SCIENTIFIC REPORTS

Received: 27 February 2015 Accepted: 22 June 2015 Published: 20 July 2015

OPEN The Balance of Expression of Dihydroflavonol 4-reductase and **Flavonol Synthase Regulates** Flavonoid Biosynthesis and Red **Foliage Coloration in Crabapples**

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Red leaf color is an attractive trait of Malus families, including crabapple (Malus spp.); however, little is known about the molecular mechanisms that regulate the coloration. Dihydroflavonols are intermediates in the production of both colored anthocyanins and colorless flavonols, and this current study focused on the gene expression balance involved in the relative accumulation of these compounds in crabapple leaves. Levels of anthocyanins and the transcript abundances of the anthocyanin biosynthetic gene, dihydroflavonol 4-reductase (McDFR) and the flavonol biosynthetic gene, flavonol synthase (McFLS), were assessed during the leaf development in two crabapple cultivars, 'Royalty' and 'Flame'. The concentrations of anthocyanins and flavonols correlated with leaf color and we propose that the expression of McDFR and McFLS influences their accumulation. Further studies showed that overexpression of McDFR, or silencing of McFLS, increased anthocyanin production, resulting in red-leaf and red fruit peel phenotypes. Conversely, elevated flavonol production and green phenotypes in crabapple leaves and apple peel were observed when McFLS was overexpressed or McDFR was silenced. These results suggest that the relative activities of McDFR and McFLS are important determinants of the red color of crabapple leaves, via the regulation of the metabolic fate of substrates that these enzymes have in common.

Flavonoids are a class of plant secondary metabolites that collectively have diverse functions, including providing protection against abiotic stresses, particularly UV-irradiation, and biotic factors, such as phytophagous insects, as well as attracting pollinators^{1,2}. They also have considerable value as components of the human diet³ and are used in the pharmaceutical industry since they have antioxidative, anticancer and anti-inflammatory properties^{4,5}. Flavonoids have been extensively studied in a range of land plants and have been divided into nine structural subclasses: chalcones, flavones, flavonols, dihydroflavonols, flavandiols, anthocyanins, proanthocyanidins, flavonoid isoflavonoids and aurones⁵. Moreover, many genes involved in the flavonoid biosynthetic pathway have been identified and characterized. For example, the genes encoding chalcone synthase (CHS), chalcone isomerase (CHI), F3H flavanone 3-hydroxylase (F3H), flavonoid 3'-monooxygenase (F3'H), dihydroflavonol 4-reductase (DFR), anthocyanidin synthase (ANS), uridine diphosphate glucose-flavonoid 3-O-glucosyl transferase (UFGT), flavonol synthase (FLS), leucoanthocyanidin reductase (LAR) and anthocyanidin reductase (ANR) have been characterized in several plant species⁶⁻¹⁴.

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Figure 1. Diagrammatic representation of the flavonoid biosynthetic pathway in plant. Enzymes required for flavonol and anthocyanin synthesis: CHS, chalcone synthase; CHI, chalcone isomerase; F3H, flavanone 3-hydroxylase; FLS, flavonol synthase; DFR, dihydroflavonol 4-reductase; F'3H, flavonoid 3'-monooxygenase; ANS, anthocyanidin synthase; GTs, glucosyltransferases; ATs, anthocyanin aromatic acyltransferases.

The flavonoid biosynthetic pathway lies downstream of the phenylpropanoid pathway and leads to the formation of anthocyanins and flavonols (Fig. 1). Chalcone synthase (CHS; EC 2.3.1.74) uses beta-coumaroyl-CoA and 3 Malonyl-CoA as substrates to form naringenin chalcone¹⁵. This condensation reaction is a key step in the pathway leading to the formation of flavonoids¹⁶. Next, the F3H converts naringenin to dihydrokaempferol. Then dihydrokaempferol and dihydroquercetin are converted to kaempferol and quercetin by FLS, respectively. DFR and FLS also catalyze competing reactions to generate products leading to the spectrum of downstream anthocyanins and flavonols^{17,18}.

MYB family transcriptional factors are known to be involved in regulating the expression expression of flavonoid biosynthesis genes¹⁹. For example, in *Arabidopsis thaliana*, AtMYBL2 interacts with TT8 (TRANSPARENT TESTA 8) to reduce anthocyanin biosynthesis²⁰. In the context of flavonoid biosynthesis, AtMYB12 is thought to effect flavonol production^{19,21} by regulation *CHS*, *CHI*, *F3H* and *FLS1* gene expression levels²². In addition, the expression of three allelic apple (*Malus × domestic*) genes (*MYB10*, *MYB1* and *MYBA*) was shown to strongly correlate with the accumulation of anthocyanins in fruit^{23–25}.

The pigmentation of many plant organs results from the presence of some of these flavonoid classes and, in many instances, a primary determinant of color is the accumulation of anthocyanins. The synthesis of anthocyanins occurs via a key branch in the flavonoid biosynthesis pathway, involving the action of the enzyme DFR on the dihydroflavonols dihydroquercetin, dihydrokaempferol or dihydromyricetin. Indeed, *DFR* genes have been shown to play an important role in determining the total anthocyanin content in *A. thaliana*^{26,27}. Additionally, the expression level of the *DFR* gene *TfDFR1* has been shown to be positively correlated with red pigment accumulation in the petals of tulip (*Tulipa fosteriana*)²⁸. It has also been reported that the expression level of *DFR* positively correlates with the abundance of anthocyanins in peanut (*Arachis hypogaea*)²⁹.

Flavonols provide another important co-pigment in the colorful organs of terrestrial plants, such as the yellow petals of *Lathyrus chrysanthus*³⁰, and they also influence pollen tube growth³¹. Flavonols are derived from 2, 3-dihydroquercetin and their formation is catalyzed by FLS, which belongs to the 2-oxoglutarate dependent dioxygenase family^{4,32}. Following the identification of an *FLS* gene from *Petunia hybrida*³³, homologous *FLS* genes have been identified from *A. thaliana, Solanum tuberosum, Matthiola incana, Malus domestica* and *Eustoma russellianum*. In addition, it has been reported that increases in transcript levels of an *FLS* gene from satsuma mandarin (*Citrus unshiu* Marc.), *CitFLS*, in the fruit peel correlate with flavonol accumulation during fruit development³⁴. Recently, it has been found that major floral color changes are a consequence of *FLS* expression in petunia (*Petunia hybrida Vilm.*), *Lisianthus (Eustoma grandiflorum*) and camellia (*C. nitidissima*)^{33,35-37}. In contrast, an indirect effect of a camellia *FLS* gene (*CnFLS1*) on anthocyanin accumulation during floral coloration was suggested following

an experiment where its overexpression in transgenic tobacco (*N. benthamiana*) plants resulted in an increase in flavonol content, but a reduction in anthocyanin levels in petals³⁷.

Leaf color is a key determinant of the commercial value of many ornamental plant species; however, much remains to be learnt about the mechanisms of color formation in leaves at the molecular level. The study of pigmentation mechanisms in leaves is therefore significant for both breeding and genetic engineering of ornamental plants. An example of an important ornamental woody plant is *Malus* crabapple, which belongs to the Rosaceae, *Malus Mill* family. The numerous plant landscape species in this family provide an excellent source of research material for studying the mechanism of color formation and accumulation, due to their colorful leaves, flowers and fruits³⁸. To date, little is known about the mechanism of anthocyanin and flavonol biosynthesis in ornamental crabapples.

In this current study, we investigated the function of the crabapple DFR (*McDFR*) and FLS (*McFLS*) genes in regulating leaf color in different cultivars. We overexpressed and silenced each gene to determine their interaction in controlling flavonol and anthocyanin biosynthesis, and evaluated the gene expression ratio of *McDFR* and *McFLS* that is required for leaf color production. We also discuss the metabolic flux between McDFR and McFLS during flavonoid biosynthesis in leaves and fruit. We propose that the finding from this study will assist future attempts to enhance anthocyanin or flavonol accumulation in the leaves of target ornamental species by altering the balance between the McDFR and McFLS enzyme activities.

Results

The anthocyanin and flavonol content of the leaves of two crabapple cultivars. Two extreme leaf color cultivars, 'Royalty' and 'Flame', have ever-red and ever-green leaves, respectively. We evaluated the abundance of anthocyanins and flavonols in the leaves of these cultivars at 5 development stages of the crabapple leaf growing season by high-performance liquid chromatography (HPLC) (Fig. 2). The chromatography results showed that cyanidin 3-O-glucoside was the predominant anthocyanin, and we found that that the major flavonols were quercetin derived compounds, such as quercetin 3-O-diglucoside, quercetin 3-O-glucoside and quercetin 3-O-glycosidase isomer (Fig. 2B). As shown in Fig. 2C, anthocyanin levels in the ever-red leaves of 'Royalty' were significantly higher than those in the ever-green leaves of 'Flame'. A gradual decrease in anthocyanin content was observed in 'Royalty' leaves during their development, while anthocyanins were almost undetectable in 'Flame' leaves. In contrast, the abundance of flavonols increased during the development of 'Flame' leaves, except at stage 5.

The expression levels of anthocyanin and flavonol biosynthetic genes in the leaves of the two crabapple cultivars. To gain insight into the relationship between the expression patterns of anthocyanin biosynthetic genes and anthocyanin/flavonol accumulation, the transcript levels of the key anthocyanin biosynthetic gene *McDFR*, and the flavonol biosynthesis gene *McFLS*, were determined by qRT-PCR. The expression of *McDFR* decreased during the development of both ever-red and ever-green leaves (Fig. 3). In contrast, the transcript levels of *McFLS* showed an increase in the early developmental stages, and then decreased in stages 4 and 5 in ever-red leaves (Fig. 2A), while only a slight decrease was seen in the last stage of the ever-green leaves (Fig. 3B). This suggested that *McDFR* expression positively correlated with anthocyanin accumulation during the development of ever-red leaves, and that *McFLS* expression and flavonol accumulation were positively correlated in leaves of both cultivars.

McDFR plays an important role in the anthocyanin biosynthetic pathway. To further investigate the function of *McDFR*, we suppressed its expression in the leaves of the red-leafed crabapple cultivar 'Royalty' by virus-induced gene silencing (VIGS) using the TRV vector³⁹. Leaves infected with the virus containing the TRV-*McDFR* construct began to exhibit a green color at 14 days post-infection (dpi), while a more severe phenotype was detected in the new buds of TRV-*McDFR* infected stem tips at 35 dpi (Fig. 4A). We next analyzed the variation in flavonoid content of the infected leaves by HPLC and observed that in the *McDFR* silenced areas of the leaves; the levels of anthocyanin were much lower than in non-silenced leaves, while the flavonol content showed a significant increase (Fig. 4B). We also confirmed, by qRT-PCR analysis, that the abundance of endogenous *McDFR* transcripts was greatly reduced in TRV-*McDFR* infected leaves. The expression levels of other flavonoid biosynthetic genes, namely *CHS* (*McCHS*) and *F3'H* (*McF3'H*) were similar in the control and infected leaves, while silencing of the *McDFR* gene increased the transcript levels of phenylalanine ammonia lyase (*PAL*; *McPAL*), *CHI* (*McCHI*), *FSH* (*McF3H*), *FLS* (*McFLS*), *ANS* (*McANS*) and *UFGT* (*McUFGT*) genes (Fig. 4C).

We also transiently over-expressed the *McDFR* gene in the stem tips of the ever-green cultivar 'Strawberry Jelly', which promoted anthocyanin accumulation at 20 dpi, and a deep green coloration in most of the new buds (Fig. 4A). The anthocyanin content showed a slight increase in the *McDFR*-overexpressing plants (Fig. 4B) and we confirmed that these plants indeed had higher *McDFR* transcript levels in the new buds (Fig. 4C). We also detected an increase in the transcript levels of anthocyanin biosynthetic genes and a decrease in *McFLS* expression, compared with the non-transformed plants (Fig. 4C).

Collectively, these results indicated that *McDFR* expression is associated with red color formation in crabapple leaves, and that changes in *McDFR* expression can affect the expression of downstream genes (e.g. *ANS* and *UFGT*) involved in the anthocyanin biosynthetic pathway.



Figure 2. Analysis of flavonoid accumulation in 5 developmental stages of leaves from the *Malus* crabapple cultivars 'Royalty' and 'Flame'. S1, 3 days after budding; S2, 9 days after budding; S3, 15 days after budding; S4, 21 days after budding; S5, 30 days after budding. (A) Five leaf developmental stages used for the analysis. (B) HPLC analysis of methanol extracts from crabapple leaf. A1, cyanidin 3-O-glucoside; F1, quercetin 3-O-diglucoside; F2, procyanidin dimer; F3, quercetin 3-O-glucoside; F4, quercetin 3-O-glycosidase isomer; F5, avicularin; F6, acetyl quercetin 3-O-glucoside; F1, phloridzin. (C) The total anthocyanins content and total flavonols content in 5 developmental stages of leaves of the two crabapple cultivars. Error bars indicate the standard error of the mean \pm SE of three replicate measurements. Different letters above the bars indicate significantly different values (P < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test.



Figure 3. The relative expression level of *McDFR* and *McFLS* transcripts at different development stages of 'Royalty' and 'Flame' leaves. (A) 'Royalty'. (B) 'Flame'. Error bars indicate the standard error of the mean \pm SE of three replicate measurements. Different letters above the bars indicate significantly different values (*P* < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test.

McFLS is involved in flavonol biosynthesis in crabapple leaves. To confirm the prediction, based on sequence homology, that *McFLS* is a key flavonol biosynthetic gene, we suppressed its expression in the leaves of 'Strawberry Jelly' using the VIGS system and the TRV vector. Approximately 14 days after *Agrobacterium* infiltration, red coloration was seen in the margin and other areas of the infected leaves (Fig. 5A). HPLC analysis confirmed that the levels of anthocyanins were significantly higher in the silenced leaves than in control leaves infiltrated with TRV alone (Fig. 5B). Finally, as seen in Fig. 4C, the expression of *McPAL*, *McDFR* and *McANS* was up-regulated in infected leaves.

We also transiently overexpressed *McFLS* in the stem tips of the ever-red leaves of 'Royalty'. Crabapple stem tips transformed with 35S::*McFLS* developed new buds with a green color (Fig. 5A), indicating that anthocyanin synthesis was suppressed. This was confirmed by HPLC analysis, which also revealed an increase in flavonol content (Fig. 5B). As expected, we observed an increase in *McFLS* transcript levels in green buds infiltrated with 35S::*McFLS* compared to control leaves. This elevated expression of *McFLS* caused an approximately 1.6-fold decrease in *McDFR* transcript abundance, and we observed lower levels of expression of other anthocyanin biosynthetic genes (Fig. 5C). To summarize, these results suggested that *McFLS* expression can promote flavonol accumulation by inhibiting the expression of *McDFR*, and that *McFLS* has a competitive relationship with *McDFR* in the flavonol biosynthetic pathway.



Figure 4. Transient expression of *McDFR* **in crabapple.** *McDFR* expression was suppressed by VIGS using the vector pTRV2-*McDFR* in 'Royalty', or the gene was overexpressed using the vector pBI121-*McDFR* in 'Strawberry Jelly'. Crabapple leaves injected with the empty TRV and pBI121 vectors and infiltration buffer were used as controls. (A) Phenotype of *McDFR* silenced or *McDFR* overexpressing 'Royalty' and 'Strawberry Jelly' leaves. (B) Anthocyanin and flavonol contents at the infiltration sites on crabapple leaves in $\mu g/g$ fresh weight (FW). (C) Relative transcript expression levels in crabapple leaves around the infiltration sites were determined using qRT-PCR. Error bars indicate the standard error of the mean \pm SE of three replicate measurements. Different letters above the bars indicate significantly different values (*P* < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test.

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Variation in the expression of *McDFR* **and** *McFLS* **in apple fruit.** To further characterize the roles of *McDFR* and *McFLS*, we infected apple (*Malus* × *domestica* 'Fuji') fruits with transgenic *Agrobacterium* harboring TRV-*McDFR*, TRV-*McFLS*, pBI121-*McDFR* or pBI121-*McFLS* constructs (Figs 6 and 7). A



Figure 5. Transient expression of *McFLS* **in crabapple.** *McFLS* expression was suppressed by VIGS using the vector pTRV2-*McFLS* in 'Royalty', or the gene was overexpressed using the vector pBI121-*McFLS* in 'Strawberry Jelly'. Crabapple leaves injected with the empty TRV and pBI121 vectors and infiltration buffer were used as controls. (**A**) Phenotype of *McFLS* silenced or *McFLS* overexpressing 'Strawberry Jelly' and 'Royalty' leaves. (**B**) Anthocyanin and flavonol contents at infiltration sites of crabapple leaves in μ g/g fresh weight (FW). (**C**) Relative transcript expression levels in crabapple leaves around the infiltration sites were determined using qRT-PCR. Error bars indicate the standard error of the mean ± SE of three replicate measurements. Different letters above the bars indicate significantly different values (*P* < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test. (A)



CK TRV-McDFR

PBI121-McDFR

CK



Figure 6. Transient expression of *McDFR* in apple fruit. *McDFR* expression was suppressed in apple fruit using the vector pTRV2-*McDFR*, or the gene was overexpressed using the vector pBI121-*McDFR*. Apple fruit injected with the empty TRV and pBI121 vectors and infiltration buffer were used as controls. (A) Phenotype of *McDFR* silenced or overexpressing *McDFR* apple peels. (B) Anthocyanin and flavonol contents at the infiltration sites of apple peels in $\mu g/g$ fresh weight (FW). (C) Relative expression levels around the infiltration sites were determined using qRT-PCR. Error bars indicate the standard error of the mean \pm SE of three replicate measurements. Different letters above the bars indicate significantly different values (*P* < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test.







TRV-McFLS

CK PBI121-McFLS



Figure 7. Transient expression of *McFLS* **in apple fruit.** *McFLS* expression was suppressed in apple fruit using the vector pTRV2- *McFLS*, or overexpressed using the vector pBI121- *McFLS*. Apple fruit injected with the empty TRV and pBI121 vectors and infiltration buffer were used as controls. **(A)** Phenotype of *McFLS* silenced or *McFLS* overexpressing apple peels. **(B)** Anthocyanin and flavonol contents at the infiltration sites of apple peels in $\mu g/g$ fresh weight (FW). **(C)** Relative expression levels were determined using qRT-PCR around the infiltration sites. Error bars indicate the standard error of the mean \pm SE of three replicate measurements. Different letters above the bars indicate significantly different values (*P* < 0.05) calculated using one-way analysis of variance (ANOVA) followed by a Duncan's multiple range test.

	Genes		Flavonoids	
	McDFR	McFLS	Anthocyanins	Flavonols
McDFR	1.00	-0.23	0.55**	-0.52**
McFLS	-0.23	1.00	-0.49**	0.27
Anthocyanins	0.55**	-0.49**	1.00	-0.73**
Flavonols	-0.52**	0.27	-0.73**	1.00

Table 1. Correlation coefficients between the levels of expression of two biosynthetic genes and

flavonoid content. *Correlation coefficient was significant at the $P \le 0.05$ level. **Correlation coefficient was significant at the $P \le 0.01$ level.

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large increase in anthocyanin accumulation was observed at the sites of TRV-*McFLS* and pBI121-*McDFR* over-expression, while the areas of TRV-*McDFR* and pBI121-*McFLS* over-expression showed a yellow or white coloration (Figs 6A and 7A). HPLC quantification of the anthocyanin and flavonol content of the infected areas confirmed that the variation in flavonoid content correlated with the observed variation in fruit color (Figs 6B and 7B). Transcript expression analysis of the transiently expressing tissues further revealed that the up-regulation of *McDFR* expression, or the down-regulation of *McFLS* expression, was accompanied by a proportional increase in the expression levels of *McDFR* as well as some of the genes involved in anthocyanin biosynthesis (*McCHS*, *McCHI*, *McF3H*, *McF3'H*, *McDFR*, *McANS* and *McUFGT*), and a decrease in the expression resulted in fading leaf color and decrease *McDFR* gene expression (Figs 6C and 7C). Finally, we compared the relationship between the expression of *McDFR* and the content of anthocyanins, and the results were consistent with the red color formation in leaves and fruit; as well as between the levels of *McFLS* and the content of flavonols (Figs 5 and 6).

Discussion

Due to the central role that the DFR enzyme plays in the anthocyanin biosynthetic pathway, *DFR* genes have been studied in several monocotyledonous and dicotyledonous species, such as *Forsythia interme*- dia^{40} , *Torenia fournieri*³¹, *Triticum aestivum*⁴¹, *Vitis vinifera*⁴² and *Ascocenda* spp.⁴³, and in some cases at the transcriptional level⁴⁴. The key structural gene, *FLS*, completes the last step of flavonol synthesis. To our knowledge, although *FLS* and *DFR* have been identified and characterized in many land plant species, these genes have yet to be studied in ornamental crabapples. There is growing evidence that anthocyanins and flavonols to contribute to the ornamental and economic value of crabapples, which have highly colorful leaves, fruits and flowers³⁸. In this study we explored the expression and function of *McDFR* and *McFLS* in the biosynthesis and accumulation of anthocyanins and flavonols.

McDFR and *McFLS* expression are coordinated and correlate with the accumulation of anthocyanins and flavonols. In this study, the content of anthocyanins in the leaves of two crabapple cultivars was evaluated, and we observed that it increased in parallel with red pigmentation in several organs/tissues. Furthermore, the anthocyanin levels showed a positive correlation with the expression of *McDFR*, but a negatively correlation with the expression of *McFLS* (Table 1). This result is congruent with a recent study showing a negative correlation of *CnFLS1* expression and anthocyanin synthesis during floral coloration in the petals of transgenic tobacco expressing this gene³⁷. We infer from these observations that there is a competitive relationship between *McDFR* and *McFLS* in flavonoid biosynthesis.

The function of *McDFR* **and** *McFLS* **in flavonoid biosynthesis.** Genetic transformation has been used to test the functions of several genes in the flavonoid biosynthetic pathway in model experimental plants, such as tobacco and *A. thaliana*, as well as in some crop species. For example, overexpression of petunia *CHI* in tomato fruit was reported to lead to an ~65% increase in flavonol levels⁴⁵, while silencing of a *CHS* gene in apple fruit resulted in changes in growth and developmental phenotypes⁴⁶. In *Gentiana triflora* and apples (*Malus* spp.), the silencing of an *ANS* gene caused a reduction in anthocyanin content and, consequently, a much weaker color¹¹. However, few studies to date have focused on genetic transformation using *DFR* or *FLS* genes and none has targeted woody ornamental species, such as crabapple.

Virus-induced gene silencing (VIGS) is a technology that allows the analysis of genes function by suppressing the expression of target genes. In a previous study, we assessed the effect of gene silencing in several plant species, including *Nicotiana benthamiana*, rose (*Rose hybrida*) and strawberry (*Fragaria* × *ananassa*)⁴⁷⁻⁴⁹. Here, we used tissue cultured seedling buds from different crabapple cultivars ('Royalty', 'Strawberry Jelly') to assess the function of *McDFR* and *McFLS* through VIGS and overexpression approaches. We determined that silencing of *McFLS* or overexpression of *McDFR* promoted the accumulation of anthocyanins, while the opposite results were obtained when *McFLS* was overexpressed or *McDFR* expression was silenced. In addition, the abundance of flavonols increased when *McFLS* was overexpressed or *McDFR* was silenced. However, it is interesting that levels of flavonols were slightly elevated in transgenic crabapple leaves that were transiently overexpressing *McDFR* or in which *McFLS*.

expression was silenced. This phenomenon may be explained if McFLS was still expressed but at lower levels degree in McDFR overexpressed or McFLS silenced crabapple plants. In this scenario the lower expression level of McFLS in these transgenic crabapple plants may have resulted in reduced accumulation flavonol, but not a complete block in flavonol biosynthesis. Moreover, correlation coefficients indicated that the levels of expression of McDFR negatively correlated with those of McFLS, as well as with the abundance of anthocyanins and flavonols (Table 1). We propose that there is a competitive relationship between the expression McDFR and McFLS that results in the production of different classes of flavonoid compounds (i.e. anthocyanins or flavonols) (Figs 4–7).

Since dihydroflavonols are substrates for DFR and FLS, they lie at an important branch point in flavonoid biosynthesis, where precursor substrates are channeled toward either anthocyanin or flavonol production. In this regard, the regulation of *DFR* expression and the competing DFR and FLS activities may be particularly important. We conclude that the expression of *McDFR* and *McFLS* may represent a key mechanism for regulating color in crabapple leaves.

A possible explanation for the expression patterns of McDFR and McFLS in two contrasting varieties. R2R3-MYB transcription factors (TFs), which belong to one of the largest plant TF families, are known to be involved in regulating the biosynthesis of anthocyanins and flavonols in A. thaliana⁵⁰. A previously study reported that over-expression of the A. thaliana TF AtMYB12 in tobacco (Nicotiana tabacum) resulted in higher levels of flavonols and increased expression of the FLS gene⁵¹. Moreover, when AtMYB12 was over-expressed in a tissue-specific manner in tomato, the flavonol biosynthesis pathway was activated⁵¹. However, expression of the DFR gene was not induced in AtMYB12 and AtMYB111 over-expressing transgenic plants, as well as in AtMYB111 transgenic lines^{1,44,51}. In apple and crabapple, MYB10, which regulates anthocyanin accumulation and coloration of various organ (e.g. fruit, petals and leaves), can activate the expression of DFR and bind to several the promoters of several anthocyanin biosynthetic genes^{52,53}. In grape berries, expression of the regulatory gene VvMYBF1 is light inducible, and is involved in the control of VvFLS1 transcription and flavonol synthesis in fruit⁵⁴. Thus, we speculated that MYB TFs may similarly regulate flavonoid biosynthesis in crabapple, and that their expression levels may vary at different development stages or in response to different environment conditions. Moreover, changes in the transcript levels of MYB TFs control the biosynthesis of flavonoids, by regulating the expression of various members of the flavonoid biosynthetic pathway. We propose that MYB TFs promote anthocyanin biosynthesis by increasing the transcript levels of DFR during fruit or leaf development, leading to red coloration. However, in response to environmental stresses, MYB TFs activate the transcription of FLS resulting in increased production of flavonols.

Conclusion

In this study, we investigated the functions of *McDFR* and *McFLS* in regulating leaf color in different crabapple cultivars. We demonstrated that the competitive relationship between the expression of *McDFR* and *McFLS* is important for anthocyanin and flavonol synthesis. In addition, the expression of *McDFR* and *McFLS* correlated with anthocyanin and flavonol accumulation, as well as with leaf color. The work described in this report may suggest strategies to genetically modify ornamental plants in order to enhance or modulate flavonoid accumulation.

Methods

Plant material and growth conditions. The plant material used included three *Malus* crabapple cultivars: (1) *Malus* cv. 'Royalty', an ever-red leaf cultivar; (2) *Malus* cv. 'Flame', an ever-green leaf cultivar. Five year old trees of these cultivars grafted on *M. hupehensis* were planted in the Crabapple Germplasm Resources Nursery in Beijing University of Agriculture; and (3) *Malus* cv. 'Strawberry Jelly', an ever-green leaf cultivar, the explants of which were harvested from one-year old branches before spring bud germination, cultured on Murashige and Skoog medium supplemented with 0.1 mg/L 6-Benzylaminopurine (6-BA) and 2 mg/L (2,4-dichlorophenoxy) acetic acid (2,4-D) at 23 °C with a 16h light ($200 \mu mol s^{-1} m^{-2}$) /8h dark period. Plants were grown in a greenhouse at 27 °C under constant illumination.

Leaves of 'Royalty' and 'Flame' were collected at five different developmental stages (Fig. 1A,B) for gene expression analyses and anthocyanin and flavonol quantification. Wild-type 'Royalty' and 'Strawberry' seedlings were grown in a greenhouse, as above. Apple (*Malus* \times *domestica* 'Fuji') fruits were used for analysis of *McDFR* and *McFLS* expression. All samples were frozen in liquid nitrogen upon collection, and stored at -80 °C until further use.

Construction of VIGS vectors and *Agrobacterium***-mediated infiltration of crabapple.** The coding sequences of *McDFR* (GenBank: FJ817487) and *McFLS* (GenBank: KF495602) have previously been deposited in the NCBI (National Center for Biotechnology Information) database. Fragments for the pTRV2-*McDFR* (540 bp) and pTRV2-*McFLS* (490 bp) constructs were amplified by PCR, using gene-specific primers, from a cDNA library derived from *Malus* crabapple leaves (cv. 'Royalty') using Taq DNA polymerase (TAKARA BIOTECH) according to the manufacturer's instructions. The PCR primers used are shown in Table S1. Virus-induced gene silencing vectors carrying the target gene fragments, as well as pTRV1 and pTRV2⁴⁷⁻⁴⁹, were transformed into *Agrobacterium tumefaciens* strain GV3101 competent cells using a freeze-thaw method⁵⁵ and selected on kanamycin- rifampicin-containing (50 mg/L)

LB (Luria Bertani media) plates. Positive clones were verified by restriction enzyme digestion and by sequencing the vector-insert junctions. The harvested bacterial cells were then resuspended to an OD_{600} of 0.5 in infiltration medium (10 mM 2-morpholinoethanesulfonic acid [MES], 200 mM acetosyringone, and 10 mM MgCl₂) and incubated at room temperature for 3h. Before infiltration, bacteria carrying pTRV1 and pTRV2 were mixed in a 1: 1 volume ratio.

For vacuum infiltration, whole plants were submerged in the Agrobacterium suspension and subjected to a vacuum (-25 kPa). When the rate of air bubbles being released from the plants started to decrease, the vacuum was released quickly to allow bacteria to enter the plant tissues. The vacuum treatment time varied from 30s up to 3 min, depending on the vacuum source used. After vacuum infiltration, plants were rinsed with sterile water, and cultured on Murashige and Skoog medium. Fifteen plants from each cultivar were treated, and 'Royalty' was used for silencing of McDFR expression, while 'Strawberry Jelly' was used for silencing of McFLS expression. All experiments were repeated three times.

Overexpression of *McDFR* **and** *McFLS* **in crabapple leaves and fruits.** The full length *McDFR* and *McFLS* open reading frames (ORFs) were cloned from the cDNA library described above and inserted into the pBI121 vector⁵³ using the *XbaI* and *SacI* sites. Primers used for these constructs are shown in Table S1. Transient expression in *Malus* crabapple leaves was performed using the 'Strawberry' Jelly cultivar and *Agrobacterium*-mediated transformation, as described above. *Agrobacterium* cells containing the different constructs were harvested and resuspended in infiltration buffer (10 mM MES, 0.2 mM acetosyringone, and 10 mM MgCl₂) to a final concentration of $OD_{600} = 0.5$. Vacuum infiltration was performed as described above, and infiltration with an empty vector was used as a negative control. Seven days after infiltration, the infected leaves and fruit were collected to observe phenotypic features and to evaluate differences in expression.

RNA extraction. To analyze the effects of VIGS and overexpression on target genes expression, tissue samples from areas showing the silencing and enhancing phenotypes were collected. For controls, corresponding samples were collected from tissues infected by *Agrobacterium* carrying vectors with no host gene fragment insert, or from non-infected plants. Samples from three independent biological replicates were analyzed. Total RNA was extracted from crabapple leaves using the RNA plant plus Reagent (TIANGEN BIOTECH) according to the manufacturer's instructions. DNase (TIANGEN BIOTECH) treatment was performed to remove any genomic DNA according to the manufacturer's instructions. First-strand cDNA was synthesized from total RNA using the Reverse Transcriptase M-MLV (RNase H⁻) kit (TaKaRa).

Quantitative RT-PCR analysis. qRT-PCR was performed using the SYBR[®] Premix Ex TaqTM II (Perfect Real Time) (TaKaRa, Ohtsu, Japan) and the CFX96TM Real Time System (Bio-Rad, USA). The PCR amplification conditions were as previously described⁵⁶, and transcript levels were determined by relative quantification using the *Malus* 18S ribosomal RNA gene (DQ341382) as the internal control and the $2^{(-\Delta\Delta CT)}$ analysis method was applied. Specific primers (Table S1) for semi-quantitative RT-PCR and qRT-PCR analysis were designed using the primer 5 software⁵⁷.

HPLC analysis. Crabapple leaf samples (approximately 0.8–1.0 g fresh weight) were subjected to extraction with 10 mL extraction solution (methanol: water: formic acid: trifluoroacetic acid=70: 27: 2: 1)⁵⁸ at 4 °C in the dark for 72 h, shaking every 6 h. The supernatant was isolated by filtration through filter paper and a further filtration through a 0.22 μ m MilliporeTM filter (Billerica, MA, USA). For the HPLC analysis, trifluoroacetic acid: formic acid: water (0.1: 2: 97.9) was used as mobile phase A and trifluoroacetic acid: formic acid: acetonitrile: water (0.1: 2: 48: 49.9) was used as mobile phase B. The gradients used were as follows: 0 min, 30% B; 10 min, 40% B; 50 min, 55% B; 70 min, 60% B; 30 min, 80% B. Detection was performed at 520 nm for anthocyanin and 350 nm for flavonol⁵⁸, respectively. All samples were analyzed in three biological triplicates (extracted from three different batches of leaves).

In this study, we employed HPLC-ESI (\pm)-MS2 analysis to identify the kinds of compounds by standards and comparing their spectroscopic data to literature^{59,60}. Cyanidin-3-O-glucoside, quercetin-3-O-glucoside, avicularin, phloridzin, quercetin (Sigma-Aldrich, Germany), Procyanidin B2 (Sigma-Aldrich, UK) was used as standards.

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Acknowledgments

We thank the Fruit Tree Key Laboratory at the Beijing University of Agriculture. We also thank the Beijing Key Laboratory for Agricultural Application and New Technique for providing experimental resources. We are grateful to all the technicians in the BUA Crabapple Germplasm Resource Garden. We thank PlantScribe (www.plantscribe.com) for carefully editing this manuscript. Financial support was provided by the National "Twelfth Five-Year" Plan for Science & Technology Support (Project No. 2013BAD02B01-4), National Natural Science Foundation of China (31301762), the Project of Construction of Innovative Teams and Teacher Career Development for Universities and Colleges Under Beijing Municipality (IDHT20140509), Beijing Municipal Commission of Education Science and Technology Promotion Plan (PXM2014-014207-000081), and Scientific Research Improvement Project of Beijing University of Agriculture (GZL2015003). The funding agencies had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Author Contributions

Conceived and designed the experiments: J.T. and Y.Y. Performed the experiments: Z.H. and Y.H. Analyzed the data: J.Z. and T.S. Contributed reagents/materials/analysis tools: Y.Y. and T.S. Wrote the paper: J.T., Y.Y. and Z.H.

Additional Information

Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Tian, J. *et al.* The Balance of Expression of Dihydroflavonol 4-reductase and Flavonol Synthase Regulates Flavonoid Biosynthesis and Red Foliage Coloration in Crabapples. *Sci. Rep.* **5**, 12228; doi: 10.1038/srep12228 (2015).

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