Identification of a Novel DFR-A Mutant Allele Determining the Bulb Color Difference between Red and Yellow Onions (Allium cepa L.)

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ABSTRACT To introduce downy mildew resistance from a yellow-colored resistant cultivar, ‘Santero’, into a yellow breeding line, OT803, the F₁ hybrid was produced by crossing Santero and OT803. The bulb color of the F₁ hybrids became light pink, suggesting involvement of complementation between the DFR-A and ANS genes in the onion anthocyanin biosynthesis pathway. Since Santero contained active DFR-A and inactive ANS alleles, OT803 was assumed to harbor active ANS and inactive DFR-A alleles. However, some yellow-colored individuals of OT803 were shown to contain the homozygous genotype of the active DFR-AR₄-like allele. Sequencing of 4,830-bp full-length sequences of this DFR-AR₄-like allele revealed that the nucleotide sequences of the DFR-AR₄ and DFR-AP₅-like alleles were identical except for a single nucleotide deletion in the last exon. This single base-pair deletion resulted in creation of a premature stop codon at 2-bp downstream of the deletion mutation. This new DFR-A mutant allele was designated DFR-APS₂. The RT-PCR results showed that transcripts of the DFR-APS₂ allele were significantly reduced, suggesting involvement of nonsense-mediated mRNA decay (NMD) mechanism. The systematic process consisting of PCR amplification and sequencing of the PCR products was modified to identify the DFR-APS₂ allele among 16 different DFR-A alleles. No additional accession was found to contain the DFR-APS₂ allele from 155 diverse onion germplasm, indicating very limited distribution of this new DFR-APS₂ allele.

Keywords Onion, Allium cepa, Bulb color, Dihydroflavonol 4-reductase (DFR), Molecular marker

INTRODUCTION

Bulb color is one of major traits in onion (Allium cepa L.) breeding programs (Brewster 1994), and diverse bulb colors such as red, yellow, white, and chartreuse have been reported (El-Shafie and Davis 1967). Flavonoid compounds are known to confer bulb colors in onions (Fossen et al. 1996). Flavonoid is one of major secondary metabolites, and more than 8,000 derivatives have been reported in plant species (Veitch and Grayer 2011). Flavonoid plays such diverse roles as protectants against UV radiation and pathogens in plants (Shirley 1996; Fini et al. 2011). In addition, as human diet, it has health-promoting effects as potent antioxidant and anticancer agents (Cook and Samman 1996; Lotito and Frei 2006; Clere et al. 2011; Cushnie and Lamb 2011; Nishiumi et al. 2011; Russo et al. 2012).

Anthocyanin, one of major flavonoid compounds, is responsible for the red bulb color, and quercetin derivatives are predominant in yellow onions (Fossen et al. 1996; Rhodes and Price 1996; Slimestad et al. 2007). Inheritance studies (Reiman 1931; Clarke et al. 1944; El-Shafie and Davis 1967) reported that five major loci were involved in determination of onion bulb colors. The I locus, known as a color inhibiting factor, determines the white bulb color when its genotype is homozygous dominant. Meanwhile,
the C locus, known as a basic color factor, also determines the white bulb color only when its genotype is homozygous recessive. When the genotype of the G locus is homozygous recessive, the chartreuse bulb color appears. The R and L loci are complementarily involved in determination of color difference between yellow and red bulb colors.

These five loci was assumed to be related with genes involved in anthocyanin biosynthesis pathway (Kim et al. 2004a). Structural genes encoding enzymes in the anthocyanin biosynthesis pathway have been well characterized in many plant species (Goodrich et al. 1992; Quattrocchio et al. 1993; Holton and Cornish 1995; Spelt et al. 2000; Yamazaki et al. 2003). In addition, regulatory genes controlling transcription of structural genes in the pathway have also been isolated in some model species such as Arabidopsis, petunia, and maize. The ternary complex consisting of R2R3-MYB, bHLH, and WD40 transcription factors is commonly identified as major regulatory genes (Ramsay and Glover 2005; Petroni and Tonelli 2011; Czemmel et al. 2012).

The R and L loci were revealed to encode dihydroflavonol 4-reductase (DFR) and anthocyanidin synthase (ANS) enzymes, respectively, in the onion anthocyanin biosynthesis pathway (Kim et al. 2004a, 2005a). One leaky and three inactive ANS alleles were identified from diverse onion germplasm (Kim et al. 2004b, 2005a, 2016). Recently, integration of a non-autonomous DNA transposon in the promoter region was revealed to cause significant reduction of the ANS gene transcription (Kim et al. 2015). Three homologous genes coding for the DFR enzyme were isolated in the previous study (Kim et al. 2005b), and they were designated DFR-A, DFR-B, and DFR-C. Among them, only the DFR-A gene was active and determined the bulb color phenotypes (Kim et al. 2005b). Since the first inactive DFR-A allele was identified (Kim et al. 2005b), eleven mutant alleles containing diverse insertion, deletion, or substitution mutations have been identified (Kim et al. 2009, 2015; Song et al. 2014). Three of them harbored transposable elements including a LTR retrotransposon and two different DNA transposons (Kim et al. 2015).

In this study, a novel DFR-A allele containing a premature stop codon, which was created by one base-pair deletion, was identified from one breeding line. The inactivation mechanism of this mutant allele was proposed, and a systematic process for identification of 16 different DFR-A alleles was updated to include the newly identified DFR-A allele.

MATERIALS AND METHODS

Plant materials

A F1 population originating from the cross between a downy mildew resistant cultivar, ‘Santero’, and a breeding line (OT803) was used to identify a new DFR-A allele. To assess the frequency of the new DFR-A allele, total genomic DNAs of 18 accessions containing the DFR-AR4 allele were analyzed (Supplementary Table 1). These 18 accessions were selected from 155 accessions collected from diverse countries in the previous study (Song et al. 2014). Individual plants of OT803 which contained the homozygous genotype of the new mutant DFR-A allele and a red-colored breeding line, H6, were used for RT-PCR amplification of the DFR-A gene.

DNA extraction, PCR amplification, and sequencing of PCR products

Total genomic DNAs were extracted from leaves of seedling or sprouted bulbs using a cetyl trimethylammonium bromide (CTAB) method (Doyle and Doyle 1987). PCR was performed in 25-μL reaction mixtures containing 0.05 μg template, 2.5 μL 1× PCR buffer, 0.2 μL forward primer (10 μM), 0.2 μL reverse primer (10 μM), 0.2 μL dNTPs (10 mM each), and 0.25 μL polymerase mix (Advantage 2 Polymerase Mix; Clontech, Palo Alto, CA, U.S.A.). The primer sequences used in this study are listed in Table 1.

PCR amplification was performed with an initial denaturation step at 94°C for 5 minutes, followed by 40 cycles of 94°C for 30 seconds, 65°C for 30 seconds, 72°C for 3 minutes, and a final 10 minutes extension at 72°C. The PCR products were visualized on a 1.5% agarose gel after ethidium bromide staining. For sequencing of PCR products, they were first purified using a QIAquick PCR Purification kit (QIAGEN, Valencia, CA, U.S.A.), and
Table 1. Primer sequences used in this study.

| Name     | Sequence (5’ to 3’) | Application                          | Reference              |
|----------|---------------------|--------------------------------------|------------------------|
| DFR-F    | ATGCCAGTGGAGTGCATGTTGAATGGT | DFR-A allele identification           | Kim et al. (2005b)     |
| DFR-R    | TGGGTAAGCGATTGTTCTATCTCTTCA   | DFR-A allele identification and RT-PCR| Kim et al. (2005b)     |
| DFR-LR3  | TTGCAAACTCCCATGCAGCCTTCTCTG  | DFR-A allele identification           | Song et al. (2014)     |
| DFR-LR4  | TTAACCATCTGCCCAAAAT       | DFR-A allele identification           | This study             |
| DFR-LF5  | GAGGACACCAAAAAGCCGAATACGAT | RT-PCR                               | Song et al. (2014)     |
| Tubulin-F| CTGGGAGCTTTAAGCCTTTG       | RT-PCR                               | Song et al. (2014)     |
| Tubulin-R| CAAGGGGACCCTGCAAAATA       | RT-PCR                               | Song et al. (2014)     |
| CHS-F    | GAGGCTACGCCACGGTGTTAGCTG   | RT-PCR                               | Kim et al. (2004b)     |
| CHS-R    | ATCAATGGCCACACTCCTAAGCACC  | RT-PCR                               | Kim et al. (2004b)     |
| F3H-F    | GAAGACGAGCGTCCAAGATGCTG    | RT-PCR                               | Kim et al. (2004b)     |
| F3H-R    | TCCACCCTTTTAGTGCTTCAGAC    | RT-PCR                               | Kim et al. (2004b)     |

Sequencing reactions were performed by a specialized company (Macrogen, Seoul, Republic of Korea). The full-length sequences of the new DFR-A allele were obtained using the method described in the previous study (Song et al. 2014).

RNA extraction, cDNA synthesis, and RT-PCR

Total RNAs were extracted from fresh leaf sheaths of four-leaf stage seedlings using a RNA extraction kit (RNeasy Plant Mini Kit, Qiagen) following the manufacturer’s instructions. cDNAs were synthesized using a commercial cDNA synthesis kit (SuperScript™ III first-strand synthesis system for RT-PCR; Invitrogen, Carlsbad, CA, U.S.A.). RT-PCR amplification was performed with an initial denaturation step at 94°C for 3 minutes followed by 30 cycles of 94°C for 30 seconds, 65°C for 30 seconds, and 72°C for 3 minutes, and a final 10-minute extension at 72°C. The primer pair used in RT-PCR is listed in Table 1. The onion tubulin sequence obtained from EST sequences (TC125) from the DFCI Allium cepa Gene Index (Antonescu et al. 2010) was used as a control.

RESULTS

Identification of a novel DFR-A mutant allele determining the yellow bulb color in onion

In order to introduce downy mildew resistance from a resistant cultivar, Santero, into a breeding line (OT803), the F1 hybrids were produced by the cross between Santero and OT803. Although the bulb color of both parental lines were yellow, the bulb color of the F1 hybrids became light pink (Fig. 1). The light pink bulb color was assumed to appear by complementation between the DFR-A and ANS genes involved in the anthocyanin biosynthesis pathway as shown in the previous study (Kim et al. 2005a). Since Santero contained an active DFR-AR allele and inactive \( \text{ANS}^{\text{R}} \) and \( \text{ANS}^{\text{S186L}} \) alleles (Kim et al. 2016), OT803 was
Table 2. Composition of DFR-A and ANS alleles of parental lines (Santero and OT803) and the F1 hybrid. The active DFR-A and ANS alleles are shown in boldfaces. The DFR-A $^{K4}$-like allele is designated as DFR-APS$^2$ allele.

| Population | Genotype                          | DFR-A | ANS             |
|------------|-----------------------------------|-------|-----------------|
| Santero    |                                   | $DFR-A^{K3}$ / $DFR-A^{K4}$  | $ANS^{P5}$ / $ANS^{S18}$ |
| OT803      | Mixed with homozygous and heterozygous $DFR-A^{P5}$ and $DFR-A^{T4}$ |       |                 |
| F1 hybrid  |                                   | $DFR-A^{P5}$ / $DFR-A^{K4}$  | $ANS^{P5}$ / $ANS^{S18}$ |

Fig. 2. Organization of the inactive mutant DFR-A alleles isolated from onions. Arrow-shaped boxes indicate coding regions and the 5′-to-3′ direction. The empty and gray boxes in the coding regions indicate exons and introns, respectively. The vertical filled arrows on the exons indicate the positions of sequence changes. The nucleotide sequences on the vertical empty arrow indicate the normal sequences of the functional alleles. The dotted lines in the promoter region indicate the deleted sequences. The dotted line outside the DFR-ADEL allele indicate entire deletion of the genic region. Nucleotide sequences and names on the inverted triangles indicate inserted sequences and transposable elements, respectively. Lengths of transposable elements are shown in parenthesis.
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Fig. 3. Alignment of partial nucleotide sequences of the DFR-A<sup>R4</sup> and DFR-A<sup>P52</sup> alleles. The vertical filled arrow indicates the position of single base-pair deletion. The rectangular box indicates the position of the premature stop codon.

Fig. 4. RT-PCR amplification of the DFR-A gene. 1-5: individuals of OT803 containing the homozygous DFR-A<sup>P52</sup> allele. 6-10: individuals of H6 containing the homozygous DFR-A<sup>R4</sup> allele. CHS-A: chalcone synthase-A, F3H: flavanone 3-hydroxylase

biosynthesis pathway were not significantly different between OT803 and H6 individuals (Fig. 4). Hereafter, this novel mutant allele is designated DFR-A<sup>P52</sup>.

**Frequency of the DFR-A<sup>P52</sup> allele in diverse onion germplasm**

For identification of specific DFR-A alleles from diverse onion germplasm, a systematic process consisting of PCR amplification and sequencing of the PCR products was devised for identification of 15 different DFR-A alleles in the previous studies (Song et al. 2014; Kim et al. 2015). For identification of the DFR-A<sup>P52</sup> allele, one additional step of sequencing of the PCR product using the DFR-LR4 primer was added in the process to distinguish the DFR-A<sup>R4</sup> and DFR-A<sup>P52</sup> alleles (Fig. 5).

In the previous study (Song et al. 2014), the DFR-A genotypes of 155 onion accessions originating from diverse countries were identified by using the process for identification of 15 DFR-A alleles. However, at that time, the DFR-A<sup>P52</sup> allele could not be distinguished from the DFR-A<sup>R4</sup> allele by such process since the position of the 1-bp deletion in the DFR-A<sup>P52</sup> allele could not be sequenced by the DFR-LR3 primer (Fig. 5). Therefore, PCR products of 18 accessions containing the DFR-A<sup>R4</sup> allele were further sequenced by the DFR-LR4 primer following the updated process (Fig. 5). However, sequencing results showed that no accession contained the DFR-A<sup>P52</sup> allele, indicating very limited distribution of the DFR-A<sup>P52</sup> allele in onion germplasm.
Fig. 5. Flowchart showing the process for identification of 16 DFR-A alleles. The SNPs are shown with numbers indicating the distances from the start codon of the DFR-A gene. Four active and 12 inactive DFR-A alleles are shown in filled and empty oval-shaped circles, respectively.

DISCUSSION

A novel inactive DFR-A allele containing a premature stop codon in the last exon was identified in this study. This is the 12th inactive DFR-A allele identified from onion germplasm. In the previous studies (Kim et al. 2005b, 2009, 2015; Song et al. 2014), eleven inactive DFR-A alleles containing diverse insertion, deletion, or substitution mutations were identified (Fig. 2). Unlike other structural and regulatory genes involved in the onion anthocyanin biosynthesis pathway, the DFR-A gene unusually has a large number of mutant alleles. No inactive mutant alleles of other structural and regulatory genes affecting color phenotypes have been identified in onions except for three ANS (Kim et al. 2005a, 2016) and one CHI (Kim et al. 2004c) mutant alleles. Although presence of multiple homologous DFR genes in the onion genome is likely to contribute to creation of a multitude of DFR-A mutants, the exact mechanism responsible for frequent mutation of the DFR-A gene is not resolved yet. Khar et al. (2008) showed that numerous copies of DFR homologs existed in the onion genome by Southern blot analysis.

The DFR-A^{PS2} allele is probably derived from the DFR-A^{R4} allele recently, since there is only a single nucleotide deletion between 4,830-bp promoter and coding regions of the DFR-A^{R4} and DFR-A^{PS2} alleles. The single nucleotide deletion changes the reading frame and create a premature stop codon. However, loss of the last 60 residues, which is caused by the premature stop codon, may not have a significant effect on the function of the DFR enzyme, since the premature stop codon is positioned at the last exon. Indeed, none of the catalytic triad and 28 residues...
involved in binding of substrate and NADPH is positioned in the last exon (Petit et al. 2007).

However, the RT-PCR result showed that the transcripts of the DFR-APS2 allele were significantly reduced, and the PCR products appeared as smear bands compared with the normal DFR-AR1 allele. Since there is no polymorphism in the 2.5-kb upstream region containing putative promoter regions between the DFR-AR4 and DFR-APS2 alleles, transcription of the DFR-APS2 allele is assumed to be normal. Therefore, the NMD mechanism is most likely to be responsible for degradation of the DFR-APS2 mRNA transcripts. The NMD mechanism, which is well conserved in all eukaryote, plays an important role in preventing deleterious effects of truncated proteins by degradation of defective mRNAs harboring premature stop codons (Hentze and Kulozik 1999; Neu-Yilik et al. 2004; Chang et al. 2007; Shaul 2015).

Although the causal genes responsible for the bulb color difference between red and yellow onions have been revealed as the genes coding for DFR and ANS enzymes (Kim et al. 2004a, 2005a), designing of universal functional markers based on mutations is not a straightforward process due to a high level of allelic variation of the DFR-A gene. In addition, complementation between the DFR-A and ANS genes makes it more complicated to identify specific DFR-A alleles of yellow onion germplasm. Therefore, systematic processes are first required for identification of specific alleles of the DFR-A and ANS genes. Once specific DFR-A or ANS alleles of parental lines in segregating populations are determined, functional markers can be designed on the basis of polymorphism between two alleles. We updated the process for identifying 16 DFR-A alleles in this study. Combined with the process for ANS allele identification (Kim et al. 2016), the updated process for DFR-A allele identification becomes an essential tool in onion breeding programs.

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