Identification of Light-Independent Anthocyanin Biosynthesis Mutants Induced by Ethyl Methane Sulfonate in Turnip “Tsuda” (Brassica rapa)

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Abstract: The epidermis of swollen storage roots in purple cultivars of turnip “Tsuda” (Brassica rapa) accumulates anthocyanin in a light-dependent manner, especially in response to UV-A light, of which the mechanism is unclear. In this study, we mutagenized 15,000 seeds by 0.5% (v/v) ethyl methane sulfonate (EMS) and obtained 14 mutants with abnormal anthocyanin production in their epidermis of swollen storage roots. These mutants were classified into two groups: the red mutants with constitutive anthocyanin accumulation in their epidermis of storage roots even in underground parts in darkness and the white mutants without anthocyanin accumulation in the epidermis of storage roots in aboveground parts exposed to sunlight. Test cross analysis demonstrated that w9, w68, w204, r15, r21, r30 and r57 contained different mutations responsible for their phenotypic variations. Further genetic analysis of four target mutants (w9, w68, w204 and r15) indicated that each of them was controlled by a different recessive gene. Intriguingly, the expression profiles of anthocyanin biosynthesis genes, including structural and regulatory genes, coincided with their anthocyanin levels in the epidermis of storage roots in the four target mutants. We proposed that potential genes responsible for the mutations should be upstream factors of the anthocyanin biosynthesis pathway in turnips, which provided resources to further investigate the mechanisms of light-induced anthocyanin accumulation.

Keywords: anthocyanin; EMS mutant; turnips; light

1. Introduction

Anthocyanins, synthesized through the flavonoid pathway, are important secondary metabolites for plants flower pigmentation, seed dispersal, fruits coloration and resistance against biotic or abiotic stresses [1–4]. Due to high antioxidant activity, anthocyanins are critical antioxidants to protect plants against accumulation of reactive oxygen species (ROS), as observed in rats and humans [5,6]. Bright organ colors derived from anthocyanin accumulation directly determine ornamental, diet and market values of fruits and ornamental crops. Therefore, a comprehensive understanding of mechanisms regulating anthocyanin biosynthesis is important for the bioengineering of anthocyanin production in ornamental and agricultural plants.

The biosynthetic pathways of anthocyanin in different species share a majority of common routes that have been well characterized [7–9]. The corresponding genes have been isolated from various anthocyanin mutants and natural variant plants [10–13]. In the biosynthetic pathway of anthocyanin, chalcone synthase (CHS) is one of the first determinants of anthocyanin biosynthesis, which
collaborates with other structural genes, such as chalcone isomerase (CHI), dihydroflavonol reductase (DFR), anthocyanidin synthase (ANS) and UDP-flavonoid glucosyl transferase (UFGT), to synthesize anthocyanins [14–16]. These genes are divided into two subgroups: early biosynthetic genes (EBGs) for production of flavonols and late biosynthetic genes (LBGs) leading to the production of anthocyanins in vegetative tissues. In Arabidopsis, activation of EBGs requires co-activator independent R2R3-MYB regulatory genes (MYB11, MYB12 and MYB111), whereas LBGs are activated by R2R3-MYB genes and basic helix-loop-helix (bHLH) proteins that form a MYB-bHLH-WD40 (MBW) transcriptional activation complex with a WD40 repeat protein [17,18]. The members of this regulatory complex consist of TRANSPARENT TESTA GLABRA 1 (TTG1); the R2R3-MYB proteins from PRODUCTION OF ANTHOCYANIN PIGMENT 1 (PAP1/MYB75), PAP2, MYB113, or MYB114; and the bHLH proteins from TRANSPARENT TESTA 8 (TT8), GLABRA3 (GL3), or ENHANCER OF GLABRA3 (EGL3). The formation of MBW complex has been demonstrated to play a key role in triggering anthocyanin accumulation [19].

The anthocyanin biosynthesis is affected by many different environmental factors, of which light is a critical one [20–22]. Light-regulated anthocyanin biosynthesis initiates by a group of plant photoreceptors perceiving light signals of different wavelengths and then transducing signals to promote expression of anthocyanin biosynthesis genes via intracellular second messenger systems. The signal transduction pathway of light-dependent anthocyanin biosynthesis involves at least three types of photoreceptors: the red and far-red light-sensing phytochromes, the blue/ultraviolet A perceiving cryptochromes, and the UV-B specific photoreceptor UV RESISTANCE LOCUS8 (UVR8) [23–27]. As a positive regulator of photomorphogenesis, ELONGATED HYPOCOTYL 5 (HY5) is required for light-induced anthocyanin accumulation in plants [28]. Although there is no physical interaction between HY5 and R2R3-MYB proteins (such as MYB12, MYB111 and PAP1), the expression level of these MYB factors is activated by light in a HY5-dependent manner [29,30]. CONSTITUTIVELY PHOTOMORPHGENIC 1 (COP1) is another main player in the integration of photomorphogenesis and anthocyanin biosynthesis. Especially, most of regulatory factors involved in anthocyanin production (such as PAP1 and PAP2) are substrates of COP1/SPA, which are E3 ubiquitin ligases. The cop1 and spa mutants, but not PAP1 over-expressing plants, are able to produce anthocyanins in dark-grown seedlings, indicating that light requirement for anthocyanin biosynthesis is a result of the light-mediated stabilization of PAP1 proteins [31]. However, besides HY5-COP1/SPA-R2R3MYB module, more knowledge about other regulators involved in anthocyanin biosynthesis in response to light needs further investigation.

Turnips (Brassica rapa) are economically important vegetable crops in the world with enriched nutrition and medicinal value in their swollen roots. Previous work has indicated that UV-A robustly induced anthocyanin accumulation in the epidermis of swollen storage root, on which single blue light showed no effect [32]. The mRNA levels of CHS remarkably increased after 6 h of UV-A treatment compared to other monochromatic lights, such as red, blue and UV-B, which is distinguished from the cryptochromes and UVR8 reactions in Arabidopsis. In addition, UV-A has been proved to exert different effects on induction of anthocyanin accumulation in hypocotyls of turnip seedlings [33]. In Arabidopsis, cryptochromes are considered as UV-A/blue light photoreceptors that play critical roles in regulating blue light-mediated photomorphogenic growth and developmental process. However, different responses to blue light and UV-A irradiation for anthocyanin accumulation in turnip “Tsuda” showed probability of distinct mechanisms, besides cryptochrome signal transduction pathway, regulating anthocyanin biosynthesis in response to UV-A [32].

In the last decade, mutagenesis was used to obtain a large number of mutants that accelerated study of many different signal transduction networks. Ethyl methane sulfonate (EMS) has been the most common mutagen of choice to establish a series of allelic mutations in all genes with desired identifiable characters in plants such as plant height, flowering time, leaf shapes and fruit color changes [34–37]. It produces random mutations in genetic material by nucleotide substitutions that primarily cause point mutations through alkylation on the O6 position of guanines. Concentration
of EMS is the most important factor that determines mutagenesis efficiency. In general, a EMS concentration yielding 50% seed lethality (LD50) is defined as an indicator for population development to obtain a large number of desirable mutations in many plants [37–39].

To identify novel genes associated with light-induced anthocyanin accumulation in turnip “Tsuda”, we mutagenized 15,000 wild-type (WT) seeds using an optimal concentration of EMS and observed several different phenotypes in M2 plants. Totally, 14 mutants with phenotypic changes of anthocyanin production in the epidermis of swollen storage roots were obtained after extensive screening. The test cross and backcross experiments in genetic analysis indicate that four light-independent anthocyanin mutants (w9, w68, w204 and r15) are caused by a different recessive gene. qRT-PCR analysis showed that anthocyanin biosynthesis genes were dramatically up-regulated in r15 and down-regulated in w9, w68 and w204, compared to those of in WT plants. In this study, we obtained a serial of stably inherited anthocyanin mutants for researches on light-induced anthocyanin accumulation, which may provide new insight into the network of anthocyanin biosynthesis in higher plants.

2. Results

2.1. Characterization of Turnip “Tsuda” (Brassica rapa)

Turnip belongs to species of Brassica and subspecies of Brassica rapa in Brassicaceae family. Purple turnip “Tsuda” is a biennial herb and mature plants need to be vernalized for blooming in winter. Vegetative stage of turnip begins approximately 5–10 days after sowing, while cotyledons of seedlings emerge above soil surface and expand to initiate leaf growth from the growing point (Figure 1A). With the sixth leaf growing, the root of turnip initiates swelling up to 4–5 cm in diameter (Figure 1B,C). During winter, low temperature contributes to vernalization of turnip and transforms turnip from vegetative stage to productive stage (Figure 1D). Besides swollen storage root and seed coat, other tissues of turnip are similar to B.rapa oilseed crops and barely accumulate anthocyanins (Figure 1E–I). Furthermore, the epidermis of swollen storage roots of turnip cultivar “Tsuda” accumulate anthocyanins in a light-dependent manner, which only pigment at the light exposed part rather than that in darkness covered with soil. Therefore, it is an attractive system for fundamental research of mechanism of light-induced anthocyanin biosynthesis in plants.

![Figure 1. Turnip cultivar “Tsuda” plant development: (A) turnip seedlings; (B) vegetative growth of plants; (C) root swelling stage; (D) main shoot of flowering plant; (E) fully expanded rosette leaf; (F) light-induced anthocyanin accumulation in swollen root; (G) bright-yellow flower; (H) silique; and (I) mature seeds. The white arrow in (A) indicates the growing point of seedlings. Bar = 0.5 cm.](image-url)
2.2. Optimization of Mutagen Dosage

Optimal mutagenesis will lead to an EMS mutant population with a high density of mutations but also vigorousness and fertility. To determine the optimal concentration of EMS for turnip mutagenesis, we used 200 seeds for each of six treatments (0%, 0.25%, 0.50%, 1%, 2%, and 5%) and calculated germination rates (GR) of treated seeds (Table 1). The results showed that the GR of seeds decreased with increasing EMS concentrations. Compared to 191 seeds of the control group (no EMS), the GR of seeds were significantly reduced to less than 20% under 1.0% EMS treatment and down to 0% with EMS concentration over 2%. These findings indicated that the EMS treatment had been effective on turnip seeds and higher EMS concentrations increased the probability of undesirable mutations with more lethality. Almost 51% seeds were germinated by 0.5% EMS treatment, the closest to LD50 among all treatment conditions. Thus, we used 0.5% EMS to induce mutagenesis in over 15,000 turnip seeds.

Table 1. The germination percentage of turnip seeds with increasing concentrations of EMS treatment.

<table>
<thead>
<tr>
<th>EMS Treatment (W/V)</th>
<th>0.00%</th>
<th>0.25%</th>
<th>0.50%</th>
<th>1.00%</th>
<th>2.00%</th>
<th>5.00%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total No. of seeds</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>No. of germinated seeds</td>
<td>191</td>
<td>148</td>
<td>97</td>
<td>29</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Germination rate (GR)</td>
<td>95.5%</td>
<td>74%</td>
<td>48.5%</td>
<td>14.5%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Relative germination rate</td>
<td>100%</td>
<td>77.5%</td>
<td>51%</td>
<td>15.5%</td>
<td>0%</td>
<td>0%</td>
</tr>
</tbody>
</table>

Germination rate (GR) was identified as the rate from number of germinated seeds compared with total number of treated seeds in each group, and relative germination rates were the rates of GR in each experimental group compared with that in control group.

2.3. Screening Light-Independent Anthocyanin Mutants Induced by EMS

We planted EMS-treated seeds in field to survey phenotypic variations in M1 generation (Figure 2A), and observed many types of abnormal phenotypes: purple color leaves, early flowering, white swollen root peels and curled leaves in different developmental stages of M1 population (Figure 2B–E). Among all treated seeds, 56.7% (8516) M1 plants were generated to reach maturity, 3.4% (291) of which did not germinate or produced viable plants in M2 population. Totally, we obtained 14 putative mutants that displayed phenotypic variations in anthocyanin accumulation in the epidermis of swollen storage roots from the M2 population. Among these mutants, eight plants (r1, r10, r15, r21, r30, r37, r53 and r57) constitutively accumulated anthocyanin even in the epidermis of swollen storage roots at underground parts and were designated as red mutants; and six plants (w1, w3, w9, w68, w146 and w204) displayed deficient anthocyanin production in the epidermis of swollen storage roots at both aboveground (exposed to sunlight) and underground parts and were designated as white mutants.

To confirm whether these mutants were affected by same genes to display phenotypic variations, we performed test cross experiments between M3 generation of mutants on same phenotypes and recorded phenotypes of their progenies. The results of F1 progeny generated by six white mutants showed that w68, w204 and w9 were caused by different mutated loci (Table S1). However, three progenies of the crossed red mutants showed mutation phenotypes as their parental strains (Table S2). The results indicated that r10, r15 and r57 probably attributed to mutations of the same gene induced by EMS. Additionally, the phenotypic variations of r15, r21, r30 and r37 were caused by mutations of different genes.

To examine the molecular basis of light-independent anthocyanin accumulation in these EMS mutants, we measured the anthocyanin content of each mutant. In WT plants, pigmentation in the epidermis of swollen storage roots showed a light inducible manner: anthocyanin accumulated only in the aboveground part of storage roots (Figure 3). In white mutants w9, w68 and w204, anthocyanin was undetectable in either the aboveground or underground parts of storage roots. In contrast to white mutants, the red mutants (r15, r21, r30 and r37) accumulated anthocyanin in both underground and aboveground parts of storage roots (Figure 3). However, we noticed that anthocyanin levels of
underground parts were lower than aboveground parts in red mutants as well as that in WT. These data suggest that anthocyanin biosynthesis of these red mutants is partially regulated by light.

Figure 2. Open field cultivation and phenotypic variations observed in M1 population: (A) open field cultivation of M1 population generated by EMS; and phenotypic characteristics of abnormal turnips with: purple leaf (B); early flowering (C); white swollen root (D); and curly leaf (E).

Figure 3. Characterization of light-independent anthocyanin mutants. Phenotypes of the WT and mutants isolated in M2 population are shown. White mutants with anthocyanin deficiency (w6, w18, w204); Red mutants with anthocyanin over accumulation in the epidermis of swollen storage roots (r15, r21, r30 and r37). The swollen roots under dash line were covered by soil, while the parts above dash line were exposed to sunlight. Total anthocyanin content in mutants and WT were measured. The white columns, aboveground part of epidermis of swollen roots; black columns, underground part of epidermis of swollen roots. The data represent mean values from three replicates with biological repeats (n ≥ 3). Error bars represent the standard error of the mean. ‘*’ represents significantly differences at the level of p < 0.01.
2.4. Genetic Analysis of the Anthocyanin Variations Mutants

We constructed F2 populations by crossing the M3 generation of each mutant line (r15, w9, w68 and w204) with WT. The derived F1 generation of four mutants showed similar phenotype to WT (Figure 4A–D). The F1 plants were self-fertilized to produce F2 progeny. In the F2 population, including over 150 individuals, two phenotypes segregated in the ratio of 3:1 (Figure 4E,F). Furthermore, the p value of χ2 test in each of four groups was more than 0.05, which fits the 3:1 ratio expected for one-locus segregation (Table 2). These results indicated that different single recessive genes controlled the mutation traits in w9, w68, w204 and r15 mutants.

![Figure 4. Phenotypes of F1 and F2 populations in different mutant lines. F1 generation of: r15 (A); w9 (B); w68 (C); and w204 (D). Ten individuals of each line were collected for observation. Partly F2 individuals of: r15 (E); and w68 (F).](image)

### Table 2. Segregations of F2 populations crossed by the WT.

<table>
<thead>
<tr>
<th>Content</th>
<th>r15</th>
<th>w9</th>
<th>w68</th>
<th>w204</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of WT plants</td>
<td>141</td>
<td>152</td>
<td>114</td>
<td>133</td>
</tr>
<tr>
<td>No. of mutant plants</td>
<td>55</td>
<td>38</td>
<td>45</td>
<td>35</td>
</tr>
<tr>
<td>Total plants</td>
<td>196</td>
<td>190</td>
<td>159</td>
<td>168</td>
</tr>
<tr>
<td>p value (χ2 = 3:1)</td>
<td>0.322</td>
<td>0.111</td>
<td>0.336</td>
<td>0.212</td>
</tr>
</tbody>
</table>

2.5. Scanning for Mutations by TILLING Analysis

TILLING (Targeting Induced Local Lesions in Genomes) is a widely used approach to determine target genes associated with specific mutation in mutant populations generated by EMS treatment. In turnip, structural genes BrCHS1, BrCHS4, BrCHS5, BrCHI, BrF3H, BrSTS, BrDFR and BrUFGT promote anthocyanin biosynthesis in response to light. The regulatory factors BrMYB12, BrPAP1, BrTT8, and BrTTG1 have positive effects on their target structural genes to regulate anthocyanin production, of which expression levels are partially influenced by light in a BrHY5-dependent manner (Figure 5). Therefore, sequences of anthocyanin biosynthesis genes (including structural genes and regulatory genes) and light responsive genes BrHY5 and BrCOP1 from mutants and WT plants were amplified for point mutation screening. From TILLING analysis, we detected a nonsense mutation occurred in the second exon of BrMYB4 in r30 [40]. Several mutations located in other anthocyanin synthesis genes were also founded. However, all of them were identified as synonymous mutations, such as BrCHS1 in w9, BrDFR in w68 and BrMYB12 in r15 (Figure 6). Taken together, these results suggested that the mutant phenotypes in these lines are likely due to mutations of factors other than the anthocyanin biosynthetic genes.
Figure 5. Anthocyanin biosynthesis pathways in turnips “Tsuda”. Enzymes encoded by flavonoid early structural genes (EBGs) and encoded by anthocyanin late structural genes (LBGs) are shown in bold. Products generated by each step are boxed. Regulatory genes are circled. These genes control structural genes expression to accumulate anthocyanins in a HY5-dependent manner under light.

Figure 6. TILLING analysis of anthocyanin biosynthesis genes and light responsive genes in white mutants and red mutants. Each sample was mixed by equal amounts of mutants and WT PCR products via specific primers amplification and digested by CelI, respectively. Seven mutants (w9, w68, w204, r15, r21, r30 and r37) of treated samples were loaded from left to right in 1.5% agarose gel for mutation scanning. Black arrows indicate CelI-digested fragments.
2.6. Expression Profiles of Anthocyanin Biosynthetic Genes

To further analyze the types of target mutants, we detected the expression levels of anthocyanin biosynthesis genes and light responsive genes of *B. rapa* in the epidermis of swollen storage roots (Table S3). In WT, anthocyanin biosynthesis genes displayed a light-dependent manner that increased 20–50 times in aboveground parts of swollen roots compared with those in soil (Figure 7). Transcript levels of all the structural and regulatory genes were much higher for *r15* than for the WT in the soil. However, in *r15*, expression levels of detected genes in aboveground parts of root were higher than those in underground parts, which also showed a light-dependent manner for anthocyanin accumulation like WT. *BrHY5* and *BrCOP1* are the upstream components of light-signaling pathway, of which the expression levels were similar between *r15* and WT. Except for *BrF3H* and *BrUFGT* in *w9*, expression of all target structural genes in *w9*, *w68* and *w204* was much lower than that in WT, corresponding to that of anthocyanin deficiency at aboveground storage roots. For the regulatory factors involved in anthocyanin biosynthesis, transcript levels of both *BrPAP1* and *BrTT8* were down-regulated in *w204*, while *BrPAP1* and *BrTTG1* expression was up-regulated and *BrTT8* was down-regulated in *w68*. The positive regulator *BrMYB12* and light responsive gene *BrHY5* exhibited higher transcript levels in *w9* and *w68*. Compared to WT, no change of light responsive genes *BrHY5* and *BrMYB12* expression was observed in *w204*, different from those in *w9* and *w68* (Figure 8). The expression level of *BrCOP1* in *w9* and *w204* were lower than that in WT, but showed no difference in *w68*. The systemic expression changes of anthocyanin related gene in these mutants further indicated that the mutated genes of *r15*, *w9*, *w68* and *w204* were likely to affect upstream components of anthocyanin biosynthesis to regulate light-induced anthocyanin accumulation.

![Figure 7](image_url). Expression analysis of anthocyanin biosynthesis genes and light responsive genes in underground and light-exposed parts of epidermis of swollen roots in *r15* and WT. The expression value for each sample was normalized to that of the *BrACTIN* gene, and the transcript level in underground parts of epidermis of swollen roots was set as 1.0. The white columns, aboveground part of epidermis of swollen roots; black columns, underground part of epidermis of swollen roots. The mean values of three biological replicates are shown with error bars (SD).
with optimal EMS concentrations as 0.3–0.4% for *B. rapa* (310 nm, 0.3 Wm⁻²) for turnip "Tsuda" mutagenesis (Table 1). Meanwhile, rapid reduction of germination was observed when the mutagen was normalized to that of the *BrACTIN* gene, and the transcript level in light-exposed epidermis of swollen roots of WT was set at 1.0. The expression levels of *w9* (gray column), *w68* (dark gray column) and *w204* (black column) were compared to WT (white column). The mean values of three technical replicates are shown with error bars (SD).

3. Discussion

Previous studies have showed that anthocyanins in the epidermis of swollen storage roots of turnip “Tsuda” accumulated specifically in response to UV-A (320–400 nm, 3.0 Wm⁻²) and blue + UV-B (310 nm, 0.3 Wm⁻²), which is also observed in seedlings [32,33]. The expression of *BrCHS1*, *BrCHS4* and *BrCHS5* increased remarkably under both UV-A and blue + UV-B light, but not in response to single blue light (470 nm, 10 Wm⁻²). The transcriptome analysis indicated that there may be some overlaps between UV-A and blue + UV-B signal transduction pathways, in which 70% up-regulated genes were co-regulated, including some structural genes (*BrCHS*, *BrCHI*, *BrF3H* and *BrANS*) and regulatory genes (*BrPAP1*, *BrTT8* and *BrMYB12*) involved in anthocyanin biosynthesis pathway [41].

To gain additional insight into the mechanism of UV-A or blue + UV-B induced anthocyanin accumulation in turnip "Tsuda", setting up a screen for mutants with light-independent anthocyanin accumulation is urgently needed. We employed a forward genetic screen approach because the genetic transformation is low efficiency with some malformed root-like structures developed when using tissue culture method. Hereby, we used EMS to generate a mutant population and screen for turnips with abnormal anthocyanin accumulation pattern in epidermis of swollen storage roots. For more effective mutagenesis, treatment with different EMS concentrations was carried out in a small scale. Several studies have been conducted to optimize EMS concentrations in constructing *Brassica* mutant libraries, with optimal EMS concentrations as 0.3–0.4% for *B. rapa* [42], 0.6% for *B. napa* [34,43,44] and 0.5% for *B. oleracea* [45]. These results indicate that 0.5% EMS is likely a suitable mutagenic concentration for *Brassica* species. Consistent with previous reports, our current study showed that 0.5% EMS were optimal for turnip “Tsuda” mutagenesis (Table 1). Meanwhile, rapid reduction of germination was observed when the mutagen concentration increased, which may be due to absorbance of mutagen by the seeds leading to damage of germ cell or defects in cell metabolic processes.
Phenotypic changes in M1 generation are reliable signals for mutagen efficiency. Two types of visible changes on leaf and root peels coloration were related to anthocyanin production, which further confirmed that a suitable effect had been reached at this EMS concentration for anthocyanin variation (Figure 2B,D). In the current study, more than 50 putative mutant lines were classified into two groups with opposite phenotypes: white mutant and red mutant. White mutants were visibly confirmed with pale or no anthocyanin production at epidermis of their storage roots under sunlight; red mutants constitutively accumulated anthocyanin in epidermis of both aboveground and underground parts of their roots. However, high proportions of these mutants were segregated and recovered to wild type in M3 generation, that may be due to dominant mutations or affected by environment in the M2 generation [37]. Therefore, along with the results of M3 generation, 14 stabled mutants were chosen for further study.

We performed TILLING analysis of anthocyanin related genes (Figure 5). Excepted for a nonsense mutation at N terminal of BrMYB4 in r30 mutant [40], we failed in identifying additional anthocyanin biosynthesis genes with another functional mutation in the rest of mutants. These results suggest that these mutants are caused by unknown genes, for which TILLING is unavailable or by known genes with unrevealed function in anthocyanin biosynthesis.

Higher level expression of anthocyanin biosynthetic genes in r15 at underground parts and aboveground parts was observed (Figure 7), which paralleled the upward trend in anthocyanin accumulation. Interestingly, we noticed that anthocyanin-related genes at aboveground parts in r15 were also evoked and accumulated more anthocyanin than that of at underground parts, which showed a light-regulated manner similar to WT. In Arabidopsis, the E3 ubiquitin ligase COP1/SPA inhibited photomorphogenesis and anthocyanin biosynthesis in darkness by degradation of HY5 and PAP proteins [46]. The capability of producing anthocyanins at underground parts of epidermis of storage roots in r15 probably due to stabilization of HY5 and PAP proteins, displaying a similar phenotype as cop1/spa mutant in Arabidopsis. Therefore, we hypothesized that the mutated gene in r15 probably involved in formation of ubiquitin ligase complex and partially impaired the activity of complex to increase PAP protein levels at the epidermis of underground parts of storage roots. Under sunlight, the induction of BrHY5 further activated BrPAP1 expression to produce more anthocyanins in r15. The differences on protein levels at underground parts and aboveground parts of epidermis of storage roots in r15 would be confirmed in further investigation. In w9 and w68, transcript levels of EBGs (BrCHS, BrCHI) and LBGs (BrDFR, BrANS1 and BrANS2) exhibited greatly reduced, which was consistent with anthocyanin deficiency phenotype. However, the expression level of their regulator BrMYB12 increased remarkably and BrPAP1 expression in w68 was enhanced five times compared to WT at the epidermis of underground parts of storage roots (Figure 8). The R2R3-MYB genes (for instance, MYB12 and MYB75/PAP1) play important roles in anthocyanin biosynthetic pathway (Figure 5). Recent data have indicated that expression levels of these MYB genes are regulated in a HY5-dependent manner under light conditions [29,30]. We noticed that BrHY5 exhibited a higher expression level in w9 and w68 mutants than WT, which probably contributes to enrichment of BrMYB12 and BrPAP1 expression. However, unexpected expression pattern of BrHY5 and BrCOP1 in these white mutants remained unknown. We predict the presence of other unknown genes that cooperate with BrHY5/BrCOP1 module to regulate anthocyanin accumulation in turnip, which need to be investigated in further studies. It seems that the causal gene of w9 is likely to function at upstream of BrCHS and BrCHI to regulate coloration in swollen roots. In turnip, Bttt8, along with BrPAP1 and BrTTG1, plays a critical role in anthocyanin accumulation via regulating expression of LBGs (BrDFR, BrANS1, BrANS2, and BrUFGT) [33]. Despite higher BrPAP1 expression, activity of MBW complex was limited due to robustly reduced Bttt8 expression in w68 (Figure 8). The gene expression analysis further showed that these white mutants, which shared anthocyanin deficiency phenotype, are identified as distinct genetic lines.

Structural genes, which are responsible for the anthocyanin biosynthesis, are well understood in plants. However, the regulatory network of anthocyanin biosynthesis is quite complicated and
regulated by many transcription factors. In this study, we chose structural genes and critical positive regulatory genes involved in anthocyanin biosynthesis pathway for analysis. However, besides these genes, negative regulatory genes such as MYB4, MYBL2 and SPL9 (Squamosa promoter binding protein-like 9) also regulate anthocyanin production in many different ways [28,47,48]. For instance, miR156 targets the SPL9 to stabilize MBW complex for anthocyanin accumulation [48,49]. The roles of microRNAs and protein levels of anthocyanin synthesis genes in these mutants need to be confirmed in future studies.

Recently, with the development of next-generation sequencing platforms, MBS technology (Mapping-by-sequencing) becomes a powerful tool for rapid identification in EMS-induced point mutations in plants. NGS platforms are now commercially available at affordable prices, and the MBS strategy has been widely used in model species such as Solanum lycopersicum, Oryza sativa and Arabidopsis [50–53]. It is sufficient for rapid simultaneously map and identify causal mutations in the genomes through sequencing genomic DNA of a pool of plants selected from an appropriate segregating population [50]. Therefore, this method is cheaper and powerful in identifying mutations in our mutants. Future work would be conducted to find out mutations, study genetic reasons for anthocyanin variations and investigate gene functions in light induced anthocyanin accumulation pathway.

In this study, we isolated two types of light-independent anthocyanin mutants and confirmed that four of them (w9, w68, w204 and r15) were controlled by different single recessive genes. These mutants provide resources for investigating the regulation of light-induced anthocyanin production in detail.

4. Materials and Methods

4.1. Plant Materials

Brassica rapa “Tsuda” are purple turnips provided from Flower Bioengineering Institute at Northeast Forestry University, Haerbin, China. Seeds were sown in August and grown in the greenhouse during winter. About 2 months after sowing, the roots of turnip initiate swelling, of which peels could be used for anthocyanin measurement and phenotypic investigation. After flowering the collected plants were hand-pollinated in the greenhouse to produce selfed progeny or crossed progeny on purpose.

4.2. EMS Mutagenesis and Plant Growth Conditions

Seeds were soaked in sterile water for 6 h and dried by filter paper. Then seeds were treated with different concentration of EMS range from 0.25% to 5% for screening and determining optimal concentration to construct mutant population. Seeds treated with phosphate buffer (pH 7.0) were used as control. After 16 h mutagenesis, seeds were washed by sterile water for 4 h. Then, treated seeds were sown and grown in 18 cm pots filled with soil in a greenhouse, in which the temperature was maintained around 22 °C. Germination rate (GR) were identified as the rate from number of germinated seeds compared with total number of treated seeds in each groups, and relative germination rates were the rates of GR in each experimental groups compared with that in control.

4.3. M1 and M2 Population Generation

EMS-treated seeds with optimal concentration of mutagen were germinated and planted in the open field. Each of the surviving M1 plants was bagged for selfing and individually harvested. At least ten seeds from each M1 line were planted in pots in the greenhouse for M2 phenotype observation. The phenotypes of anthocyanin accumulation in the epidermis of swollen storage roots were recorded. Then mutants in M2 plants were selfed for homozygous identification and screened for further research.
4.4. Anthocyanin Measurement

The surface part approximately 1–2 mm thick of the swollen roots in WT and mutants were peeled by punchers and weighed using scales. Three pieces of roots epidermis on each turnip were grinded to a fine powder in liquid nitrogen and soaked in 1 mL of methanol containing 1% HCl for 24 h at 4 °C. Before measurement, samples were centrifuged at 12,000 rpm at 4 °C for 15 min. Collected 0.5 mL supernatant into a new tube and diluted samples with same volume of measured buffer. Anthocyanin content was expressed as optical density (OD$_{530}$) per gram of peels of the swollen roots. Three biological replications were used to calculate anthocyanin content in mutants and WT.

4.5. Genetic Analysis on Light-Independent Anthocyanin Mutants

The mutant lines were backcrossed to WT to generate F$_1$ progeny, and the F$_2$ progeny were derived from self-pollination of the F$_1$. At least 10 individuals of F$_1$ and 200 F$_2$ plants were sown for genetic analysis. Anthocyanin content of each F$_1$ and F$_2$ plant was recorded for analysis.

4.6. TILLING Analysis

Cell was extracted from celery to recognize and cut mismatched bases in double-stranded DNA specifically. Mutations in anthocyanin-related genes were amplified by PCR using specific primers from WT and mutant genomes. The amplified products derived from WT, mutants and the mixture of these two samples were then subjected to a complete denaturation-slow annealing program to produce heteroduplexes: 99 °C for 10 min, 70 cycles of 70–64 °C for 20 s (−0.3 °C per cycle), then held at 15 °C. The annealed PCR products were digested with Cell nuclease at 45 °C for 30 min, in a 10 µL reaction mixture containing 8 µL PCR product, 0.5 µL Cell, 10 mM HEPES pH 7.5, 10 mM KCl, 10 mM MgSO$_4$, 0.002% Triton X-100 and 0.2 µg/mL BSA. The reaction was stopped by adding 2 µL 0.5 M EDTA (pH 8.0) to the reaction. Cell-digested PCR products were separated using 2% agarose gel electrophoresis.

4.7. RNA Extraction and qRT-PCR

The epidermis of swollen roots of 2-month old WT and mutants were collected then grinded to a fine powder by liquid nitrogen. Total RNA was isolated with TRIzol-Reagent. Poly (dT) cDNA was reverse-transcribed and removed gDNA from 1 µg of the total RNA by TransScript One-Step gDNA Removal and cDNA Synthesis SuperMix kit (TransGen Biotech, Beijing, China). Diluted cDNA was used as the template DNA for the quantitative RT-PCR of anthocyanin biosynthetic genes (Table S4). The PCR amplification conditions were described previously [40]. The reaction was performed on ABI7500 real-time PCR system with the Power SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA, USA), according to the manufacturer’s instructions. Melting curve analysis was performed to control the purity and specificity of amplification. The transcript levels were determined by relative quantification using the turnip BrACTIN (AF111812.1) as the internal control and calculated using the 2$^{-\Delta\Delta C_t}$ comparative threshold cycle (Ct) method. At least three biological and three technological replicates were used for analysis.

Supplementary Materials: Supplementary materials can be found at www.mdpi.com/1422-0067/18/7/1288/s1.

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Conflicts of Interest: The authors declare no conflict of interest.
Abbreviations

EMS Ethyl methane Sulfonate  
CHS Chalcone synthase  
CHI Chalcone isomerase  
F3H Flavanone 3-hydroxylase  
DFR Dihydroflavonol reductase  
ANS Anthocyanidin synthase  
UFGT UDP-flavonoid glucosyl transferase  
PAP1 Production of anthocyanin pigment 1  
TT8 Transparent testa 8  
TTG1 Transparent testa glabra 1  
HY5 Elongated hypocotyl 5  
COP1 Constitutively photomorphogenic 1

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