Differential regulation of the anthocyanin profile in purple kiwifruit (Actinidia species)

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Abstract
Anthocyanins are a group of secondary metabolites that colour fruit and flowers orange, red, purple or blue depending on a number of factors, such as the basic structure, co-pigmentation, metal ion complexation and vacular pH. The biosynthesis of anthocyanin is regulated at the transcriptional level by a group of transcription factors, the MYB–bHLH–WD40 (MBW) complex. In this study, the purple colouration in several kiwifruit (Actinidia) species was identified and characterised as red cyanidin-based and blue delphinidin-based anthocyanins. The differential pigmentation in the skin and flesh can be attributed to the differential ratio of cyanidin and delphinidin derivatives accumulated in the total anthocyanin profile. The expression of anthocyanin biosynthetic genes chalcone synthase (CHS), flavonoid 3-O-glucosyltransferase (F3GT), flavonoid 3′-hydroxylase (F3′H) and flavonoid 3″′-hydroxylase (F3″′H) is crucial for anthocyanin accumulation. However, the balance of expression of the F3′H and F3″′H genes appears responsible for the ratio of cyanidin and delphinidin derivatives, while a lack of CHS, F3GT and MYB110 expression is responsible for a lack of total anthocyanins. The transcriptional regulation of the F3′H and F3″′H promoters by the R2R3 MYB transcription factor MYB110 is markedly different in tobacco transient assays. When kiwifruit MYB10 or MYB110 are over-expressed in Actinidia chinensis both cyanidin-based and delphinidin-based anthocyanins are elevated, but F3′H and F3″′H genes are not strongly correlated with MYB expression. These results suggest that the core kiwifruit anthocyanin pathway genes are dependent on characterised MYB transcription factors, while other regulatory proteins are more directly responsible for the expression of the F3′H and F3″′H genes.

Introduction
Anthocyanins are flavonoids that belong to the polyphenols group. These secondary metabolites are synthesised in plants and are essential for plant growth and development1. Anthocyanin is well recognised for colouring fruits and flowers in a range of hues from orange, red, to blue and violet, making them attractive to pollinators and dispersers, while also serving as an indicator of fruit quality and ripeness for consumers1,2. In addition to co-pigmentation, different pH level and metal ion complex formations can moderately change the visual colours of anthocyanins, the biosynthetic genes flavonoid 3′ hydroxylase (F3′H) and flavonoid 3′, 5′-hydroxylase (F3″′H) are responsible for the type of anthocyanin accumulated8. F3′H belongs to the CYP75B subfamily of the cytochrome P450-dependent monoxygenase (P450) superfamily and converts dihydrokaempferol to dihydroquercetin which is the precursor of cyanidin-based anthocyanin. F3″′H belongs to the CYP75A subfamily of the P450 superfamily and converts dihydrokaempferol and dihydroquercetin to dihydromyricetin, which is the

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precursor of delphinidin-based anthocyanin. It appears that the F3′5′H enzyme has evolved from F3′H precursors several times in flowering plants.

Studies in a broad range of plant species have shown that the overall anthocyanin accumulation is regulated at the transcriptional level by R2R3 MYB transcription factors (TF), such as PAP1 (AtMYB75) in Arabidopsis, MYB1 and MYB10 in apple, AN2 in Petunia, VvMYB1A in grape and FaMYB10 in strawberry. These MYBs often partner with a bHLH TF to form a MBW complex involved in the induction of anthocyanin production and are characterised by the conserved R2R3 DNA-binding domains as well as the conserved [A/S/G] NDV motif for anthocyanin-promoting MYB TFs in dicots, the C-terminus KPRPR[S/T]T motif and the bHLH-interacting motif. Over-expression of PAP1 in Arabidopsis resulted in elevated transcript level of bHLH TF TT8 and Petunia bHLH TF AN1 interacts with MYB TF AN2 to regulate anthocyanin pathway. In the model plant Nicotiana tabacum, transient over-expression of kiwifruit AcMYBF110 revealed that the endogenous bHLH TF NtAN1 is an activator of anthocyanin biosynthetic genes and is regulated by the other endogenous bHLH TF NtJAF13.

Kiwifruit belongs to the genus Actinidia which comprises more than 70 species, with fruit that show different colours and texture in both skin and flesh. The pigment in the red-fleshed kiwifruit (Actinidia chinensis) has been previously identified as cyanidin-based anthocyanin and the anthocyanin biosynthetic gene flavanoid 3-O-glycosyltransferase F3GT1 was responsible for accumulation of anthocyanin. An R2R3 MYB TF MYB110 was identified from Actinidia hybrid population that determines red petal colour and activates F3GT1 to produce anthocyanin. In terms of fruit colour, AcMYB75 and AcMYBF110 have been identified from A. chinensis for their activation of anthocyanin biosynthetic genes. Delphinidin was first reported in two taxa: A. melanandra and A. arguta var. purpurea but the genes regulating its accumulation remain to be studied. Recently it has been reported that small amounts of delphinidin derivatives are present in the red-inner pericarp of A. chinensis ‘Hongyang’. The final ratio of 3′-hydroxylated derivatives to 3′5′-hydroxylated anthocyanins has been a focus of attempts to engineer flower colour, such as the blue rose, which required both introduction of transgenes for F3′5′H and DFR as well as suppression of the endogenous DFR. In chrysanthemum, over-expression of a daisy F3′5′H and suppression of the endogenous F3′H did not result in more delphinidin derivatives, while a pansy F3′5′H under the regulation of a CHS promoter resulted in delphinidin production in chrysanthemum flower petals. Differential regulation of these genes at the transcriptional level has been studied. In grape, genes encoding F3′5′H and F3′H are expressed in tissues accumulating flavonoids, especially in skin of ripening red berries; there appear to be at least 15 paralogues of the F3′5′H and two F3′H genes. The transcript profiles during véraison (colour change) correlate with accumulation of cyanidin-based and delphinidin-based anthocyanins. Climatic effects have been studied in grape, where the ratio of F3′5′H to F3′H may be lowered due to hot temperatures close to harvest. UV levels appear to differentially affect expression of F3′H/F3′5′H genes in Antarctic moss, Pohlia nutans, while in grape, light exclusion increases the ratio of dihydroxylated to trihydroxylated anthocyanins and the ratio of expression of F3′H/F3′5′H.

In grape, two copies of the gene encoding the F3′H enzyme are present in the grape genome; only one is expressed in vegetative and reproductive tissues. A detailed study of this F3′H gene showed it to be well expressed in many tissues, but elevated in berry skins. Its promoter was isolated and used to identify transcriptional regulators. Interestingly, VvMYBA1 (the MYB activator of anthocyanin biosynthesis) was not identified as an interactor, but the interactors isolated included grape MYB repressors, NACs, EIN3 and homeodomain TFs. Furthermore, the grape MYBA1 was responsible for F3′5′H activation and induced the production of trihydroxylated anthocyanins whereas the two closely related MYB TFs, MYBA6 and MYBA7, were unable to activate the trihydroxylated branch of the anthocyanin pathway. In kiwifruit, a F3′5′H gene has been isolated and shown to generate delphinidin-based anthocyanins when transiently expressed in tobacco and kiwifruit. Furthermore, insertion of more MYB-binding sites into the promoter of this F3′5′H upregulates delphinidin-derivative content when transiently over-expressed in the presence of kiwifruit MYB110. Here, we aim to characterise the key biosynthetic genes that are necessary for accumulation of anthocyanin in kiwifruit, and to ascertain if there is differential regulation of F3′H and F3′5′H that contributes to the ratio of cyanidin and delphinidin in different kiwifruit species.

Results

Colour development during ripening

The fruits were harvested at the mature green stage as defined by 100% black seed coat colour, average firmness between 636 and 980 kgf, average dry matter of 10% and average degrees Brix of 4. The fruits were then ripened at 20°C. Hue angles as measured by Minolta CR300 showed that all four Actinidia species had similar green/yellow skin colour at harvest (Fig. 1a, b). As the fruit ripened, the intensity of red hues increased on the skin and eventually blue hues developed, as detected by Minolta chromameter. The deepest colour was in the skin of A. arguta var.
purpurea (Ap), followed by A. macrosperma × A. melanandra (MaMe) Red and A. melanandra (Me). Only the MaMe Yellow remained at low a*/b* ratio which indicates yellow hues. A similar colour development was observed in the fruit flesh, changing from green to deep red at the ripe stage (Supplemental Fig. 1). The exception was the MaMe Yellow kiwifruit where both the skin and flesh colour developed from green to yellow at the colour-change stage, and then to orange at the ripe stage.

Detection of cyanidin and delphinidin in fruit skin and flesh

Anthocyanins were analysed by HPLC and LCMS in the skin and flesh of the ripening fruit lines. There was no detected anthocyanin at the mature green fruit stage. At both colour-change stage and ripe stage, anthocyanin accumulation was higher in the skin than in the flesh, hence the intense pigmentation observed in the skin. Cyanidin-3-O-β-galactoside (cy-gal), cyanidin 3-O-[2-O-(β-xyllosyl)-β-galactoside] (cy-xylgal) and delphinidin 3-O-[2-O-(β-xyllosyl)-β-galactoside] (dp-xylgal) were detected in the purple kiwifruit (Supplemental Fig. 3) as previously reported. The most abundant cyanidin-based compound was cy-xylgal, comprising between 20 and 99% of total anthocyanins.

The two cyanidin-based anthocyanins comprised over half of the total anthocyanin profile in the skin and flesh tissue of Me and Ap, while the delphinidin-based anthocyanin was predominant in the skin and flesh tissue of MaMe Red (Fig. 2). Delphinidin was detected in Me skin in relatively small amounts. The highest total anthocyanin accumulation at ripe stage was measured for Ap, which accumulated 976 μg/g FW in skin tissue and 250 μg/g in flesh, followed by MaMe Red (478 μg/g in skin, 81 μg/g in flesh) and Me (150 μg/g in skin, 159 μg/g in flesh). No anthocyanin was detected in MaMe Yellow kiwifruit.

At the ripe stage, MaMe Red skin comprised 73% delphinidin-based and 27% cyanidin-based anthocyanin (Supplemental Table 1). The proportion of delphinidin derivatives and cyanidin derivatives in the flesh was 46 and 54%, respectively. In Ap, cyanidin-based anthocyanins contributed to 64% of the total anthocyanin in the skin and 97% in the flesh. In contrast, cyanidin-based anthocyanins contributed to almost all the measured anthocyanin in skin and flesh in Me.

It appears that the total anthocyanin level increased as the fruit transitioned from colour-change stage to ripe stage but the proportion of cyanidin and delphinidin remained at a similar ratio (Fig. 2).

Expression analysis of anthocyanin biosynthetic genes during fruit colour change

Two key enzymes responsible for the synthesis of cyanidin-based and delphinidin-based anthocyanins are
the closely related flavonoid 3’-hydroxylase (F3’H) and flavonoid 3’, 5’-hydroxylase (F3’5’H). In the recently published genome of *Actinidia chinensis*<sup>30</sup>, there are three F3’H-like and two F3’5’H-like genes. Previously, gene expression of two of the AcF3’H genes was detected in the red-fleshed *A. chinensis*, which contains cyanidin 3-O-[2-O-(β-xyllosyl)-β-galactoside]<sup>21</sup>. Recently, the AcF3’5’H gene has been shown to produce delphinidin-based anthocyanin in *Agrobacterium*-mediated transient assays in tobacco and kiwifruit<sup>38</sup>. The DNA sequence of F3’H and F3’5’H genes was acquired from the reference *A. chinensis* genome and was cloned from all four *Actinidia* species tested. F3’5’Hs clustered with the *A. chinensis* F3’5’H within the F3’5’H clade from other plant species.

Two F3’H-like genes and one full-length F3’5’H-like gene were expressed in the skin and flesh of purple fruit lines. Transcript of the F3’5’H was detected in the skin at all developmental stages in all kiwifruit species (Fig. 4), but less so in the flesh. Moreover, the expression of the F3’Hs and F3’5’H was detected at the mature green stage when there was no anthocyanin detected. The expression of F3’5’H was the highest in Ap skin at the colour-change stage. MaMe Red skin also showed high levels of F3’5’H expression at the colour-change stage and the ripe stage. F3’5’H expression was lower in Me compared to that in Ap and MaMe Red. Of the two F3’H-like genes, F3’H2 appeared to have higher expression than F3’H1 in Me, MaMe Red, Yellow and Ap, except for when its low expression at colour-change stage coincided with the slightly higher F3’H1 expression in Ap.

Pearson’s correlation showed that the expression of F3’5’H is correlated with total anthocyanin accumulation (*r* = 0.87, *p* = 0.0024<sup>**</sup>) at colour-change stage (Table 1), and that the expression of F3’H1 correlated strongly with the accumulation of cyanidin-based anthocyanin (*r* = 0.85, *p* = 0.0041<sup>**</sup>). Considering the multiple stop codons throughout the deduced amino acid sequence of F3’H1s, F3’H2 appears to be the functional enzyme converting substrates to cyanidin-based anthocyanins. However, when considering all tissues, F3’H2 has a statistically negative association with total anthocyanin accumulation. This is due to the
Fig. 3 (See legend on next page.)
expression of F3’H2 in the absence of anthocyanin in MaMe Yellow.

The expression of other biosynthetic genes in the anthocyanin pathway was analysed in both skin and flesh tissue across the three developmental stages (Supplemental Figs. 2A and 2B). Gene sequences were mined from the recently published A. chinensis genome40. Chalcone synthase (CHS) and flavonon 3-O-glucosyltransferase (F3GT) were well expressed in both the skin and flesh of purple kiwifruit but barely detected in MaMe Yellow skin and flesh.

Pearson’s correlation shows that the expression of CHS and F3GT was strongly correlated with the total anthocyanin accumulation ($r = 0.94, p < 0.001^{***}$) and $r = 0.96, p < 0.001^{***}$, respectively) (Table 1). Furthermore, chalcone isomerase (CHI) and dihydroflavonol reductase (DFR) were also associated with total anthocyanin accumulation ($r = 0.87, p = 0.0028^{**}$ and $r = 0.88, p = 0.0021^{**}$, respectively), in which DFR shows strong correlation with delphinin-derivative accumulation ($r = 0.97, p < 0.001^{***}$).

Flavonone 3-hydroxylase (F3H) and leucoanthocyanidin dioxygenase (LDOX) were highly expressed in all the kiwifruit (even fruit without anthocyanin). While DFR and CHI were correlated with anthocyanin accumulation, the lack of expression of CHS and F3GT in MaMe Yellow suggests these two genes may limit final anthocyanin levels, while the ratio of the expression of F3’H2 and F3’S’H determines the profile of the accumulated anthocyanins.

Expression analysis of anthocyanin-associated MYB transcription factors

Two R2R3 MYB transcription factors were analysed for their expression associated with anthocyanin accumulation across the three developmental stages. MYB10 has been implicated in fruit coloration in A. chinensis43 while MYB110 controls petal colour in A. eriantha44. MYB10 expression was very low at the mature green stage for all kiwifruit and became undetectable in the later stages, suggesting its transcription is not associated with anthocyanin accumulation (Fig. 5). However, MYB110 was highly expressed in all three purple kiwifruit, particularly the skin, but absent in the MaMe Yellow kiwifruit.

A very strong correlation existed between MYB110 expression and anthocyanin accumulation at ripe stage ($r = 0.95, p < 0.001^{***}$) (Supplemental Table 2). Two potential bHLH TF partners of the anthocyanin-associated MYB were also analysed21. The expression of bHLH2 was undetectable in any of the developmental stages; however, bHLH5 was expressed in all stages and in all kiwifruit including MaMe Yellow.

Pearson’s correlation shows expression of bHLH5 is correlated with total anthocyanin accumulation at colour-change stage ($r = 0.82, p = 0.0065^{**}$) but not at ripe stage ($r = 0.26, p = 0.27$) (Supplemental Table 2).

MYB110 is an anthocyanin-related R2R3 MYB transcription factor

The TF MYB110 was cloned from Me, Ap and MaMe Red kiwifruit for sequence comparison with the known A. chinensis MYB110. Alignment showed the protein sequence of Me, Ap and MaMe Red were 100% identical to each other and shared 96% similarity with the AcMYB110 (Fig. 6a).

Alignment with identified anthocyanin-related R2R3 MYB proteins revealed R2R3 DNA-binding domains and the highly conserved bHLH co-factor interacting signature motif ([D/E]Lx2[R/K]x3Lx6Lx3R) in the R2R3 domain of MYB11017. Also the conserved [A/S/G]NDV motif for anthocyanin-promoting MYB TFs in dicot and the C-terminus KPRPR[S/T]F motif present in anthocyanin-related regulators was present in the kiwi-fruit MYB11018. Furthermore, infiltration of MYB110 (the identical sequence from either Me, Ap and MaMe Red) induced anthocyanin production in Nicotiana tabacum leaves (Fig. 6b), producing 240 ± 41 μg/g FW of delphinidin-based anthocyanin and 434 ± 82 μg/g FW of cyanidin-based anthocyanin (Supplemental Fig. 4). This supports the hypothesis that MYB110 from these species is an anthocyanin-related R2R3 MYB TF and is similar to AcMYB11021.
Transcriptional regulation of \( F3' \)H and \( F3'S' \)H by MYB110

MYB110 was examined for its ability to trans-activate the Actinidia \( F3' \)H and \( F3'S' \)H promoters in dual luciferase transient assays in \( N. \) benthamiana leaves. AcMYB110 was included in the transient assay as a positive control as it shows high similarity with MYB110 from purple kiwifruit and has shown strong activation on anthocyanin biosynthetic gene \( F3GT \) in kiwifruit flower petals\(^{24} \). The reporter gene \( \beta \)-Glucuronidase (GUS) driven by the 35S promoter was the negative control for background promoter activity.

The results showed that MYB110 strongly activated certain \( F3' \)H promoters, 33-fold higher in Me and 25-fold higher in Ap (Fig. 7a). This was species specific; there was no activation of the two MaMe \( F3' \)H promoters by either AcMYB110 or MYB110, as their promoter activity readings were lower than the GUS control. The same applied for the other published kiwifruit MYB AcMYB10 (data not shown). The potential partner of the MYB, bHLH5, gave no additional activation, possibly due to the strong endogenous activity of the tobacco bHLH, NtAN1. There was no activation of any of the cloned Actinidia \( F3'S' \)H promoters by infiltration with AcMYB110 or MYB110, even in the presence of bHLH5 (Fig. 7b). Surprisingly each of the promoters showed high background activity (twice as much luciferase produced as 35S-renilla without additional transcription factors) and this activity could not be further enhanced by cloned MYBs or bHLHs. Therefore, MYB110 showed strong activation of the Actinidia \( F3' \)H promoters cloned from Me and Ap but seemed unable to activate MaMe \( F3' \)H promoters or any of the \( F3'S' \)H promoters. This suggests the \( F3' \)H and \( F3'S' \)H promoters are differentially regulated; in some instances the promoter of the \( F3' \)H can be upregulated by MYB110, while the \( F3'S' \)H is not directly affected by known kiwifruit MYBs.

The activity of the \( F3' \)H and \( F3'S' \)H in A. chinensis plants expressing 35s:MYB110 or 35s:MYB10

To further examine the regulation of kiwifruit delphinidin biosynthesis, three independent lines of A. chinensis over-expressing either 35s:AcMYB10 or 35s:AcMYB110 were examined. Unlike control empty vector plants in the same greenhouse conditions, the leaves of 35s:AcMYB110 were completely red at all stages, while the young and expanded leaves of 35s:AcMYB10 showed red colour in the centre (Fig. 8a).
A small amount of cyanidin derivative (8 μg/g FW) was detected in the young leaves of the empty vector control lines, but not in expanded or mature leaves. In AcMYB10 over-expression lines, anthocyanin was visible in the centre of the young (33 μg/g FW) and expanded leaves (17 μg/g FW), and was detected along the veins of the mature leaves but this was visibly masked by green colouration (76 μg/g FW) (Supplemental Table 3).

Anthocyanin production was significantly higher in the AcMYB110 over-expression lines, accumulating 643 μg/g of cyanidin derivative in young leaves, 771 μg/g in expanded leaves and up to 1105 μg/g FW of cyanidin derivative in the large mature leaves.

Delphinidin-derivative was detected in all leaves from AcMYB10 and AcMYB110 over-expression line ranging from 1–2 μg/g FW and 25–32 μg/g FW, respectively. Real time quantitative PCR (RT-PCR) revealed that the expression of MYB110 correlated with the expression of CHS as well as F3GT in red leaves, which is consistent with previous findings (Fig. 8c). 23

Both copies of F3’H were expressed regardless of the MYB over-expression and the expression of F3’5’H was elevated in MYB110 over-expression line but not to the same extent as the expression of CHS and F3GT, hence the low-delphinidin accumulation. This suggests that in A. chinensis, the regulatory effect of MYB11 is more prominent on CHS and F3GT and less direct on F3’H and F3’5’H for anthocyanin accumulation.

Discussion

The intense pigmentation on both the skin and flesh of the purple kiwifruit is due to the accumulation of cyanidin-based and delphinidin-based anthocyanin. The presence of both cyanidin and delphinidin has been observed in some A. melanandra and A. arguta var. purpurea genotypes. The data presented here show that the key change during the ripening process of A. melanandra, A. arguta var. purpurea and the two MaMe kiwifruits is the increase in total anthocyanin concentration while the composition and the ratio of cyanidin to delphinidin remained the same between the colour-change stage and the ripe stage.

A comparison of the anthocyanin biosynthetic gene transcription between the purple A. melanandra, A. arguta var. purpurea and MaMe Red, with MaMe Yellow kiwifruit showed that CHS and F3GT expression were strongly correlated with the accumulation of total anthocyanin. CHS is responsible for initiating flavonol biosynthesis and F3GT has been shown to be responsible for anthocyanin biosynthesis in red-fleshed kiwifruit. The anthocyanin biosynthetic genes F3H and LDOX have high expression in the purple fruits but there was no statistical evidence that their gene expression correlated with the accumulation of anthocyanin. F3H catalyses the hydroxylation of the substrate naringenin to form dihydrokaempferol (DHK), which is the precursor of anthocyanin as well as the metabolic intermediate for flavonols. In a recent publication, LDOX was proposed to have a critical role in anthocyanin biosynthesis in A. arguta species. LDOX is undoubtedly required for processing substrate flux down the anthocyanin biosynthetic pathway, but was less correlated with anthocyanin accumulation than CHS, F3GT and MYB110. F3’H and F3’5’H are important for anthocyanin accumulation as F3’H catalyses the hydroxylation on the intermediate substrate DHK into dihydromyricetin (DHM) at the
Fig. 5 Expression of MYB110, bHLH5 and MYB10 in Actinidia melanandra, A. arguta var. purpurea, MaMe Red and MaMe Yellow. Expression was measured in the skin and flesh tissue across the three developmental stages: mature green, colour change and ripe. Error bars are SEM for three biological replicates.
### Fig. 6 Comparison of kiwifruit MYB sequence and function.

**a** Deduced amino acid sequence alignment of AcMYB75, AcMYB110, MYB10 from *Actinidia chinensis* and other plant species in comparison to MYB110 from *A. chinensis* and the purple kiwifruit lines. The bHLH interaction domain are highlighted in grey box and box A indicates the conserved amino acid residues specific to anthocyanin-promoting MYB in dicot and box B indicates the C terminus motif that is specific to anthocyanin-related regulators.

**b** Infiltration site 7 days after transient transformation of AcMYB110 on the left side of the *N. tabacum* leaf and MYB10 from *A. arguta var. purpurea* on the right side of the leaf.

| Sequence       | AcMYB75 | AcMYB110 | MYB10 A. chinensis | MYB10 A. arguta var. purpurea | MYB10 A. arguta var. purpurea MaLe red | AcMYB110 (A. chinensis) | AcMYB110 (A. arguta var. purpurea) | AcMYB110 (A. arguta var. purpurea) MaLe red |
|----------------|---------|----------|------------------|-----------------------------|----------------------------------------|-------------------------|----------------------------------|-----------------------------------------|
| AcMYB75        |         |          |                  |                             |                                        |                         |                                  |                                         |
| AcMYB110       |         |          |                  |                             |                                        |                         |                                  |                                         |
| MYB10 A. chinensis |       |          |                  |                             |                                        |                         |                                  |                                         |
| MYB10 A. arguta var. purpurea |       |          |                  |                             |                                        |                         |                                  |                                         |
| MYB10 A. arguta var. purpurea MaLe red |       |          |                  |                             |                                        |                         |                                  |                                         |
| AcMYB110 (A. chinensis) |       |          |                  |                             |                                        |                         |                                  |                                         |
| AcMYB110 (A. arguta var. purpurea) |       |          |                  |                             |                                        |                         |                                  |                                         |
| AcMYB110 (A. arguta var. purpurea) MaLe red |       |          |                  |                             |                                        |                         |                                  |                                         |

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3’ position of the B-ring which forms the precursor of cyanidin while F3′5′H catalyses the hydroxylation on both the 3’ and 5’ position which forms the precursor of delphinidin. The expression of these two genes was present in all kiwifruit tissue tested, even in the MaMe Yellow where no anthocyanin was accumulated. In the previously characterised red-centred A. chinensis, the red colour was due to the accumulation of cyanidin-based anthocyanin only and no delphinidin-based anthocyanin was detected. In this cultivar of A. chinensis, only F3’H1 and F3’H2 gene expression was detected while no transcription of F3′5′H was observed. This suggests that while CHS and F3GT are crucial for total anthocyanin accumulation, F3’H and F3′5′H are required for determining anthocyanin composition.

In this study, the ratio of cyanidin to delphinidin appeared to be determined as soon as the colour develops at the colour-change stage. During colour-change stage, the expression of F3’H1 was absent and the expression of F3’H2 was very low, but it appeared to be complemented by the high F3′5′H expression at the colour-change stage and the ripe stage, hence a high amount of delphinidin was accumulated in the MaMe Red kiwifruit. In A. arguta var. purpurea the expression of F3’H2 was very low at the colour-change stage but was complemented by the high expression of F3′5′H which may explain the similar level of cyanidin-based and delphinidin-based anthocyanin. In addition, F3’H and F3′5′H were expressed at the mature green stage of all kiwifruit despite the lack of red colour at that stage. There were multiple stop codons at consistent amino acid positions throughout the deduced amino acid sequence of F3’H1 cloned from purple kiwifruit lines, suggesting that F3’H2 was the most likely candidate for the functional enzyme responsible for cyanidin accumulation.

R2R3 MYB transcription factors also play a role in regulating anthocyanin biosynthesis in kiwifruit. Previously, AcMYB110 has been shown to determine red petal colour in kiwifruit flowers and activated F3GT expression. Here, MYB110 was highly expressed in both the skin and flesh of the purple kiwifruit lines but absent in the MaMe Yellow kiwifruit. The expression showed correlation with total anthocyanin accumulation and coincided with the colour development, as well as the expression of the biosynthetic genes CHS and F3GT. Protein alignment showed 96% similarity with AcMYB110, while the presence of anthocyanin regulator specific motifs and the ability to induce anthocyanin in tobacco leaves suggests a functionally similar MYB activator of anthocyanin biosynthesis. In promoter activation assays in the model plant N. benthamiana, MYB110 induced activation of the F3’H promoter isolated from A. melanandra and A. arguta var. purpurea, but did not induce activation of the MaMe F3’H promoters. Moreover, there was no activation of any of the cloned F3′5′H promoters even in the presence of the MBW complex partner bHLH5. The TF bHLH5 was most highly
Fig. 8 The activity of F3′H and F3′5′H in Actinidia chinensis over-expressing empty 35s vector, 35s:AcMYB10 and 35s:AcMYB110. a Digital image of young, expanded and mature leaves from the transformed A. chinensis lines. b Measurement of cyanidin-based and delphinidin-based anthocyanin in the leaves. Error bars indicate SEM for three independent plant lines. c Expression of MYB10, MYB110 and anthocyanin biosynthetic genes. F3′H flavonoid 3′ hydroxylase, F3′5′H flavonoid 3′ 5′ hydroxylase, CHS chalcone synthase, F3GT flavonoid 3-O-glucosyltransferase. Error bars indicate SEM for three independent plant lines.
expressed at the ripe stage in all lines, but as it showed low correlation to anthocyanin accumulation and MYB110 expression, and the transcript appeared to be independent of MYB110 expression.

Surprisingly both F3'H and F3'S'H are well expressed in MaMe Yellow kiwifruit despite the absence of MYB110 expression, which implies MYB110 it is not the key regulator of F3'H and F3'S'H. MYB110 was found to activate two of the cloned F3'H promoters, and no F3'S'H promoters, so it is apparent that these two genes are differentially regulated and with other transcription factors likely to regulate anthocyanin profile. In a recent publication, AcMYB110 was only able to induce significant delphinidin accumulation in A. eriantha kiwifruit only when the F3'S'H promoter was engineered with the MYB-binding site R6 motif\textsuperscript{28}. Otherwise, the AcMYB110 has very little activation on the native AcF3'S'H promoter. It appeared that when AcMYB110 was transiently expressed in A. eriantha kiwifruit, there was significantly more cyanidin-based anthocyanin and little delphinidin-based anthocyanin.

In leaves of stably transformed A. chinensis lines, F3'S'H expression was elevated with over-expression of MYB110, and hence some delphinidin derivatives were produced (2–4% of total anthocyanin). However, the magnitude of induction was significantly less than those of CHS and F3GT. MYB110 over-expression facilitated significant accumulation of cyanidin as the F3'Hs transcripts were already abundant, even in wild-type tissue. Therefore, from results of both stable transgenics and transient assays MYB110 (or MYB10) does not appear to be the main regulator of F3'H and F3'S'H. In two recent publications, AcMYB75 and AcMYBF110 were reported to be the key R2R3 TFs for anthocyanin regulation in kiwifruit\textsuperscript{25,26}. Protein alignment showed that AcMYB75 and AcMYBF110 are both 99.5% identical to AcMYB10 examined in this study (Fig. 6a). MYB10 shows no correlation with anthocyanin biosynthesis in purple kiwifruit lines. In A. chinensis AcMYB75/MYB10 and AcMYBF110 are important in controlling flesh colour. MYB110 appears to be the activator TF of anthocyanin biosynthesis in purple kiwifruit lines A. melanandra, A. arguta var. purpurea and MaMe species. It is crucial for anthocyanin accumulation and may partially determine the composition of cyanidin-based and delphinidin-based anthocyanin. However, while it promotes the accumulation of anthocyanin, other TFs or regulatory mechanisms appear responsible for the accumulation of delphinidin-based compounds.

**Conclusion**

The anthocyanin biosynthesis pathway has been extensively studied and key transcriptional regulators have been identified. Previous research on anthocyanin biosynthesis in kiwifruit has focused on total anthocyanin accumulation rather than the specific profile of anthocyanins. While it is important to understand the mechanism that switches on the overall pathway, the regulation involved in determining the type of anthocyanin that accumulates will be useful in metabolic engineering. Further research is now required to extend the understanding of the regulation of the anthocyanin profile to achieve novel colours and maximise health benefits.

**Materials and methods**

**Plant material and storage condition**

Actinidia melanandra (Me), A. arguta var. purpurea (Ap) and A. macroasperma × A. melanandra (MaMe) fruits were grown at the Plant and Food Research Orchard, Motueka, New Zealand. Sixty fruits from each vine were collected at the mature green stage. At mature green stage, harvested fruits showed 100% black seed coat colour, average firmness between 636 and 980 kgf, average dry matter between 10 and 12% and average degrees Brix of 4. Twenty fruits were sampled at each stage. For the colour-change stage and ripening stage, all fruits were stored at 20 °C for ripening. Skin peel and flesh tissue were separated, snap-frozen and collected into three biological replicates, each consisting of three fruit.

**Skin and fruit colour measurement**

Skin colour change was measured using a Minolta CR-300 chromameter (Konica Minolta, Mahwah, NJ, USA) by the L*a*b* colour space (CIE, 1976)\textsuperscript{47}. Measurements were taken on two flat surfaces of each of the four fruits for each stage and repeated twice.

**Anthocyanin identification and quantification**

Skin peel and flesh tissue were ground to powder under liquid nitrogen. Anthocyanin concentrations were measured by ultra-high performance liquid chromatography (UHPLC) and the identity of the anthocyanins detected was confirmed by liquid chromatography/mass spectrometry analysis (LC-MS). Approximately 300 mg of skin peel powder and 500 mg of flesh powder for each biological replicate from Me, Ap and two MaMe were freeze-dried overnight. Anthocyanins were extracted with acidified methanol (100% MeOH + 0.1% HCl, 5× volume per material weight) for 2 h at room temperature. A measure of 1 mL of supernatant was spin-dried followed by re-suspension in 500 μL of 20% methanol. The supernatant was filtered through a syringe filter and diluted with 20% methanol prior to analysis. The system used was a Dionex UltiMate 3000 Series UHPLC (ThermoFisher Scientific, San Jose, CA, USA) with PDA (photodiode array) detection at 520 nm. Compound separation was achieved using an Acclaim PA2 C18 column, 3 μ 2.1 × 150 mm (Dionex, ThermoFisher Scientific), maintained at 35 °C. Solvents
were (A) 0.1% formic acid and (B) acetonitrile + 0.1% formic acid and the flow rate was 350 μL/min. The initial mobile phase, 100%A was ramped linearly to 92%A at 5 min, then 85%A at 10 min, 80%A at 20 min and 100%B at 34 min. The total run time per sample was 40 min and the sample injection volume was 5 μL. Standards of cyanidin-3-O-β-glucoside were used to quantify anthocyanin concentrations and the results for individual and total anthocyanins are reported as cyanidin-3-O-β-glucoside equivalents per gram fresh weight (FW). For anthocyanin quantification in A. chinensis leaves, 300 mg of freeze-dried leaf tissue powder were extracted with 2 mL of acidified methanol and used the same procedure as described above.

Identification of detected anthocyanins was confirmed by LC-MS using an LTQ linear ion trap mass spectrometer fitted with an ESI interface (ThermoFisher Scientific, San Jose, CA, USA) coupled to an Ultimate 3000 UHPLC and PDA detector (Dionex, Sunnyvale, CA, USA). Anthocyanin separation was achieved using a Poroshell 120 SB-C18 column, 2.7 μ × 150 mm (Agilent, Torrance, CA, USA), maintained at 70 °C. Solvents were (A) 5:3:92 acetonitrile:formic acid:water v/v/v and (B) acetonitrile + 0.1% formic acid and the flow rate was 200 μL/min. The initial mobile phase, 100%A was held for 2 min before being ramped linearly to 88%A at 14 min, 5% A at 15 min and held for 4 min before resetting to the original conditions. Sample injection volume was 10 μL.

MS data were acquired in the positive mode using a data-dependent LC-MS³ method. This method isolates and fragments the most intense parent ion to give MS² data (daughter ions), then isolates and fragments the most intense daughter ion (MS³ data). The capillary voltage, capillary temperature, sheath gas pressure and auxiliary gas were set at 43 V, 275 °C, 35 psi and 10 psi, respectively.

Real time quantitative PCR expression analysis

RNA was isolated from the ground skin and flesh tissue using the Spectrum Plant Total RNA kit (Sigma-Aldrich, www.sigmaaldrich.com) and reverse transcribed into cDNA using the QuantiTect Reverse Transcription kit (Qiagen, www.qiagen.com) according to the manufacturer’s protocol. Genes of the anthocyanin pathway were identified by BLAST with genes of known function on the Actinidia chinensis genome and the Actinidia arguta RNA-seq data⁴⁰,⁴⁸ (Supplemental Table 4). Gene-specific primers were designed using Vector NTI 11.0 (http://www.invitrogen.com/) and are summarised in supplementary table 5 (Additional file). RT-PCR was carried out using the LightCycler 480 with LightCycler 480 SYBR Green I Master (Roche Diagnostics, USA). Each reaction volume was 5 μL and reactions were run in quadruplicate and a non-template control and water control were included in each run. The thermal cycling conditions were 95 °C for 10 s, 60 °C for 10 s and 72 °C for 20 s, then a melting temperature cycle with continuous fluorescence data acquisition from 65 to 95 °C. The data output was analysed by the lightcycler480 software 1.5 using the Target/Reference ratio to compare the expression level of the target genes normalised to the reference gene, A. chinensis orthologue of At1g07940 elongation factor 1-a (EF1α)⁴⁹.

Isolation, cloning and sequence alignment of candidate genes

Sequences of candidate genes were identified by BLAST from the A. chinensis genome⁴⁰ and PCR amplified from cDNA by specific primers (Supplemental Table 5). PCR product of MYB110 were cloned into pSAK277 over-expression vector using In-Fusion cloning (Takara Bio USA, Inc.). F3’H and F3’5’H genes were amplified using specific primers from the four Actinidia species cDNA and cloned in pHEX2 vectors using Gateway cloning (ThermoFisher Scientific, USA). Genes were sequenced by Macrogen, Korea. Multiple nucleotide and amino acid sequence alignment and phylogeny tree were created by Geneious 10.0.3.

Transgenic over-expression of MYB110 in tobacco leaves

The vector pSAK277 with 35s:MYB110 construct was transformed into Agrobacterium tumefaciens strain GV3101 by electroporation followed by incubation on plate before infiltration. N. tabacum plants were grown under glasshouse conditions using natural light with daylight extension to 16 h as described by Espley et al.¹². Three leaves of 6-week-old N. tabacum were used for infiltration and kept under the same growth conditions. Leaves were photographed and harvested 7 days after infiltration in liquid nitrogen and stored at −80 °C until analysis.

Dual luciferase transient assay in tobacco leaves

Genomic DNA was extract from skin tissue of A. melanandra, A. arguta var. purpurea and MaMe using the DNeasy Plant Mini kit (Qiagen, http://www.qiagen.com/). The 2.3 kb promoter region of F3’H and F3’5’H were isolated by polymerase chain reaction (PCR), cloned into pGreen 0800-LUC vector using In-Fusion HD cloning (Clontech, Takara Bio USA) and transformed into A. tumefaciens GV3101⁵⁰. The promoter sequence was confirmed by sequencing (Macrogen, Korea). Promoter activation dual luciferase assays were performed on leaves of 6-week-old N. benthamiana grown under above mentioned glasshouse condition. The promoters of F3’H and F3’5’H were co-infiltrated with MYB110²⁴. Three days after inoculation, four leave discs from each treatment were sampled to assay the firefly luciferase and renilla luciferase activities using the dual luciferase assay reagent (Promega, Madison, WI, USA). The promoter activities were expressed as a ratio of LUC to REN activity²⁴,⁵⁰.
Stable over-expression of MYB10 and MYB110 in A. chinensis

Newly initiated leaves of A. chinensis “Hort16A” from in vitro grown shoots were inoculated with A. tumefaciens strain EHA105 containing the vectors of pSAK277 with 35s:MYB10 and 35s:MYB110, respectively. The transformation procedure was based on the previous report34. Calli which formed in the regeneration and selection medium containing 150 mg/L of kanamycin were excised individually and transferred to fresh regeneration and selection medium for bud induction. The shoots generated from these independent transformant calli were excised and cultured in the selection medium containing 50 mg/L of kanamycin. After they had rooted, these transgenic plants were potted and grown in the containment greenhouse.

Statistical analysis

The correlation relationship between the amount of anthocyanin concentration and expression of the anthocyanin biosynthetic genes were calculated and presented in Pearson’s correlation coefficient r (Supplemental Table 2).

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A.C.A and R.V.E conceived the study and led the research. Y.P. performed the experiments and with A.C.A and R.V.E wrote the manuscript. T.W. and K.L.W generated, maintained and pollinated the transgenic plants. J.C. performed the metabolite analysis. All authors discussed the results and reviewed and contributed to the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

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Supplementary Information

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