs showed similar expression trends in the dynamics growing stages of fleshy roots, which was found consistently with anthocyanidin profiles of fleshy root in development of carmine radish, such as series functional enzymes acted as important regulators in
anthocyanins biosynthesis, including dihydroflavonol 4-reductase (*DFR*: Cluster_13775), flavonoid 3’-monooxygenase (*F3′H*: Cluster_4431), leucoanthocyanidin dioxygenase (*ANS*: Cluster_3903) and Chalcone synthase (*CHS*: Cluster_39833), as well as some regulation enzymes comprised of anthocyanidin 3-O-glucoside 2′″′-O-xylosyltransferase (*F3GGT1*: Cluster_9270), coumaroyl-CoA:anthocyanidin 3-O-glucoside-6″-O-coumaroyltransferase 1-like (*3AT2*: Cluster_46827), UDP-glycosyltransferase 75C1-like (*UGT75C1*: Cluster_2736) and UDP-glycosyltransferase 78D2-like (*UGT78D2*: Cluster_11854). In addition, sets of transport proteins and transcription factors, such as, glutathione S-transferase F12 (Cluster_24268), MYB transcription factor (Cluster_28373), as well as Zinc finger, RING-type protein (Cluster_7186) (Fig. 2C, Table S3). Moreover, these DEGs related to the dynamics growing stages of fleshy roots that involved in different biological processes were validated using qRT-PCR and the results showed higher consistent with expression profiles of RNASeq data (Fig. 3, Table S4).

**Functional annotation of DEGs related to the dynamics growing stages of fleshy roots in carmine radish**

To explore the regulatory mechanisms of DEGs related to the dynamics growing stages of fleshy roots in carmine radish, GO annotation and KEGG pathway enrichment of those putative DEGs were conducted. The results illustrated that GO terms comprised of “anthocyanin-containing compound biosynthetic process” and “anthocyanin-containing compound metabolic process” were commonly overrepresented in the other dynamics growing stages of fleshy roots after initial expansion of fleshy roots (IE-40 days after planting), “flavoriod biosynthetic process” and “flavoriod metabolic process” were found overrepresented in fleshy roots of IFS, FBS and PS; but for “pigment biosynthetic process” and “pigment metabolic process”, which was only found overrepresented in fleshy roots of IFS and FBS; moreover, we found that GO terms comprised of “glucosinolate biosynthetic
process” and “glucosinolate metabolic process” were only overrepresented in fleshy roots of IE. (Fig. 4A, Table S5). By conducting pathway enrichment analysis, these results indicated that five significantly enrichment pathways DEG were identified for the dynamics growing stages of fleshy roots in carmine radish, including Flavonoid biosynthesis, Flavone and flavonol biosynthesis, Diterpenoid biosynthesis, Anthocyanin biosynthesis, as well as Benzoxazinoid biosynthesis (Fig. 4B, Table S6).

Discussion

Anthocyanins have reported as important regulators for many of the red, purple and blue pigments in plants as a subgroup of flavonoids. Based on the anthocyanidin structures cyanidin, delphinidin and pelargonidin, more than 550 different anthocyanins have been isolated from diverse plant and identified by the extent of hydroxylation in the flavonoid B ring [12]. The researches have demonstrated that anthocyanin compounds were synthesized from methylation, glycosylation and acylation of the basic flavonol structure. Several structural genes and enzymes have been reported involved in anthocyanin and flavonoid biosynthetic pathway in fruits of most of plant species. Previous studies have demonstrated that anthocyanins are firstly formed from phenylalanine by series enzymes through phenylpropanoid metabolism, such as Chalcone synthase (CHS) contributed in flavonoid synthesis as the first important enzyme, followed by Chalcone isomerase (CHI) that acted as a central branching point of the flavonoid pathway, thereby leading to the formation of flavanones through closing the C-ring[13]. After that, Dihydroflavonol 4-reductase (DFR) and anthocyanidin synthase (ANS) were used to create proanthocyanindins for formation of anthocyanidins, or anthocyanin through the action of UDP-flavonoid glycosyltransferases (UFGTs)[14]. Here, dihydroflavonol 4-reductase (DFR: Cluster_13775), flavonoid 3'-monooxygenase (F3’H: Cluster_4431), leucoanthocyanidin dioxygenase (ANS: Cluster_3903) and Chalcone synthase (CHS: Cluster_39833) were
identified and validated using qRT-PCR, which was significantly up-regulated in the dynamics growing stages of fleshy roots among other different development periods from seedling stage to full-bloom stage, but decreased in podding stage; and showed consistently with dynamics anthocyanidin profiles of fleshy root in development of carmine radish. In addition, the expression of UFGT genes has also been identified as important regulator involved in the anthocyanin biosynthetic pathway [15, 16] that attached sugar moieties to the anthocyanin aglycone for stabilizing the anthocyanidin. In this study, UDP-glycosyltransferase 75C1-like (UGT75C1: Cluster_2736) and UDP-glycosyltransferase 78D2-like (UGT78D2: Cluster_11854) were significantly up-regulated in differential fleshy roots types with the development of dynamic growing stages of fleshy roots from seedling stage to full-bloom stage, but decreased in podding stage. After that, further modifications comprised of glycosylation, acylation or methylation existed for anthocyanins modifications, such as the formation of anthocyanin through a reaction catalysed by a cyanidin 3-O-glycosyltransferase. Here, anthocyanidin 3-O-glucoside 2'''-O-xylosyltransferase (F3GGT1: Cluster_9270), coumaroyl-CoA:anthocyanidin 3-O-glucoside-6''-O-coumaroyltransferase 1-like (3AT2: cluster_9270) were demonstrated as key regulation enzyme for formation of anthocyanin in carmine radish. In addition, sets of transport proteins and transcription factors, such as, glutathione S-transferase F12 (Cluster_24268), MYB transcription factor (Cluster_28373), as well as Zinc finger, RING-type protein (Cluster_7186). To date, GSTs were found involved in anthocyanin transport based on genetic and biochemical evidence[17]. Bz2, firstly demonstrated in Zea mays by its mutant bronze-2 as a GST-encoding gene, which was found involved in vacuolar transfer of anthocyanins (bz2)[18]. Anthocyanin accumulation and pigment mislocalization were found reduced in Arabidopsis, caused by Mutations in the GST-encoding genes[19]. In this study, GSTs F12 was significantly up-regulated in differential fleshy roots types with
the development of dynamic growing stages of fleshy roots. These findings provided further evidences for the role of GSH in anthocyanin transport mechanisms. Moreover, MYB is a key component of the central regulatory to determine variation of anthocyanin production [20]. More importantly, MYB transcription factors have been extensively studied for their roles in the regulation of pigmentation in plants. It is known that R2R3-type MYB proteins and the MYB-bHLH-WD40 complex directly activate the transcription of structural genes in the anthocyanin pathway, such as transcription of the early (CHS, CHI, F3’H and FLS) and late (DFR, ANS and ANR) flavonoid biosynthesis genes, respectively[21].

In this study, we also demonstrated that MYB transcription factors was significantly dynamically up-regulated and showed remarkable positive and significant correlation to red pigment content in differential fleshyroots types (Table S3). So we inferred that MYB transcription factors might specifically activate early flavonoid biosynthesis genes comprise of CHS, CHI and F3’H, as well as late flavonoid biosynthesis genes consisted of DFR and ANS in carmine radish, thereby directly playing important roles in anthocyanin biosynthesis. However, their molecular regulation mechanism awaits further investigation.

Methods

Plant material and experiment design

Five local cultivars of carmine radish (‘Hongxin 1’, ‘Guanguan’,‘Longquan 1’, ‘Yanzhi 1’ and ‘Yanzhi 2’) collected from Fuling were selected as experiment materials with containing natural red pigment (red radish pigment). For identify the pigment contents of five local cultivars of carmine radish, dynamics Anthocyanidin profiles of fleshy root in development of carmine radish were investigated by HPLC analysis, including fleshy root from seedling stage (SS-15 days after planting); initial expansion of fleshy roots (IE-40 days after planting); full-expansion of fleshy roots (FE-70 days after planting); bolting stage (BS-120 days after planting); initial flowering stage(IFS-140 days after planting);
full-bloom stage (FBS-160 days after planting); podding stage (PS-200 days after planting). Briefly, fleshy roots were collected from three homozygous of five local cultivars of carmine radish individuals and pooled together, respectively. The fleshy root tissues were grinded with liquid nitrogen and then we extracted the red pigment with a solvent mixture containing methanol (40%, v/v), formic acid (0.1%, v/v) and acetone (40%, v/v). We used a 10 μL injection volume for a VDS C-18 column (4.6 × 250 mm, 5 μm, VDS Optilab, Germany) with 0.8 mL min⁻¹ flow rate. Based on results from HPLC analysis, young fleshy roots obtained from the development stage of carmine radish ‘Hongxin 1’ was used for RNA-Seq in this study (Fig. S1).

The carmine radish ‘Hongxin 1’ was cultivated in a greenhouse at the experimental farm of the Yihe (Yangtze normal university experiment base) in 2018. Firstly, we sowed seeds of ‘Hongxin 1’ in sterilized soil for 2 weeks under normal growth conditions (23°C, 16 h light/8 h dark). After that, 2-week-old plants were transferred and kept for 15 days in the cold room (5 ± 1°C, 12 h light/12 h dark) for vernalization treatment. After the vernalization periods, the plants were grown in a normal growth room under normal growth conditions (23°C, 16 h light/8 h dark). At least three independent biological replicates for fleshy roots obtained from the development stage of carmine radish ‘Hongxin 1’ were collected for RNA-Seq with two replicates. All harvested tissue were immediately frozen in liquid nitrogen and stored at -80°C for RNA-seq analysis respectively.

**Sample preparation and library construction**

Library construction was conducted following NEBNext Ultra RNA Library Prep Kits for Illumina (NEB, USA), the mRNA was isolated through magnetic beads with Oligo (dT) using approximately 5 μg of total RNA and subsequently converted into short fragments by fragmentation buffer. After that, short fragments were converted into the first strands of
cDNA used as templates with random hexamers, as well as the second strands of cDNA were also synthesized, and then the desired synthesized cDNA fragments were purified (QiaQuick PCR kit) for PCR amplification, and the quantify and qualify of each sample library were checked by agilent 2100 Bioanaylzer and ABI StepOnePlus Real-Time PCR System. Ultimately, 200bp paired-end reads were generated from the prepared library with 2 replicates using Illumina HiSeqTM 2000.

**Reads processing and differentially expressed genes (DEGs) identification**

Using Trimmomatic software, clean reads were obtained through filtering out adaptor-only reads, trimming reads and low-quality reads (base quality ≤10). After that, high-quality Reads were aligned to the SSU and LSU rRNA sequences download from silvva database using bwa with parameters “-n 4 -o 1 -e 1 -i 0 -l 50 -k 2” and the mapped reads were removed by a home-made Perl script as rRNA reads. Subsequently, we used Trinity software to de novo assembled clean reads into transcripts, and all their responding unigenes were annotated through searching in national center for biotechnology information (NCBI) non-redundant protein (Nr) databases and Swiss-Prot protein databases using BLASTx search tool with threshold E-value set as less than 10. After that, the functions of assembled unigenes were annotated through gene ontology (GO, http://www.geneontology.org/) database and Kyoto encyclopedia of genes and genomes (KEGG, http://www.kegg.jp/) database. To assess the abundances of assembled transcript, we firstly mapped the clean reads of seven different fleshy roots libraries to the de novo assembled transcriptome using Bowtie2, and then assessed with RSEM through transcript quantification of the de novo assembly, only transcripts (FPKM ≥ 1) were considered as significant expressed transcripts[7]. At last, DEGs (Differential expressed genes) were then screened by noiseqbio [8] and then identify using a corrected P-value <0.05 between each set of compared samples. (The fold change of gene expression of six cultivars of
radish comprised of ‘IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’ were identified by comparing with ‘SS_root’ respectively, including ‘IE_root’ Vs ‘SS_root’, ‘FE_root’ Vs ‘SS_root’, ‘BS_root’ Vs ‘SS_root’, ‘IFS_root’ Vs ‘SS_root’, ‘FBS_root’ Vs ‘SS_root’ and ‘PS_root’ Vs ‘SS_root’). Furthermore, DEGs in the dynamic growing stages of carmine radish were analyzed and plotted using Neighbor-Joining cluster through homemade R script.

**GO functional annotation and KEGG pathway analysis of co-modulated differently expressed genes (DEGs) in growing stages of carmine radish**

Co-modulated DEGs (Common DEGs in the dynamic growing stages of fleshy roots in carmine radish) were identified based on venny graph. Subsequently, we conducted those co-modulated DEGs for GO annotation through Gene Ontology Database (http://www.geneontology.org/) and KEGG pathway enrichment analysis using KOBAS software, respectively [9]. In addition, the degree of KEGG enrichment was evaluated as the rich factor, q-value, and the number of genes in the enriched pathway. The rich factor refers to the ratio of the number of DEGs to the number of total annotated genes in a certain pathway. The q-value is a multiple hypothesis-corrected P value. The q-value can take on values between 0 and 1; values closer to 0 indicate more significant enrichment. After that, R script was used to construct their relative graphs.

**Validated of candidate DEGs invovled in the growing stages of carmine radish using real-time qRT-PCR**

To confirm the results obtained from the RNA-Seq assay, 11 DEGs with great alteration that related to growing stages of carmine radish were chosen and validated by qRT-PCR. The primers are designed by Primer 5.0 software for qRT-PCR experiments and radish gene (Actin) is used as a standard control (Table S1). The amplification programs were performed according to the standard protocol of the ABI7500 system, and conducted in
triplicate as mentioned by Jian et al.[10]. The relative quantitative method \(2^{-\Delta \Delta CT}\) was used to calculate the fold change in the expression levels of target genes[11].

**Data deposition**

All the raw read sequences were deposited in the NCBI sequence read archive under the accession number PRJNA565866.

**List Of Abbreviations**

ASRGs: Anthocyanin synthesis-related genes; PAL: Phenylalanine ammonia-lyase; C4H: Cinnamate 4-hydroxylase; 4CL: 4-coumarate: CoA ligase; CHS: Chalcone synthase; CHI: Chalcone isomerase; F3H: Flavanone 3-hydroxylase; F35H: Flavonoid 3,5 –hydroxylase; DFR: Dihydroflavonols 4-reductase; ANS: Anthocyanin synthase; UFGT: UDP-glucose: Flavonoid 3-glucosyltransferase; FNS: Flavone synthase; FLS: Flavcnol synthase; MT: Metallothionein-like protein. SS: seedling stage; IE: Initial expansion; FE: Full-expansion; BS: Bolting stage; IFS: Initial flowering stage; FBS: Full-bloom stage; PS: Podding stage; DEGs: Differential expression genes; KEGG: Kyoto encyclopedia of genes and genomes; FPKM: Fragments per kilobase of transcript per million mapped reads; GO: Gene ontology; NCBI: National center for biotechnology information; BP: Biological process; MF: Molecular function; CC: Cellular component.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All data generated or analysed during this study are included in this published article [and
its supplementary information files].

**Competing interests**

The authors declare that they have no competing interest.

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**Authors’ Contributions**

Conceptualization, Jian Gao and Wenbo Li; Methodology, Mao Luo and Jian Gao; Software, Jian Gao and Mao Luo; Formal Analysis, Wen-Bo Li and Jian Gao; Resources, Hua Peng; Writing – Original Draft Preparation, Jian Gao; Writing – Review & Editing, Jian Gao and Mao Luo; Supervision, WenBo Li; Funding Acquisition, Wenbo Li. All authors have read and approved the manuscript for publication.

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Table

Table 1. Summary of raw reads for 14 samples (two replicates) of fleshy roots of seven growth phases
| Sample   | Total_Read_Count | rRNA_Read_Count | Clean_Read_Count | Clean_Rate |
|----------|-----------------|----------------|-----------------|------------|
| FBS_root_1 | 33990858        | 7871020        | 26119838        | 76.84%     |
| FBS_root_2 | 33990858        | 8386243        | 25604615        | 75.33%     |
| BS_root_1  | 33401149        | 7702113        | 25699036        | 76.94%     |
| BS_root_2  | 33401149        | 7587241        | 25813908        | 77.28%     |
| IE_root_1  | 28574219        | 4463319        | 24110900        | 84.38%     |
| IE_root_2  | 28574219        | 4663468        | 23910751        | 83.68%     |
| SS_root_1  | 20191382        | 504415         | 19686967        | 97.50%     |
| SS_root_2  | 20191382        | 560164         | 19631218        | 97.23%     |
| IFS_root_1 | 24547530        | 5534772        | 19012758        | 77.45%     |
| IFS_root_2 | 24547530        | 5976321        | 18571209        | 75.65%     |
| FE_root_1  | 14316434        | 3200500        | 11115934        | 77.94%     |
| FE_root_2  | 14316434        | 3092331        | 11224103        | 78.40%     |
| PS_root_1  | 20673951        | 6219688        | 14454263        | 70.08%     |
| PS_root_2  | 20673951        | 5789807        | 14884144        | 72.01%     |

Additional Files

Supplementary figures

**Fig. S1.** Dynamics Anthocyanidin profiles of fleshy roots in development of carmine radish

**Fig. S2.** Length distribution of contigs and Unigenes from 'Hongxin 1' carmine radish transcriptome.

Supplementary tables

**Table S1:** List of primers for qRT-PCR analysis of 11 candidate DEGs involved in the dynamics growing stages of fleshy roots in carmine radish

**Table S2:** The abundance of global expression genes identified using FRKM in the dynamics development of different fleshy roots types in carmine radish.

**Table S3:** Expression levels of co-modulated DEGs identified in the dynamics development of different fleshy roots.

**Table S4:** RNA-seq expression data and qRT-PCR data for each candidate DEGs involved in the dynamics growing stages of fleshy roots in carmine radish
Table S5: GO annotation of Co-modulated DEGs in carmine radish.

Table S6: KEGG pathway enrichment of Co-modulated DEGs in carmine radish.

Figures

Figure 1

Normalized expression levels for all global expressed genes involved in the dynamics growing stages of fleshy roots in carmine radish.
Figure 2

Transcriptional changes of DEGs involved in the dynamics growing stages of fleshy roots in carmine radish. A. Statistic of differentially expression genes (including up-regulated and down-regulated in each comparison groups) in the dynamics growing stages of fleshy roots (‘IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’), compared with ‘SS_root’ group. B. Venny graph of co-modulated DEGs (Common DEGs in the dynamic growing stages of fleshy root in carmine radish). C. Clustering and heat map of common differentially expressed (Co-modulated genes) based on the expression profiles in the dynamics growing...
stages of fleshy roots (‘IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’), compared with ‘SS_root’ group.

Figure 3

Validate of candidate Co-modulated DEGs involved in the dynamics growing stages of fleshy roots in carmine radish using qRT-PCR and then correlation between RNA-seq and qPCR data were conducted. Each RNA-seq expression data was plotted against that from quantitative real-time PCR and fit into a linear regression. Both x- and y-axes were shown in log2 scale and each color represented a different gene.
Figure 4

Functional enrichment analysis of differentially expressed genes (DEGs) related to the dynamics growing stages of fleshy roots in carmine radish. A. Enriched GO terms of DEGs related to the dynamics growing stages of fleshy roots in radish. GO terms are plotted on the ordinate, and the enrichment factor (rich factor) is plotted on the abscissa. The colour of points represents the q-value, and the size of points represents the number of DEGs mapped to the reference pathway. Legends for the colour scale of q-values and size-scaling of the number of DEGs are shown to the right of the plot. B. Pathway enrichment analysis among
differentially expressed genes related to anthocyanin synthesis in radish.

Enriched KEGG pathway terms divided by the dynamics growing stages (IE_root’, ‘FE_root’, ‘BS_root’, ‘IFS_root’, ‘FBS_root’ and ‘PS_root’), compared with ‘SS_root’. Red color indicates statically overrepresented.

**Supplementary Files**

This is a list of supplementary files associated with the primary manuscript. Click to download.

- Fig S2.png
- Fig S1.png
- 5-Supplemental_tables.xlsx