Cloning and Expression Analysis of RrMYB113 Gene Related to Anthocyanin Biosynthesis in Rosa rugosa

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Abstract
Anthocyanin is one of water-soluble natural pigments widely existing in flowers, fruits, stems, leaves and seeds of plants, and it is the major factor conferring pink or red to the petals of Rosa rugosa. MYB TFs play an important role in the anthocyanin synthesis in plants. This work aimed to clone the MYB gene related to anthocyanin synthesis in the petals of Rosa rugosa, and explore the relationship between them to lay a good foundation for gene engineering improvement of R. rugosa. Based on the transcriptional data, a full-length cDNA sequence of MYB Gene, RrMYB113 (GenBank accession Nos MG720012), was cloned at the first time from the petals of Rosa rugosa “Zi zhi” with RT-PCR and RACE methods. The full-length cDNA is 885 bp with an open reading frame of 654 bp, encoding 216 amino acids. The derived RrMYB113 protein has a molecular weight of 25,297.64 Da, a calculated pI of 9.61, a R2R3-MYB domain and bHLH binding domain, and it also has the signature motifs ((A/S/G)NDV and KPRPR(T/S)), thus belonging to Sg6 R2R3-MYB subfamily. In the secondary structure of RrMYB113 protein, there is 37.04% α-helix, 39.81% random coil, 14.81% extended peptide chain, and 8.33% β-corner. There is no transmembrane domain and no signal peptide cleavage site, seventeen Ser phosphorylation sites, fifteen Thr phosphorylation sites, four Tyr phosphorylation sites, and no O-glycosylation sites. The expression of RrMYB113 increased with the color deepening in petals, and it expressed at a higher level in petals than in other tissues of Rosa rugosa “Zi zhi”. These results are meaningful to reveal that RrMYB113 might be an important regulator in anthocyanin biosynthesis and coloration in the petals of R. rugosa.

Keywords
Rosa rugosa, Anthocyanin, R2R3-MYB, Gene Expression

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1. Introduction

*Rosa rugosa* is a deciduous shrub of genus *Rosa* in the *Rosacea* family with highly ornamental value, and it plays an important role in landscaping. The flower color of *R. rugosa* is very single, most of which is red, pink, and white, while other colors are rarely seen, which has seriously limited its application in landscaping. Anthocyanin is one of water-soluble natural pigments widely existing in flowers, fruits, stems, leaves and seeds in natural plants [1] [2] [3] [4], and plays an important role in the color of *R. rugosa*. At present, there are few studies on the molecular regulation mechanism of anthocyanin synthesis in *R. rugosa*. Therefore, cloning the MYB TFs from *R. rugosa* related to anthocyanin synthesis is important for understanding the regulation mechanism of anthocyanin accumulation and changing the colors. Anthocyanin is synthesized through the synthetic pathway of flavonoids in phenylpropane pathway, and it is usually catalysed with a series of synthetase and transport proteins, which most have been cloned from model plants [1] [2] [3] [4], and it has been studied extensively in many plants such as *Petunia hybrid* [5], *Zea mays* [6] and *Malus pumila* [7]. Studies show that R_{2}R_{3}-MYB, bHLH and WD40 are three important TFs of regulating anthocyanin synthesis in higher plants, and these TFs play a role by forming a transcriptional complex, which is named MYB-bHLH-WD40 (MBW) [1] [4] [8] [9]. As the most widely used transcription factor in anthocyanin synthesis, R_{2}R_{3}-MYB protein can activate one or more structural genes expression, thereby promoting anthocyanin synthesis [1] [4] [10]-[15]. The *MdMYB1* isolated from *Malus domestica* could induce a large number of anthocyanin to synthetise in cells [16]. The *GhMYB10* isolated from *Gerbara hybrid* is related to the anthocyanin synthesis in petals and leaves, and it can induce the anthocyanin synthesis in pollen sac in the transgenic tobacco [17]. The *VvMYBA1* isolated from *Vitis vinifera* can specifically express in pericarp and could induce anthocyanin biosynthesis [18]. The overexpression of MYB protein encoded by ANT1 in *Lycopersicon esculentum* could activate the expression of CHS, CHI, DFR and other structural genes, thus promoting the anthocyanin synthesis [19].

In this study, we cloned one MYB gene from the petals of *R. rugosa*, and analysed its bioinformatics and expression patterns. These results would provide a theoretical foundation for molecular mechanism of anthocyanin biosynthesis and could be severed as the basis for further comprehension of the pigmentation mechanism in *R. rugosa*.

2. Materials and Methods

2.1. Plant Materials

The plant materials, Chinese representative *Rosa rugosa* “Zi zhi”, “Fen zizhi”, “Bai zizhi”, were from the rose germplasm resources garden at Shandong Agricultural University. *R. rugosa* “Zi zhi” is the most representative traditional rose in China. The stems, leaves, stamens, pistils and petals of these varieties were
collected as samples for expression analysis. All samples were collected directly frozen with liquid nitrogen, and finally stored at −80°C until used.

2.2. Methods

2.2.1. Total RNA Extraction and cDNA Synthesis
An EASY spin Plant RNA Kit from Adlai Biotechnology Co., Ltd. was used to extract the total RNA from the tissue in Section 2.1. Agarose gel electrophoresis and spectrophotometer were used to determine the quality and concentration of the RNA. Abm’s 5× All-In-One RT MsterMix was used to synthesize the first-strand cDNA.

2.2.2. PCR Cloning of Anthocyanin Biosynthesis Related Gene
Based on the related unigene sequences from transcriptome in petals of R. rugosa, the specific primers for the anthocyanin biosynthesis related gene were showed in Table 1, which were designed with Primer Premier 5.0. PCR amplification was conducted using the synthesized cDNA in Section 2.2.1 as a template and the primers in Table 1. The reaction system included 1 µL cDNA, 1 µL F1 primer (10 µmol/L), 1 µL R1 primer (10 µmol/L), and 12.5 µL PCR MIX, with ddH₂O added to a total volume of 25 µL. The reaction conditions were: 94°C for 5 min; 94°C for 30 s, 53°C for 30 s, and 72°C for 1 min for a total of 35 cycles; and then extension at 72°C for 10 min. Next, 1% agarose gel electrophoresis was used to detect the PCR products. The target PCR fragment was recovered with the Hipure Gel Pure DNA Mini Kit (Magen). The recovered fragment was ligated to the pMD18-T vector and then transformed into E. coli DH5a. The positive clones were selected and sent to BGI for sequencing.

2.2.3. Bioinformatics Analysis of Gene
BLASTX (NCBI) was used to study the homology of the nucleotide sequence and the deduced amino acid sequence. DNAMAN5.2.2 was used to conduct multiple sequence alignment. The ORF finder (NCBI) was used to search for an open reading frame, and the Conserved Domains database (NCBI) was used to analyze the conserved domains. ExPaSy-SOPMA was used to predict protein secondary structure. The ProtParam Tool was used to analyze protein physical and chemical properties. Furthermore, the ProtScale was used to predict hydrophilic or hydrophobic protein properties. The NetPhos 3.1 Server was used to predict potential protein phosphorylation sites, and the NetOGlyc 4.0 Server was

Table 1. Primers used to clone and expression analysis of RrMYB113 in R. rugosa.

<table>
<thead>
<tr>
<th>Name</th>
<th>Primer sequence (5’ to 3’)</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>113-F</td>
<td>ATGGAGGTGAGAAAAAGGTACA</td>
<td>Cloning of the</td>
</tr>
<tr>
<td>113-R</td>
<td>TTGCTTGTCTCTCTCTGTAG</td>
<td>Middle Fragment</td>
</tr>
<tr>
<td>113-3’F</td>
<td>CCACGAACTCTTCACCAAAATG</td>
<td>3’RACE PCR</td>
</tr>
<tr>
<td>B26</td>
<td>GACCTCGACATCGAATTTTTTTTTTTTTTTTTTTTTTTTTTT</td>
<td></td>
</tr>
<tr>
<td>RrMYB113-F</td>
<td>ATGGAGGTGAGAAAAAGG</td>
<td>ORF PCR</td>
</tr>
<tr>
<td>RrMYB113-R</td>
<td>TTATTCGTCTCTTCTCTTG</td>
<td></td>
</tr>
</tbody>
</table>
2.2.4. Real-Time Quantitative PCR Analysis

Total RNA extraction and cDNA synthesis were referenced to Section 2.2.1. The expression levels of *RrMYB113* gene involved in anthocyanin biosynthesis were analyzed using quantitative real time PCR. Real time PCR reactions were conducted using two-step PCR System with SYBR Green for detection, using specific primers RrMYB113-Q-F (CCACAGTAATAAGACCTCGA) and RrMYB113-Q-R (GGTGGTGATGTTGATGATG). The reaction volume was comprised of 20 ul containing 10 ul SYBR®Premix Ex Taq™, 0.4 ul primer (RrMYB113-Q-F and RrMYB113-Q-R) and 1 ul cDNA, with ddH₂O added to a total volume of 20 µL. The reaction conditions were as follows: pre-heating at 94˚C for 5 min; 39 cycles at 95˚C for 10 s, at 60˚C for 30 s. Signals were monitored by the Chromo3 real-time PCR system, finally 30 s at 60˚C and 30 s at 95˚C for the melting curve. The cycle threshold (Ct) value for each PCR reaction was calculated. After completion of the amplification steps, the melting curve was determined for each analysis. Gene transcripts were quantified using the comparative Ct method, which compares the transcript level of the target gene with that of the reference gene.

3. Results and Analysis

3.1. Cloning and Sequence Analysis of *RrMYB113* Gene

One R2R3-MYB transcription factor, *RrMYB113* (GenBank accession number: MG720012), was cloned from the petals of *Rosa rugosa*, and the blast analysis confirmed that all its homologous genes were R2R3-MYB TFs. The cloned middle fragment is 625 bp, the cloned 3'-terminal fragment is 498 bp. These two fragments were spliced together with DNAstar in order to obtain an 885 bp cDNA sequence and the ORF is 651 bp, encoding a polypeptide of 216 amino acids (Figure 1).

Amino acid sequence alignment between *RrMYB113* and other MYB TFs with

![Figure 1. PCR amplification of *RrMYB113*. M: Marker; C1, C2: Full-length fragment; B1, B2: Intermediate fragment; A1, A2: 3'-RACE.](image-url)
higher homology revealed that RrMYB113 consisted of both R2 and R3 DNA-binding domains. Besides, the alignment showed that the bHLH motif, which interacted with bHLH proteins, appeared in the R3 domain. What’s more, RrMYB113 had the signature motifs \([(A/S/G)NDV\) and \(KPRPR(T/S)\)] of Sg6 R R3-MYB subfamily (Figure 2).

In order to study the evolutionary relationship between RrMYB113 and MYB TFs protein in other species, the evolution tree was constructed and analyzed by BLAST with 10 species with homology from high to low. The evolution tree was constructed by MEGA5.0 software, and the system evolution tree was tested by bootstrap, which was repeated 1000 times. The results showed that RrMYB113 was closely related to the members belonging to Rosaceae family, such as Rubus idaeus, Rubus hybrid, and so on, while it was relatively distant from other MYBs in different families. In addition, eight MYB TFs such as RrMYB113 and PaMYB90 were clustered into one branch, and EjMYB10 and PpMYB10 gathered into another branch (Figure 3).

Figure 2. Multiple alignment of the RrMYB113 with other MYB TFs. Notes: The red line indicate the conserved R2-domain and R3-domain, the black line indicate the conserved residuals interacting with bHLH proteins. Box (A) a conserved motif of [A/S/G]NDV, box (B) a conserved motif of KPRPR (T/S).

Figure 3. The phylogenetic tree derived from the alignment of amino acid sequences of RrMYB113 and other MYB TFs.
3.2. Bioinformatics Analysis of RrMYB113 Gene

The RrMYB113 protein encoded 216 amino acids, 35 basic amino acids (Arg + Lys), 27 acid amino acids (Asp + Glu), and 154 neutral amino acids, and the prediction molecular formula was $C_{1109}H_{1747}N_{333}O_{329}S_9$. The derived protein had a molecular weight of 25,297.64 Da, a calculated $pI$ of 9.61. It belonged to the unstable protein with an unstable index at 59.15, and it was also a hydrophilic protein with the total average hydrophobic index at $-0.906$. The secondary structure prediction result demonstrated that there were 37.04% α-helix, 39.81% random coil, 14.81% extended peptide chain, and 8.33% β-corner. The phosphorylation site prediction results demonstrated that there were 17 Ser phosphorylation sites, 15 Thr phosphorylation sites, 4 Tyr phosphorylation sites, and no O-glycosylation sites.

3.3. Expression Patterns of RrMYB113 in Different Tissues and Different Varieties

The expression analysis of RrMYB113 in different tissues showed that RrMYB113 expressed differentially among stems, leaves, stamens, sepals, pistils, and petals. RrMYB113 was more abundant in petals than stems, leaves, stamens, sepals and pistils. The highest expression level of RrMYB113 was observed in petals, while it expressed slightly in pistil, stamen and leaves, and almost didn’t express in sepals and stems. In addition, the results showed that the expression of RrMYB113 increased with the color deepening among the three cultivars, highest in R. rugosa “Zi zhi”, followed by R. rugosa “Fen zizhi”, and the lowest in R. rugosa “Bai zizhi” (Figure 4).

4. Discussion

In this study, a MYB gene named RrMYB113 has been isolated from R. rugosa.
The amino acid sequence alignment showed that *RrMYB113* contained R2 and R3 DNA-binding domains, and it also had a bHLH interaction motif in the R3 domain, which provided corresponding binding sites for the formation of the three element complex (MYB-bHLH-WD40) [1] [6]. Furthermore, it had the signature motifs ((A/S/G)NDV and KPRPR(T/S)), and belonged to Sg6 R2R3-MYB subfamily. Previous studies show that the R2R3-MYB proteins of the Sg6 subfamily are mainly involved in the regulation of the synthesis and accumulation of anthocyanins in plants [4]. Many R2R3-MYB TFs are known to control anthocyanin biosynthesis by regulating structural genes in the anthocyanin pathway [14] [22] [23]. At present, R2R3-MYB TFs of Sg6 subfamily have been cloned from *Rosa chinensis*, *Lycopersicon esculentum*, *Dioscorea esculenta*, *Malus domestica*, *Citrus sinensis*, and so on [12] [14] [24] [25]. In several other plant species, the expression of many R2R3-MYB TFs in the anthocyanin pathway is strongly correlated with anthocyanin accumulation. For example, *MdMYB10* express highly in red-fleshed apple, but is virtually undetectable in the white-fleshed apple [26]. Evolutionary analysis showed that *RrMYB113* was highly homologous to the MYB TFs of the Sg6 subfamily in other species. Therefore, it’s conjectured that the *RrMYB113* gene was related to anthocyanin synthesis.

Through the bioinformatics analysis, we found that the alpha helix and random coil accounted for a considerable proportion in the secondary structure of *RrMYB113* protein, while the extended strand and beta turn occupied small percentage. A previous study has reported that the alpha helix plays an important role in R motif of the MYB domain, and each R motif is generally composed of three alpha helices, and the second and third R motif form a HTH structure and then combine with the first R motif, further forming a HTH domain with a hydrophobic core. What’s more, the third alpha helices in R motif has a role of identifying DNA, so that the MYB protein has high specificity. Therefore, it was predicted that the *RrMYB113* gene belonged to the R2R3-MYB [9]. Besides, the random coil is beneficial to the combination of cells with water, and *RrMYB113* belongs to the hydrophilic protein, so we presumed that *RrMYB113* gene may play a protective role in osmotic stress of plants [27].

The results of Real-time quantitative PCR showed that the expression of *RrMYB113* gene exhibited a decreasing trend in the petals of *R. rugosa* “Zi zhi”, *R. rugosa* “Fen zizhi” and *R. rugosa* “Bai zizhi”, and in the expression of different tissues in *R. rugosa* “Zi zhi”, the *RrMYB113* gene highly expressed in petals, while in a very low level in other tissues. Previous studies indicate that there are positive and negative mechanisms of MYB protein on anthocyanin regulation in plants [9]. For example, in apples, the *MdMYB1* is positively related to anthocyanin synthesis, and it is regulated by light. And overexpression of *MdMYB10* which cloned from leaf and pulp could increase the accumulation of anthocyanin in seedlings, while overexpression of *MdMYB16*, *MdMYB17* and *MdMYB111* in tobacco could inhibit the activity of DFR promoter, and then influence the
anthocyanin synthesis [16] [28]. And the FaMYB10 gene isolated from *Fragaria × ananassa* is similar to *MdMYB10*, which could promote the anthocyanins synthesis, while the FaMYB1 gene exhibit oppositely [15]. In the present study, the expression level of RrMYB113 increases with the color deepening, and it's highest expressed in the petals of *R. rugosa* “Zi zhi” in different tissues. Therefore, the author believed that the RrMYB113 gene positively regulate the anthocyanin synthesis in *R. rugosa*.

5. Conclusion

In conclusion, one R2R3-MYB TF, RrMYB113, was isolated from *R. rugosa* and was found to be involved in regulating anthocyanin biosynthetic pathway. The results of this study provided important information on the anthocyanin synthesis of *R. rugosa*. In future work, we will test whether the overexpression of RrMYB113 leads to anthocyanin accumulation in *Arabidopsis thaliana* and *Nicotiana tabacum*.

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References


