Over-expression of the Hybrid Aspen Homeobox PttKN1 Gene in Red Leaf Beet Induced Altered Coloration of Leaves

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Abstract

PttKN1 (Populus tremula × tremuloides KNOTTED1) gene belongs to the KNOXI gene family. It plays an important role in plant development, typically in meristem initiation, maintenance and organogenesis, and potentially in plant coloration. To investigate the gene functions further, it was introduced into red leaf beet by the floral dip method mediated via Agrobacterium tumefaciens. The transformants demonstrated typical phenotypes as with other PttKN1 transformants. These alterations were very different from the morphology of the wild type. Among them, morphological modification of changed color throughout the entire plant from claret of wild type to yellowish green was the highlight in those transgenic PttKN1-beet plants. The result of spraying selection showed that the PttKN1-beet plants had kanamycin resistance. PCR assay of the 3SS-Promoter, NPTII and PttKN1 gene, PCR-Southern analysis of the NPTII and PttKN1 gene showed that the foreign PttKN1 gene had successfully integrated into the genome of beet plant. Furthermore, the results of RT-PCR analysis showed that the gene was ectopic expressed in transgenic plants. These data suggested that there is a correlation between the ectopic expression of PttKN1 gene and morphological alterations of beet plants. Pigment content assay showed that betaxanthins concentrations shared little difference between wild type and transgenic lines, while betacyanins content in transgenic plants was sharply decreased, indicating that the altered plant coloration of the transgenic beet plants may be caused by the changed betacyanins content. The tyrosinase study suggested that the sharply decreased of betacyanins content in transgenic plants was caused via the decreased tyrosinase level. Therefore, the reason for the altered plant coloration may be due to partial inhibition of betacyanin biosynthesis that was induced via the pleiotropic roles of PttKN1 gene.

Keywords: betalains, coloration, floral dip, PttKN1 gene, tyrosinase

Introduction

Flavonoids/anthocyanins, betalains and carotenoids are three main pigments for plant coloration (Tanaka et al., 2008). Among them, carotenoids often have a wide variety in plant tissues, while anthocyanins and betalains do not coexist (Stafford, 1994; Tanaka et al., 2008). In general, anthocyanins are widely distributed in seed plants, but betalains are found in plant species only of the Caryophyllales order (Stafford, 1994). In the past decades, the biosynthesis and transcriptional regulation of anthocyanins (Koes et al., 2005; Tanaka et al., 2008) and carotenoids (Hirschberg 2001; Howitt Pogson, 2006) are well studied. But the knowledge on betalains is much less than the other two pigments (Tanaka et al., 2008). Betalains are classified into red betacyanins and yellow betaxanthins. They are responsible for brilliant color from yellow to red (Strack et al., 2003) in Caryophyllales plant including red leaf beet (Beta vulgaris var. cicla L.), which is one of the varieties of leaf beet (Xu et al., 2009). Published data indicated that betalains are derived from tyrosine and then form two pivotal precursors of L-(3, 4-dihydroxyphenyl) -alanine (Dopa) and cyclo-Dopa, which reaction was catalyzed by tyrosinase (Gandia-Herrero et al., 2005; Steiner et al., 1999). These are the first steps in the biogenesis of betalamic acid (then formed betaxanthins) and betacyanins (Gandia-Herrero et al., 2005). Thus, tyrosinase is thought to be the first key enzyme in betalains biosynthetic pathways (Van Gelder et al., 1997). A clear correlation between the tyrosinase activity and betacianin content in young Beta vulgaris L. subsp. vulgaris (Garden Beet Group) plants was also reported (Steiner et al., 1999).

PttKN1 (Populus tremula × tremuloides KNOTTED1), which is a hybrid aspen homeobox gene (Hu et al., 2005; Meng et al., 2009a) plays an important role in plant development, especially in meristem initiation, maintenance and organogenesis (Xu et al., 2011). It is well reported that the PttKN1 transformants of Begonia maculata (Xu et al., 2011), carnation (Meng et al., 2009a), cockscomb (Meng et al., 2009b), Petunia hybrida (Hu et al., 2005) and tobacco (Xu et
al., 2014) produced various morphological alterations compared with their wild type plant. Those phenotypes were mainly focused on altered leaf arrangement (Hu et al., 2005; Meng et al., 2009b; Xu et al., 2011) and leaf morphology like lobed leaf (Hu et al., 2005; Xu et al., 2011; Xu et al., 2014), cup-shaped leaf (Xu et al., 2011) and distorted leaf (Hu et al., 2005; Meng et al., 2009a; Xu et al., 2011; Xu et al., 2014) etc. However, transgenic PttKN1 P. hybrida showed color modification of flower (Hu et al., 2005) and B. maculata showed color modification of leaf (Xu et al., 2011). These phenomena suggested that PttKN1 gene may regulate the biosynthesis of anthocyanins. In fact, transgenic PttKN1 coleus (Solenostemon scutellarioides) showed a decreased anthocyanins level (Xu et al., 2013).

The present work describes the transformation of red leaf beet using the floral dip method mediated by Agrobacterium tumefaciens. The over-expression of the foreign PttKN1 gene produced pleiotropic morphological alterations including leaf color alteration from claret to yellowish green. This phenomenon suggested that PttKN1 gene may also impact the biosynthesis of betalain for it is responsible for the coloration of red leaf beet. Further analysis suggested that the alteration was related to the ectopic expression of PttKN1 gene and the decreased tyrosinase level.

Materials and methods

Plant transformation

Seed balls of red leaf beet (Beta vulgaris L. var. cida L.) were purchased from Lanzhou Botanical Garden, Gansu Province, China. They were planted and grown in the experimental field of Lanzhou University. The plants were geography separated to maintain humidity and keep them out of direct sunlight. After 3 weeks young leaves were introduced into the 25 ml liquid medium: BA 0.5 mg l−1, 0.01% Silwet LP77 and acetosyringone (AS) either or both into a solution with an optical density at OD600 of approximately 0.3. The tissue culture was transferred to YEB medium supplemented with 25 mg l−1 kanamycin (Km) and 75 mg l−1 carbenicillin under the condition of 27 °C and 180 rpm overnight. After that, 1/2 MS medium containing BA 0.5 mg l−1 and 3% sugar was prepared as the basic dipping medium. Then the cells were collected by centrifugation (2,000 g, 10 min) and diluted with the above basic medium, or the modified medium that comprised of the basic medium added 0.01% Silwet L-77 and acetosyringone (AS) either or both into a solution with an optical density at OD600 of approximately 0.3. Then the plants with developed inflorescence and many immature floral buds were selected to be inoculated with the above prepared A. tumefaciens for 1/2, 1, 3, 5, 10 and 15 min, respectively. The dipped inflorescences were covered with black plastic for 24 h to maintain humidity and keep them out of direct sunlight. After 3-5 wks, seed balls were harvested and vernalized in water solution with 100 mg l−1 km at 4 °C for 48 h. Then the seed balls were sown in mixture of vermiculite, perlite and sphagnum moss (1:1:1) and sprayed 300 ml km solution (100 mg l−1) per 0.3 × 0.45 m2 twice a week. One month later, kanamycin-resistant seedlings were recorded and the morphologies were observed and photographed. Transformation efficiency expressed as “percentage transformation”, was calculated as percentage of kanamycin-resistant seedlings within the total seedlings tested.

PCR assay

DNA was extracted from the leaves of wild type and putative transgenic plants by improved CTAB method (Dole, 1990). Transgenic plants were verified via PCR of the PttKN1 gene, NPTII gene and 35S-Promoter. The primers were designed for PttKN1 gene (Forward: 5’-CCATCAAGGCAAGATCATAG-3’ and Reverse: 5’-GGACCATGGCAGATCATATT-3’) to yield 311 bp fragments, NPTII gene (Forward: 5’-GTCATCTACCTGCTCCCTG-3’ and Reverse: 5’-GGCTTCCAATGGAATGACG-3’) to yield 142 bp fragments and 35S-Pro (Forward: 5’- AAGGAAGGGCCATGCTTAAGG-3’ and Reverse: 5’-ATAGAGGAAAGGGCTTCGAGG-3’) to yield 192 bp fragments.

PCR was performed in a total volume of 25 μl containing 10×PCR Buffer (Mg2+ plus) 2.5 μl, dNTP (2.5 mM each) 1 μl, each primer (10 μM) 1 μl, DNA 100–300 ng, TaKaRa Taq (5 U/μl) 0.2 μl. PCR conditions were as follows: 94 °C for 2 min; 32 cycles of 94 °C for 1 min; and finally elongated at 72 °C for 5 min. PCR products were run on 0.8% agarose gel and photographed with Alpha Imager™ 2000 Documentation Analysis System.

RNA extraction

Total RNA was isolated from leaves of putative transgenic and wild type plants using the RNA Easyspin Isolation System (Aidlab biotech, Beijing, China). RT-PCR was performed with one step RNA PCR kit (TaKaRa Biotechnology, Dalian, China) according to the manufacturer’s instructions. Conditions were the same as the above PCR assay. The PttKN1 gene amplification was carried out using the primers of outer (5’-CCACCAATTCTCGCCCTTATG-3’ and 5’-CTTCGATGAGCCGACACC-3’) and inner (5’-CCATCAAAGCCAAAGATCATAGG-3’ and 5’-GGACCATGGCAGATCATATTG-3’).

PCR-Southern blot

The PttKN1 and NPTII gene were amplified from pPCV702 plasmid, non-transformed control plant and the putative transgenic plant as the above PCR protocol. Therein, pPCV702 plasmid was regarded as positive control and non-transformed plant as negative control. The primer was the same as the above. PCR products were then separated on a 0.8% agarose gel and transferred to nylon membranes (Roche) with 20×standard saline citrate by a standard capillary transfer method (Sambrook and Russell, 2001). The probe preparation, hybridization and detection were performed using the DIG-High Prime DNA Labeling and Detection Starter Kit I (Roche) according to the manufacturer’s instruction.

Betalains content

Betalains were extracted from the second to fourth true-leaf stage, using about 0.5 g of wild type and transgenic plants in 25 ml sodium phosphate-citrate buffer (pH 5.0) for 30 min at room temperature. Then the extracts were diluted to 100 ml with the same buffer and centrifuged at 2,500 g for 10 min. The absorbance at 476 nm and 535 nm were measured for betaxanthin and betacyanin, respectively. The pigment concentration was calculated using extinction value E (1 cm/%)
of 750 for betaxanthin and 1120 for betacyanin (Nilson, 1970). The results were expressed as mg g\(^{-1}\) fresh weight. All experiments were performed in triplicate.

Tyrosinase assays

True leaves from the wild type and transgenic plants, about 0.5 g, were grinded and extracted for 2 h in 7.5 ml sodium phosphate buffer (pH 7.2). After centrifugation (2,000 g, 10 min), 6 ml of the extract was diluted to 10.0 ml with the same buffer. The enzyme activity was measured by a modified spectrophotometry as described by Li et al. (2009) with L-dopa (Sigma) as the substrate. The reaction mixture (5.0 ml of final volume) containing 2.0 ml L-dopa, 0, 0.1, 0.2, 0.3 or 0.4 ml tyrosinase extract and sodium phosphate buffer (pH 6.8) to total volume 5.0 ml. Then the mixture was incubated at 30 °C for 10 min and the absorption at 475 nm due to the formation of dopachrome (\(\varepsilon_{475} = 3700 \text{ M}^{-1}\text{cm}^{-1}\)) was measured with a Shimadzu UV-1700 spectrophotometer. All experiments were performed in triplicate.

Statistical analysis

All data were reported as means ± SE. Statistical analysis was performed using SPSS version17.0 (SPSS Inc., USA). T-test was performed to evaluate the effectiveness of operation. A P < 0.05 was considered statistically significant.

Results

Transformation of red leaf beet with floral dip method

Red leaf beet was transformed via floral dip method and then the seedlings of transformed progeny were selected via kanamycin spraying method. One month later, kanamycin-resistant seedlings were recorded and observed. Of the total of 1,600 seed balls sowed for test, 132 exhibited kanamycin resistance. It could be seen that dipped time and medium composition play important roles on the transformation efficiency (Table 1). When 1/2MS added 5 mg l\(^{-1}\) BA and 3% sugar was used as the basic A. tumefaciens inoculation medium, the red leaf beet plants with many immature floral buds were inoculated 1/2, 1, 3, 5, 10 and 15 min, respectively. Therein, instant treatment with 1/2 min could induce putative transformation, and the transformation efficiency was improved along the added dipped time to 5 min with the highest 2.63%, while further prolonged time was displayed unfavorable. When Silwet L-77 or acetosyringone was added to the basic inoculation medium respectively, they altered transformation efficiency by little compared with the basic medium used. While the two were used in pair, transformation efficiency was upgraded from 1.83% to 2.57%.

Phenotypes of the transgenic plants

Normal red leaf beet with two symmetrical cotyledons and clarret hypocotyl is a dicotyledonous plant (Fig. 1A). The plants have heart-shaped and clarret true leaves (Figs. 1F, 1I), raceme with a columned main inflorescence stem (Fig. 1N). The transformants obtained from the above floral dip method showed different phenotypes compared to their wild type plants. Of the total of 132 kanamycin resistant plants, 67 showed the same color as the wild type with some morphological changes

<table>
<thead>
<tr>
<th>Treatment</th>
<th>% transformation</th>
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<tbody>
<tr>
<td>Time (min)</td>
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<tr>
<td>1/2</td>
<td>0.56 ± 0.97a</td>
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<tr>
<td>1</td>
<td>0.67 ± 1.15a</td>
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<tr>
<td>3</td>
<td>1.83 ± 1.92c</td>
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<tr>
<td>5</td>
<td>2.63 ± 1.19d</td>
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<td>10</td>
<td>1.30 ± 1.12b</td>
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<td>15</td>
<td>1.87 ± 0.11c</td>
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<td>Medium composition**</td>
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<tr>
<td>Basic medium</td>
<td>1.83 ± 1.92c</td>
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<tr>
<td>Basic medium + Silwet L-77</td>
<td>1.33 ± 2.31b</td>
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<tr>
<td>Basic medium + AS</td>
<td>1.99 ± 0.02c</td>
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<tr>
<td>Basic medium + Silwet L-77 + AS</td>
<td>2.57 ± 1.07d</td>
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Note: basic medium (1/2MS + 5 mg l\(^{-1}\) BA+ 3% sugar) was used; "dipped 3 min.

The significant difference is 0.05

Fig. 1. Phenotypes of the wild type plant of red leaf beet and its transformants. A - wild type plant showed two cotyledons; B-E: transgenic beet plant; B - ectopic bud formed from hypocotyl (arrow); C - anisomereous cotyledons; D - cup-shaped cotyledon; E - three cotyledons with altered plant coloration; F - wild type plant with heart-shaped and clarret true leaves; G-H: transgenic beet plant; G - plant with similar morphology of wild type except for coloration; H - plant with several inflorescence main stems (arrows); I - leaf of wild type; J-M: leaves of transgenic beet plant; J - lobed leaf; K - distorted leaf; L - split leaf with two main veins; M - aberrant leaf with altered color; N - columned inflorescence stem of wild type plant; O-R: transgenic beet plant; O - flattened inflorescence stem; P - columned inflorescence stem with altered color; Q - flattened inflorescence stem with altered color; R - ectopic bud formed from inflorescence stem
like ectopic bud formed from hypocotyl (Fig. 1B), anisomeric cotyledon (Fig. 1C), cup-shaped cotyledon (Fig. 1D), lobed leaf (Fig. 1E), distorted leaf (Fig. 1F), split leaf with two main veins (Fig. 1G) and flattened inflorescence stem (Fig. 1H), etc. Among them, lobed leaf and distorted leaf shared the most percent of 0.96% and 1.62% respectively, and the other phenotypes appeared sporadically. Simultaneously, the other 65 plants showed altered color from claret to yellowish green throughout the entire plant including cotyledons, hypocotyl, leaves, stem, inflorescence stem and flowers. These transformants showed some novel phenotypes such as plant possesses three cotyledons compared to the wild type’s two (Fig. 1I), similar morphology as the wild type except for plant color (Fig. 1J), plants with several columned inflorescence main stems (Fig. 1K), aberrant leaves (Fig. 1L), columned (Fig. 1M) or flattened (Fig. 1N) inflorescence stem, ectopic bud formed from inflorescence stem (Fig. 1O), etc. Among them, color changed plants with normal morphology as the wild type holds the most of 3.18%.

Identification of transgenic lines

Of the above morphological alterations, the interest was on those with changed color for they scarcely appeared in the KNOXI included PttKN1 gene over-expressed plants. In order to confirm the phenotype alterations were induced by the ectopic expression of the PttKN1 gene, PCR, RT-PCR, and PCR-Southern were conducted. As shown in Fig. 2A, PCR assay of 35S-Promoter, NPTII and PttKN1 gene showed specific bands of 192 bp, 142 bp and 311 bp, respectively, while no signal was detected for the wild type. Further analysis of PCR-Southern bands of 192 bp, 142 bp and 311 bp, respectively, and the other phenotypes that had changed color. A: PCR analysis of 35S-Promoter, the 192 bp indicated 35S-Promoter specific band; lane 1, 4, 6, putative transgenic plants; lane 2, 3, 5, wild type plant; lane 1-2, 35S-Promoter, the 192 bp indicated 35S-Promoter specific band; lane 5-6, PttKN1 gene, the 311 bp indicated PttKN1 specific band. B: RT-PCR analysis of PttKN1 gene; lane 1, wild type; 2-5, the color changed plants; the 311 bp indicated PttKN1 specific bands. C: PCR-Southern hybridization of NPTII gene; lane 1-2, the color changed plants. D: PCR-Southern hybridization of PttKN1 gene; lane 1-2, the color changed plants. The pPCV702 vector was regarded as positive control and wild type plant as negative control. PC: positive control; NC negative control.

Pigment contents

The pigments of betacyanins and betaxanthins which have been shown to be a major source of natural colorants in red leaf beet were extracted and measured from the second to fourth true-leaf of wild type and transgenic plants. The results showed that the betacyanins content in transgenic color changed type of 0.15 ± 0.01 mg g⁻¹ was far below compared with the wild type data of 1.71 ± 0.14 mg g⁻¹. Even so, the betaxanthins content in transgenic color changed type and wild type plant had no significant difference of 1.10 ± 0.15 mg g⁻¹ and 1.09 ± 0.22 mg g⁻¹, respectively. The results of t-test showed that the differences of betacyanins between wild type and transgenic lines were very significant (P < 0.05), while the differences of betaxanthins between wild type and transgenic lines were not significant (P < 0.05).

Fig. 2. Molecular assays from the leaves of the putative transgenic plants that had changed color. A: PCR analysis of 35S-Promoter, NPTII and PttKN1 gene; lane 1, 4, 6, putative transgenic plants; lane 2, 3, 5, wild type plant; lane 1-2, 35S-Promoter, the 192 bp indicated 35S-Promoter specific band; lane 5-6, PttKN1 gene, the 311 bp indicated PttKN1 specific band. B: RT-PCR analysis of PttKN1 gene; lane 1, wild type; 2-5, the color changed plants; the 311 bp indicated PttKN1 specific bands. C: PCR-Southern hybridization of NPTII gene; lane 1-2, the color changed plants. D: PCR-Southern hybridization of PttKN1 gene; lane 1-2, the color changed plants. The pPCV702 vector was regarded as positive control and wild type plant as negative control. PC: positive control; NC negative control.
Tyrosinase content

The tyrosinase activity was measured via spectrophotometry with L-dopa as the substrate. The conversion kinetic curve was established with the change rate (ΔA/Δt) of spectro-absorption degree to time that was taken as the reaction rate, and then the tyrosinase activity was calculated (Fig. 3). The results showed that the tyrosinase content in the wild type and transgenic color changed type plant was 5.96 ± 0.62 [×10^6 A / (g·s)] and 1.63 ± 0.31 [×10^6 A / (g·s)], respectively. The tyrosinase level in wild type was about 4 times than it was in transgenic color changed type. The results of t-test showed that the difference of tyrosinase level between wild type and transgenic lines was very significant (P < 0.05).

Discussions

Floral dip method had been used widely for Arabidopsis thaliana transformation (Clough and Bent, 1998; Hay and Tsiantis, 2006) and occasionally for Capsella bursa-pastoris (Bartholmes et al., 2008), Cardamine hirsuta (Hay and Tsiantis, 2006), Medicago truncatula (Curtis, 2004) and Triticum aestivum (Zale et al., 2009) etc. In this report, the method was used to obtain transgenic red leaf beet. The results of kanamycin spraying selection, phenotypes observation and molecular identification indicated that the foreign PttKN1 gene had successfully integrated and expressed into the genome of beet plant. In the past studies, the impact factors of this method on transformation efficiency, like medium composition and additives, had been discussed in detail (Clough and Bent, 1998). Here, the focus was on the roles of dipped time and some additives. It could be seen from Table 1 that the transformation efficiency improved with increased dipped time to 5 min with the highest 2.63% and then reduced. These phenomena told us that sufficient inoculation was critical to improve transformation efficiency, while excessive inoculation might be unfavorable. When surfactant Silwet L-77 or chemotactic acetosyringone were added alone, transformation efficiency showed a little difference with the basic medium used. But when the two reagents were used in pair, transformation efficiency was improved. These results suggested that those additives were not necessary for the success of transformation, but important for enhanced transformation; meanwhile, activation of the host plant and A. tumefaciens shared the same importance to get higher transformation efficiency.

The successful transformation of the floral dip method on red leaf beet led to several novel phenotypes different from their wild type plants. These phenotypes were mainly focused on the ectopic meristem formation, altered stem morphology and altered leaf morphology such as lobed leaf, distorted leaf etc. These morphological modifications were in highly accordance with other PttKN1 transformants (Hu et al., 2005; Xu et al., 2011). Besides these typical phenotypes, the transgenic PttKN1-beet plant displayed changed color throughout the entire plant from claret to yellowish green. This alteration was rare in those PttKN1 gene or KNOXI gene transformants.

The plant coloration of red leaf beet was determinate by the chlorophyll, carotenoid and betalain pigments. As there exists only a slight difference of the chloroplast pigment content between the wild type and transgenic color changed plant, it could not be the reason to make the color changed throughout the entire plant (data not show). So, the changed color could only be caused via the betalain pigments alteration. The sharply decreased betacyanins content in transgenic plant confirmed this supposition. Further studies on tyrosinase content indicated that the betacyanins biosynthesis was partly inhibited in transgenic plants. As tyrosinase had two kinds of activities of hydroxylating activity to form Dopa and oxidizing activity to form cyclo-Dopa (Schliemann et al., 1999), and its activity was measured with L-dopa as the substrate, we could infer that the oxidizing activity of tyrosinase was reduced. Consequently, it could concluded that the ectopic expression of PttKN1 gene in red leaf beet that bring to the changed color from claret to yellowish green might be through regulating the tyrosinase activity. However, the cDNA encoding tyrosinase had not been identified from betalain-producing plants (Tanaka et al., 2008), so how the over-expression of PttKN1 gene acts on the process need further investigation. Hatlestad et al. (2012) provided potential dues as they identified a novel cytochrome P450 that is absolutely required to make betacyanin and without this P450, beets can only make betaxanthin (Hatlestad et al., 2012).

In fact, besides the biosynthesis of betalain pigments could be regulated via the over-expression of PttKN1 gene in red leaf beet, anthocyanins biosynthesis could also be regulated via the PttKN1 gene in transgenic P. hybridra (Hu et al., 2005), B. maculata (Xu et al., 2011) and coleus (Xu et al., 2013). These suggested an interesting hypothesis that the betalain pathway may have co-opted the anthocyanin regulators (Lloyd et al., 2011). Excitingly, analysis of transcriptome data revealed a MYB type transcription factor homologous to the MYBs that could regulate anthocyanin pigments (Lloyd et al., 2011) and activate the biosynthetic genes in the betalain network. Actually, MYB-domain protein was considered the regulator of the KNOX gene expression (Marja et al., 1999). So the expression pattern of MYBs gene induced via the over-expressed PttKN1 gene may be the further investigation.

Conclusions

In conclusion, the present work describes the successful application of floral dip method on the genetic transformation of red leaf beet. The expression of the foreign PttKN1 gene produced pleiotropic morphological typically with the same of other PttKN1 transformants and surprisingly the plant coloration changed from claret to yellowish green. This phenomenon was induced via the partly inhibited tyrosinase activity and further a decreased betacyanin level result from it. But how the tyrosinase activity was regulated by over-expressed PttKN1 gene remains unclear, as the cDNA encoding the enzyme had not been identified from betalain-producing plants. Moreover, the data of PttKN1 gene on regulating anthocyanins biosynthesis suggested an interesting hypothesis of that the betalain pathway may have co-opted the anthocyanin regulators. However, it remains to be investigated whether the PttKN1 gene roles on the two pathways through the interaction with MYB transcription factor.

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References


