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Colour variation in red grapevines (Vitis vinifera L.): genomic organisation, expression of flavonoid 3'-hydroxylase, flavonoid 3',5'-hydroxylase genes and related metabolite profiling of red cyanidin-/blue delphinidin-based anthocyanins in berry skin

Simone D Castellarin1, Gabriele Di Gaspero*1, Raffaella Marconi1, Alberto Nonis1, Enrico Peterlunger1, Sophie Paillard2,3, Anne-Francoise Adam-Blondon2 and Raffaele Testolin1

Address: 1Dipartimento di Scienze Agrarie e Ambientali, University of Udine, via delle Scienze 208, 33100 Udine, Italy, 2UMR de Génomique Végétale, INRA-CNRS-UEVE, 2, Rue Gaston Crémieux, CP5708, 91057 Evry Cedex, France and 3UMR118 INRA-AgroCampus Rennes, Amélioration des Plantes et Biotechnologies Végétales, Domaine de la Motte, BP 35327, 35633 Le Rheu Cedex, France

Email: Simone D Castellarin - castellarin.simone@spes.uniud.it; Gabriele Di Gaspero* - gabriele.digaspero@uniud.it; Raffaella Marconi - marconi@uniud.it; Alberto Nonis - Alberto.nonis@uniud.it; Enrico Peterlunger - enrico.peterlunger@uniud.it; Sophie Paillard - paillard@rennes.inra.fr; Anne-Francoise Adam-Blondon - adam@evry.inra.fr; Raffaele Testolin - testolin@uniud.it

* Corresponding author

Abstract

Background: Structural genes of the phenyl-propanoid pathway which encode flavonoid 3'- and 3',5'-hydroxylases (F3'H and F3'5'H) have long been invoked to explain the biosynthesis of cyanidin- and delphinidin-based anthocyanin pigments in the so-called red cultivars of grapevine. The relative proportion of the two types of anthocyanins is largely under genetic control and determines the colour variation among red/purple/blue berry grape varieties and their corresponding wines.

Results: Gene fragments of VvF3'H and VvF3'5'H, that were isolated from Vitis vinifera 'Cabernet Sauvignon' using degenerate primers designed on plant homologous genes, translated into 313 and 239 amino acid protein fragments, respectively, with up to 76% and 82% identity to plant CYP75 cytochrome P450 monooxygenases. Putative function was assigned on the basis of sequence homology, expression profiling and its correlation with metabolite accumulation at ten different ripening stages. At the onset of colour transition, transcriptional induction of VvF3'H and VvF3'5'H was temporally coordinated with the beginning of anthocyanin biosynthesis, the expression being 2-fold and 50-fold higher, respectively, in red berries versus green berries. The peak of VvF3'5'H expression was observed two weeks later concomitantly with the increase of the ratio of delphinidin-/cyanidin-derivatives. The analysis of structural genomics revealed that two copies of VvF3'H are physically linked on linkage group no. 17 and several copies of VvF3'5'H are tightly clustered and embedded into a segmental duplication on linkage group no. 6, unveiling a high complexity when compared to other plant flavonoid hydroxylase genes known so far, mostly in ornamentals.

Conclusion: We have shown that genes encoding flavonoid 3'- and 3',5'-hydroxylases are expressed in any tissues of the grape plant that accumulate flavonoids and, particularly, in skin of ripening red berries that synthesise mostly anthocyanins. The correlation between transcript profiles and the kinetics of accumulation of red/cyanidin- and blue/delphinidin-based anthocyanins indicated that VvF3'H and VvF3'5'H expression is consistent with the chromatic evolution of ripening bunches. Local physical maps constructed around the VvF3'H and VvF3'5'H loci should help facilitate the identification of the regulatory elements of each isoform and the future manipulation of grapevine and wine colour through agronomical, environmental and biotechnological tools.
Background
Vacuolar accumulation of anthocyanins in berry skin is responsible for the pigmentation of red/blue grapevine (Vitis vinifera) cultivars. Anthocyanin biosynthesis is developmentally triggered at the onset of berry ripening (8–10 weeks after blooming) and lasts till harvest [1]. Expression profiling of structural genes and transcription factors has claimed a major role for the gene UFGT encoding a UDP-glucose:flavonoid 3-O-glucosyltransferase and its regulatory Myb-type gene VvMybA in conveying the flux of flavonoid intermediates towards the synthesis of anthocyanins [2,3]. The enzyme UFGT is synthesised since véraison under the control of VvMybA and catalyses the 3’-glycosidation of cyanidin- and delphinidin-based anthocyanidins [4]. The resulting stably coloured anthocyanins may undergo methylation upon the activity of methyl-transferases that convert cyanidin into peonidin and delphinidin into petunidin or malvidin, similarly to how it occurs in other anthocyanin-producing species [5,6].

Cyanidin- and delphinidin-derived anthocyanins diverge from each other regarding the number of the substituted groups that are found on the B-ring of the flavonoid skeleton (either two or three, respectively) and have their chromatic properties affected accordingly. Cyanidin-based anthocyanins exhibit a reddish colour whereas delphinidin-based anthocyanins are purple to blue. They jointly determine the tonality of red berry grape varieties and their corresponding wines. The relative proportion of the five anthocyanins is largely under genetic control and unique to each cultivar [7,8]. Sunlight exposition and water deficit have been shown to modulate to some extent the anthocyanin profile [9,10].

In plant species, the biosynthesis of cyanidin- and delphinidin-type precursors is driven, upstream of the enzyme UFGT, by the activity of two flavonoid 3’- and 3’,5’-hydroxylases (F3’H and F3’,5’H) that add either a single hydroxyl group at the 3’ position or two hydroxyl groups at the 3’ and 5’ positions to dihydrokaempferol and/or dihydroquercetin. Once converted into 3’-dihydroquercetin or 3’,5’-dihydroxymyricetin, these intermediates flow through common downstream enzymes to form di-substituted and tri-substituted anthocyanins when the gene UFGT is expressed and to form other polyphenols (flavonols, leucoanthocyanins, catechins, proanthocyanins, tannins) at different developmental stages. F3’H and F3’,5’H genes code for CYP75 cytochrome P450 hydroxylases that have been well characterised in the ornamental

Table 1: List of the PCR primers used.

| Target DNA | Name       | 5’→3’ sequencea |
|------------|------------|-----------------|
| Plant F3’H | F3’H_S     | CCIGTAAANTNGNCARYTIBIA |
| Plant F3’H | F3’H_AS    | GCICKYTGIAIGRINACRTA |
| Plant F3’5’H | F3’5’H_S  | ATGCTGACNTAYGCATGGNCINAYTATG |
| Plant F3’5’H | F3’5’H_AS | CGAARTCRTTTGCKKNONGTNCDAYTT |
| Grape F3’H  | VvF3’H_S   | ATTCGACCCTGAAATG |
| Grape F3’H  | VvF3’H_AS  | CAAGAAGACTGGAGCAGT |
| Grape F3’5’H | Vv3’5’H-1A_S | GAGGTGGACTGTATTTATTTAAAGAT |
| Grape F3’5’H | Vv3’5’H-1A_AS | AAGAAATGGACCAATGTTGCAAG |
| Grape F3’5’H | Vv3’5’H-2A_S  | AGGCCCTTAAATGGTTGAAAG |
| Grape F3’5’H | Vv3’5’H-2A_AS | CTTGAGACTCGTCGAGGAGT |
| Grape F3’H  | VvF3’H-intron_S | ATCATTTTCAGGTGCGGCAAT |
| Grape F3’H  | VvF3’H-intron_AS | GCTCACTATACCAACATTTAAAGCA |
| Grape F3’5’H | Vv3’5’H-2A-intron_S | AAGGTCCTATTACACATTTAAAG |
| Grape F3’5’H | Vv3’5’H-intron_AS | GGCAGTAGACTTCAGCAGAG |
| BES38C24-FM | 38C24-FM_S | GTCATCTCAGGGCAGAG |
| BES38C24-FM | 38C24-FM_AS | TTCCGGCCCTTGGTTGAG |
| BES38C24-RM | 38C24-RM_S | GCGATTGATCATGCCCTTGG |
| BES38C24-RM | 38C24-RM_AS | TCATATTGGGAGGAGGACA |
| BES17K4-FM | 17K4-FM_S  | CCAATATTGGGAGGAGGACA |
| BES17K4-FM | 17K4-FM_AS | CATCAGGGTGGAGGAGGACA |
| BES9C17-FM | 9C17-FM_S  | AGGAAGCCTTGTGCAAGA |
| BES9C17-FM | 9C17-FM_AS | GGAAGCCTTGTGCAAGA |
| BES40A22-FM | 40A22-FM_S | CATTTGAGGCTACAGGTTTC |
| BES40A22-FM | 40A22-FM_AS | CGTGCATATCTTTTCAGG |
| BES5A23-FM | 5A23-FM_S  | TCCAAGAAGGAGGAGGTCG |
| BES5A23-FM | 5A23-FM_AS | AGCAGTACTGCGTGCTTCA |

a Degenerate IUB code: I = inosine; R = A or G; M = A or C; W = A or T; Y = C or T; S = C or G; D = A, G or T; N = A, C, G, or T.
plants Antirrhinum majus [11], Campanula medium [12], Dianthus caryophyllus [13], Gentiana triflora [14], Ipomoea purpurea, [15], Petunia hybrida [16], Phalaenopsis [17] and Vinca major [18]. In all these species, F3'H and F3',5'H are found as low/medium copy number genes and have been genetically manipulated to produce novel flower pigmentation. In grapevine, only tentative consensus (TC) sequences for F3'H and F3',5'H genes are available in the TIGR grape gene database http://www.tigr.org. TCs were built based on partially overlapping ESTs from cDNA libraries obtained from different genotypes and tissues. This makes it unfeasible to predict or rule out the presence of different isogenes.

This paper reports (1) the isolation of flavonoid 3'- and 3',5'-hydroxylase gene members in grapevine via sequence homology, (2) the assessment of their chromosomal location by genetic and physical mapping, (3) their transcript profiles in different tissues, (4) the time-course of expression in berry skin of the red cultivar ‘Merlot’ from prévéraison to harvest compared to the curves of accumulation of anthocyanins, catechins and total polyphenols.

**Results and discussion**

**Isolation of flavonoid 3'- and 3',5'-hydroxylase genes in grapevine**

Gene isolation was accomplished via sequence homology with the help of newly designed degenerate primers. PCR bands that were amplified from genomic DNA of Vitis vinifera ‘Cabernet Sauvignon’ were longer than predicted on the basis of plant flavonoid hydroxylase transcripts (Table 1 and 2).

Five different sequences were isolated with the F3'H degenerate primers following SSCP analysis and were given the designation VvF3'H-1a, -1b, -1c, -1d, and VvF3'H-2 [GenBank:DQ298196-DQ298200]. Once the primer sites had been removed, the sequences VvF3'H-1a to -1d were 1,034 bp long with a 95 bp intron; VvF3'H-2 was 1,011 bp long with a 95 bp intron. Sizing of genomic amplicons performed by capillary electrophoresis confirmed that the two classes of amplicons diverge from each other for a 23-bp indel. Sequence identity was between 99.8 and 99.9% (99.7–99.8% for the coding region) among the sequences VvF3'H-1. The presence of mutations introduced by Taq polymerase during the PCR process and immortalised through the cloning of single PCR-amplified molecules could not be ruled out as the origin of differences among the four VvF3'H-1 sequences. The sequences VvF3'H-1a to -1d translated into protein fragments of 313 amino acids with 311 positions conserved among the sequences when pair-wise comparisons were performed (Figure 1). VvF3'H-2 contained a frame shift mutation due to a 23-bp deletion in the coding region. Function was putatively assigned to VvF3'H-1 based on its highest identity match in GenBank at the protein level (76% identity, E = 8e-134) that corresponded to a flavonoid 3'-hydroxylase of Petunia × hybrida [GenBank:AAD56282]. The nucleotide coding sequence of VvF3'H-1 matched the grapevine tentative consensus sequence TC42042 that has been predicted from three partially overlapping ESTs sequenced in ‘Cabernet Sauvignon’ and ‘Chardonnay’ (TIGR grape gene index). The VvF3'H-1 protein fragments spanned the 1–245 amino acid region of TC42042 and extended it on the 5’ end of the gene by 67 amino acids. Amino acid sequence identity between the four VvF3'H-1 protein fragments and TC42042 was 95–96%.

The degenerate primers targeting F3’5’H yielded five sequences that were given the designation VvF3’5’H-1a, -1b, -1c, VvF3’5’H-2a, -2b [GenBank:DQ298201-

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**Table 2: Primers pairs and PCR conditions used for the amplification of grapevine flavonoid hydroxylase genes.**

| Code   | Sense primer | Anti-sense primer | Expected size (bp) | Annealing temp. (°C) |
|--------|--------------|-------------------|--------------------|----------------------|
| F3'H   | F3'H_S       | F3'H_AS           | 825*               | 60                   |
| F3'5'H | F3'5'H_S     | F3'5'H_AS         | 1020*              | 56                   |
| F3     | VvF3'H_S     | VvF3'H_AS         | 196                | 58                   |
| F3-1   | VvF3'H-indel_S | VvF3'H_AS      | 166                | 58                   |
| F3-2   | VvF3'H-2A_S  | VvF3'H-2A_AS      | 152                | 65                   |
| F3int  | VvF3'H-intron_S | VvF3'H-intron_AS | 255                | 58                   |
| F3-1int| VvF3'H-1-A-intron_S | VvF3'H-intron_AS | 558                | 58                   |
| F3-2int| VvF3'H-2-A-intron_S | VvF3'H-intron_AS | 593                | 58                   |
| 38C24-FM | 38C24-FM_S | 38C24-FM_AS       | 330                | 58                   |
| 38C24-RM | 38C24-RM_S     | 38C24-RM_AS     | 312                | 58                   |
| 17K14-FM | 17K4-FM_S | 17K4-FM_AS       | 327                | 58                   |
| 9C17-FM    | 9C17-FM_S     | 9C17-FM_AS       | 358                | 58                   |
| 40A22-FM | 40A22-FM_S     | 40A22-FM_AS     | 327                | 58                   |
| 5A23-FM    | 5A23-FM_S     | 5A23-FM_AS       | 218                | 54                   |

* in absence of introns.
Once the degenerate primer sites had been removed, the sequences VvF3’5’H-1a, -1b and -1c were 1,120 bp long with a 403 bp intron; VvF3’5’H-2a and VvF3’5’H-2b were 1,122 bp long with a 405 bp intron. The translated coding region matched several plant F3’5’H genes in GenBank. VvF3’5’H-1a, and VvF3’5’H-1b shared the highest identity score at the amino acid level with the accession AAP31058 of *Gossypium hirsutum* (81%, E = e\(^{-111}\)) as well as VvF3’5’H-2a and -2b did with the accession AAM51564 of *Glycine max* (82%, E = e\(^{-112}\)). Translated sequences of VvF3’5’H-1a and VvF3’5’H-1b diverged from each other for two out of 239 amino acids. The same occurred between the sequences VvF3’5’H-2a and VvF3’5’H-2b (Figure 1). The sequences VvF3’5’H-1a and VvF3’5’H-1b had 93–94% identity at the amino acid level with VvF3’5’H-2a and VvF3’5’H-2b. All sequences matched the tentative consensus TC45860 obtained from 19 ESTs at the TIGR grape gene index. The 'Cabernet Sauvignon' gene fragments identified in this work spanned the 187–424 amino acid region of TC45860 over a predicted ORF of 508 amino acids. The sequences VvF3’5’H-1a and VvF3’5’H-1b peptide sequences had 95% identity with TC45860. The VvF3’5’H gene fragments partially overlapped also with a EST (ID CTG1028815) of a 'Cabernet Sauvignon' berry cDNA library already analysed by MPSS http://mpss.udel.edu/grape. In order to predict how many functional copies of the gene VvF3’5’H are present in the grape genome we looked at the number of different ESTs available at the grape gene databases. Twelve EST singletons held at the TIGR and the EST CTG1028815 spanned
the 5' end of the VvF3'5'H gene fragments whilst four ESTs were alignable to the 3' end. All 17 ESTs, irrespective of their tissue-specific localisation and the source genotype, were aligned with the 'Cabernet Sauvignon' genomic sequences VvF3'5'H-1a, -1b, -1c, -2a and -2b (Figure 2). Six ESTs clustered with the 'Cabernet Sauvignon' VvF3'5'H-1a and VvF3'5'H-1b gene fragments. Those ESTs were detected in leaves of the white variety 'Chardonnay' and in petioles and berries of the red variety 'Cabernet Sauvignon'. Eleven ESTs grouped together with the 'Cabernet Sauvignon' VvF3'5'H-2a and VvF3'5'H-2b genomic sequences. The latter ESTs were present in various tissues (leaves and berries) of red ('Cabernet Sauvignon' and 'Syrah') and white ('Chardonnay') cultivars. Two major groups of F3'5'H transcripts including either VvF3'5'H-1 or VvF3'5'H-2 can be detected in different tissues of both red and white grapevines, suggesting that at least two genes encoding CYP75A cyt P450 hydroxylases are expressed in flavonoid-synthesizing tissues. Due to the putative presence of SNPs in ESTs from different cultivars, it was not possible to ascertain the number of functional copies belonging to each group, but together with the number of highly similar sequences obtained from 'Cabernet Sauvignon' genomic DNA, this pointed to the presence of multiple gene members coding for different isoforms of flavonoid hydroxylases in the grape genome.

In order to test the phylogenetic relationship between grapevine hydroxylases and the multi-functional family of plant monoxygenases, a Neighbor-Joining tree was constructed using translated sequences of VvF3'H, VvF3'5'H-1, VvF3'5'H-2 and 670 protein entries of plant cyt P450 monoxygenases (Figure 3). VvF3'H and VvF3'5'H grouped solely with P450 proteins specifically annotated as flavonoid hydroxylases belonging to the family-specific branch of CYP75 monooxygenases. VvF3'H and VvF3'5'H split into two terminal branches, each one hosting either CYP75B or CYP75A monooxygenases. Although most plant P450s are of polyphyletic origin and some P450 families are lineage-specific [19], CYP75 P450s are ancient genes that existed even before the separation between gymnosperms and flowering plants [20]. The ancient origin of CYP75 monooxygenases is also supported by the finding that intron position of VvF3'H and VvF3'5'H was conserved when compared to other plant genes coding for flavonoid hydroxylases such as the Arabidopsis gene TT7 [21].

**Genetic mapping**

Genetic loci harbouring genes that code for flavonoid hydroxylases were mapped using the single strand conformational polymorphism (SSCP) analysis of F3'H and F3'5'H sequences in an F1 pseudo-testcross mapping population originated from the cross 'Cabernet Sauvignon'
Vitis 'breeding line 20/3'. The two genes segregated independently and were mapped to linkage groups LG17 and LG6, respectively (Figure 4). The linkage groups were annotated according to [22] and included also reference microsatellite markers and SSCP markers generated from BAC end-sequences (BES) of a 'Cabernet Sauvignon' BAC library. The locus containing the gene that codes for flavonoid 3’-hydroxylase was identified by linkage analysis following the segregation of the SSCP of two co-segregating bands amplified from the parent 'Cabernet Sauvignon' by means of the primer pair F3int (Table 2). The VvF3'H locus mapped to the middle of LG17 and co-segregated with the BES marker 5A23-FM and the microsatellite marker VMC9g4. The locus containing the genes that code for flavonoid 3',5'-hydroxylases was mapped by means of a polymorphic SSCP band amplified from the parent 'Cabernet Sauvignon' using the primer pair F35-1int. The primer pair F35-1, specific for the genes VvF3'5'H-1a, identified five BAC clones. The primer pair F35-2, specific for VvF3'5'H-2b, identified two different BAC clones. Twelve

Figure 3
The CYP75 clan of plant cyt P450s. The branch containing the CYP75 family is focused on a Neighbor-Joining tree constructed using 670 plant cytochrome P450 monooxygenases. Flavonoid 3'-hydroxylases VvF3'H-1a to -1d and flavonoid 3',5'-hydroxylases VvF3'5'H-1a to -2b that were identified in grapevine are in bold. They split into two lower hierarchical branches that identified the CYP75B (flavonoid 3'-hydroxylases) and CYP75A (flavonoid 3',5'-hydroxylases) sub-families, respectively.

Vitis 'breeding line 20/3'. The two genes segregated independently and were mapped to linkage groups LG17 and LG6, respectively (Figure 4). The linkage groups were annotated according to [22] and included also reference microsatellite markers and SSCP markers generated from BAC end-sequences (BES) of a 'Cabernet Sauvignon' BAC library. The locus containing the gene that codes for flavonoid 3’-hydroxylase was identified by linkage analysis following the segregation of the SSCP of two co-segregating bands amplified from the parent 'Cabernet Sauvignon' by means of the primer pair F3int (Table 2). The VvF3'H locus mapped to the middle of LG17 and co-segregated with the BES marker 5A23-FM and the microsatellite marker VMC9g4. The locus containing the genes that code for flavonoid 3',5'-hydroxylase was mapped by means of a polymorphic SSCP band amplified from the parent 'Cabernet Sauvignon' using the primer pair F35-1int. This marker mapped to the middle of LG6 and co-segregated with microsatellite marker VrZag30 and BES markers 9C17-FM and 17K4-FM (Figure 4). Other SSCP bands that were co-amplified by the same primer pair F35-1int could not be scored in the progeny due to faintness or homozygosis. SSCP marker F35-2int could not be mapped in the 'Cabernet Sauvignon' parental map, but co-segregated with VrZag30 and SSCP marker F35-1int in the male parent of the same cross (data not shown).

Physical mapping and anchoring of BAC contigs to the genetic map
A total of 34 BAC clones containing the genes VvF3'H and VvF3'5'H were identified by screening a set of 18,432 BACs corresponding to a 6X genome coverage [23]. The primer pair F3, specific for grapevine flavonoid 3'-hydroxylase, detected 1.75 gene copies per haploid genome equivalent. Two bands of 196 bp and 173 bp, corresponding to the VvF3'H-1 and the VvF3'H-2 gene fragments, respectively, were co-amplified from all but two of the BACs, while the 196 bp band alone was amplified from the other two BAC clones. The genes coding for flavonoid 3',5'-hydroxylases were searched in the BAC library using the primer pairs F35-1 and F35-2 (Table 2). The primer pair F35-1, specific for the genes VvF3'5'H-1a, identified five BAC clones. The primer pair F35-2, specific for VvF3'5'H-2b, identified two different BAC clones. Twelve
additional BAC clones were identified by both primer pairs. The number of positive BAC clones per haploid genome equivalent was on average 2.4 for the gene VvF3'5'H-1 and 2.75 for the gene VvF3'5'H-2. Local physical maps around the VvF3'H and VvF3'5'H loci were constructed using all positive BAC clones identified. These BAC clones were used to query 2,035 BAC contigs resulting from a whole genome physical map. This map was assembled using FPC software [24] at a 10-50 cut-off from the genome end-sequences of BAC clones positive for VvF3'H and VvF3'5'H and mapped using the SSCP technique. VMC, VVI, UDV, VVMD, VVS, and VrZag are microsatellite markers. Numbers on the left indicate the genetic distance between the markers expressed in centiMorgan (cM).

Two flavonoid 3',5'-hydroxylase genes, VvF3'H-1 and VvF3'5'H-2, were tightly linked in contig ctg253 that consisted of 28 BAC clones and approximately covered 700 kb (Figure 5). Six BAC clones contained both isogenes, whilst two BAC clones (6A08 and 5A23) contained only VvF3'H-1. BAC clone 5A23 was initially not included in the ctg253 following computational contig assembly because it did not meet the cut-off due to a low number of fingerprint bands. However, it was later placed in the contig upon PCR anchorage to other BAC clones using the BES-derived primers (BES5A23-FM). The same marker BES5A23-FM was used to anchor contig ctg253 to the 'Cabernet Sauvignon' genetic map. The marker based on the SSCP of the VvF3'H gene fragments and the BES-derived marker co-segregated at the level of resolution of the genetic map.

Flavonoid 3',5'-hydroxylase genes were present in four contigs (ctg313, ctg871, ctg1320, and ctg2373) that covered approximately 450, 600, 280 and 140 kb, respectively. Each one of the contigs ctg313, ctg1320 and ctg2373 contained both VvF3'5'H-1 and VvF3'5'H-2 gene fragments. Contig ctg871 contained only the VvF3'5'H-2 gene fragment. Contigs ctg313, ctg871, ctg1320 and ctg2373 included 26, 29, 10 and 3 BAC clones, respectively, of which 14, 10 and 3, respectively, contained either one or both isoforms of flavonoid 3',5'-hydroxylases as shown in Figure 6. VvF3'5'H-1 and VvF3'5'H-2 fragments were amplified from each individual positive BAC clone of contigs ctg313, ctg871, ctg1320 and ctg2373 using the primers pairs F35-1int and F35-2int, respectively, and analyzed by the SSCP technique. Based on BAC order and the number of SSCP haplotypes, at a first approximation we hypothesize the occurrence of four loci containing VvF3'5'H-1-type sequences and seven loci containing VvF3'5'H-2-type sequences over the four contigs (Figure 6). Two paralogous copies of gene VvF3'5'H-1 were identified in ctg313. The first paralog had two allelic variants, that could be detected by SSCP analysis, one present in three BAC clones 6H20, 9C17, 31P20 (corresponding to the SSCP band genetically mapped from 'Cabernet Sauvignon') and the other in the remaining five clones (data not shown). Two BAC clones (28B15 and 99117) spanned the region that included both paralogs. A third potential paralogous copy of the gene VvF3'5'H-1 was identified in contig ctg1320 whose BAC clones contained one of the two SSCP haplotypes present in the BACs from contig ctg313. An additional different haplotype of the gene VvF3'5'H-1 was identified in contig ctg2373 that included just three BAC clones. We could detect a potential of seven paralogous copies of the gene VvF3'5'H-2 in contigs ctg313, ctg871, ctg1320 and ctg2373. The primers pair F35-2int identified three types of gene fragments distinguishable by size, hereafter referred to as short, medium and long (Figure 6). All three gene fragments were found in ctg1320 in the close vicinity of the gene copy of VvF3'5'H-1. Three BAC clones contained all three forms at the same time, confirming the paralogous nature of these VvF3'5'H-2 gene copies. The medium sized gene fragment VvF3'5'H-2 was also amplified in ctg313 and ctg2373 where it showed a SSCP haplotype different from that found in ctg1320 (data not shown). A different locus carrying the longer fragment of
VvF3'5'H-2 was present in ctg871 with a SSCP haplotype not distinguishable from that found in ctg1320.

**Integration of genetic and physical maps around the locus VvF3'5'H**

Contigs ctg313, ctg871, ctg1320 and ctg2373 were anchored to the 'Cabernet Sauvignon' genetic map by using SSR markers present in the BAC clones and single strand conformational polymorphism of the BAC-end sequences (Figure 7). The SSR marker VrZag30 that was identified in five BACs of ctg313 allowed this contig to be anchored to LG6. Map position of ctg313 was confirmed by mapping the SSCP marker BES9A23-FM developed on a BAC clone embedded into ctg313 that co-segregated with SSR VrZag30. The contig ctg871 was anchored to LG6 by the SSR marker VMC3a8. The contig ctg1320 was anchored to LG6 using the SSR VMC3f12 and the BAC end-derived SSCP marker BES38C24-RM. The contig ctg2373 was anchored to LG6 by the BAC end-derived SSCP marker BES17K4-FM. The SSCP band of the marker

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**Figure 5**

**Physical map of the VvF3'H locus.** Map of BAC contig ctg253 carrying the VvF3'H-1 and VvF3'H-2 genes and its anchorage to LG17 of 'Cabernet Sauvignon' and *V. vinifera* consensus genetic maps. BAC clone 5A23 bears the VvF3'H-1 isogene and was included after PCR anchoring onto the contig using marker BES9A23-FM. Names written in horizontal refer to BAC clones and contigs; those written in vertical refer to markers and genes. The orientation of the contig is arbitrary. Genetic and physical distances are not drawn to scale. The presence of a given marker on a BAC clone is highlighted with the symbol '●'. The symbol 'X' stands for lack of amplification of a given marker in the corresponding BAC clone. BAC clones missing any symbol at marker BES9A23-FM were not tested. Physical localisation of VvF3'H-1 and VvF3'H-2 loci is shown in grey boxes.
BES17K04-FM inherited from 'Cabernet Sauvignon', which corresponded to the SSCP band amplified from the BAC clone 17K04, was amplified in the mapping population and completely co-segregated with SSR VrZag30 and BES9C17-FM. The marker BES38C24-RM mapped 2.2 cM apart from those three markers and co-segregated with SSR VMC3a8. All contigs ctg313, ctg871, ctg1320 and ctg2373 were located on LG6 in the region covered by the markers VrZag30, VMC3a8, BES9C17-FM, BES17K4-FM and BES38C24-RM. This interval spanned 2.2 cM in the 'Cabernet Sauvignon' parental map. Contigs ctg313, ctg871, ctg1320 were also cross-referenced to the V. vinif-
era consensus map [A. Doligez, unpublished data] by the SSR markers VrZag30, VMC3a8 and VMC3f12, respectively. The region spanned 3.0 cM and the median VMC3a8 marker was settled 0.6 cM and 2.4 cM away from VrZag30 and VMC3f12, respectively (Figure 7).

The BES of positive BAC clones provided new markers that were closely linked to the flavonoid 3',5'-hydroxylase gene copies and served to characterize that region more precisely. The marker BES9C17-FM was located on the physical map downstream of SSR VrZag30 and identified only BAC clones belonging to the contig ctg313. BES38C24-RM matched uniquely the BAC clones 38C24 and 21H16 belonging to ctg1320. Conversely, BES40A22-FM and BES38C24-FM primers were designed on BES belonging to ctg313 and ctg1320, respectively, but cross-identify BAC clones assembled in both contigs. SSCP analysis of BES40A22-FM and BES38C24-FM fragments revealed that for each marker two haplotypes are present in ctg313 and two different haplotypes are present in ctg1320. Yet, marker BES17K04-FM matched BACs 6H20, 9C17 and 31P20 of ctg313 and BACs 29J23, 38C24, 4K10, 30K14 of ctg1320. SSCP analysis identified four different haplotypes at the regions amplified by BES17K04-FM primers: two haplotypes were detected in the BAC clones of ctg1320, one more different haplotype was present in ctg313, and yet another SSCP haplotype was found in both BAC17K04 and BAC24N06 belonging to ctg2373. The occurrence of more than two haplotypes of the flavonoid 3',5'-hydroxylase gene fragments and of the BES markers in a diploid genotype confirmed the presence of paralogous copies of the flavonoid 3',5'-hydroxylase genes and rule out the possibility that the individual contigs represent allelic regions that have not been properly

Figure 7
Integrated genetic and local physical map spanning the complex locus VvF3'5'H on LG6. Contigs were assembled using FPC and BAC clones were further aligned using BES-derived markers. Contigs were anchored to the 'Cabernet Sauvignon' and the V. vinifera consensus genetic maps using cross-referenced SSR markers. The orientation of each contig is arbitrary. Genetic and physical distances are not drawn to scale. Names written in horizontal refer to BAC clones and contigs; those written in vertical refer to markers and genes. The presence of a given marker on a BAC clone is highlighted with a symbol and different SSCP haplotypes for each marker are shown with different symbols (■ ◆ ● ○ △). The symbol ‘X’ stands for lack of amplification of a given marker in the corresponding BAC clone. BAC clones missing any symbols were not tested with the corresponding marker. Physical localisation of VvF3'5'H-1 and VvF3'5'H-2 loci is shown in grey boxes.
assumed. A segmental duplication spanning the markers F35-1int, F35-2int, BES40A22-FM, BES17K04-FM and BES38C24-FM that occurred within 2.2-3.0 cm on LG6 could explain the physical organization of flavonoid 3',5'-hydroxylase isogenes as it has been outlined in Figure 7. The contig ctg2373 that comprised only three BAC clones could not be unambiguously positioned in relation to the other contigs. BES markers from the other contigs did not match any BAC clones of ctg2373 but marker BES17K04-FM matched both ctg313 and ctg1320. Marker BES17K04-FM is therefore located within the segmental duplication. It remains uncertain whether ctg2373 partially overlaps the region of ctg313 and split from it due to high heterozygosity or if it actually covers an adjacent locus. Under the assumption of BAC alignment between BAC clones 9C17 and 17K04 that is supported by genetic co-segregation of the corresponding BES and by co-amplification by marker BES17K04-FM, ctg2373 would carry the haplotype allelic to BACs 6H20, 9C17, 31P20 at the BES17K04-FM marker BES17K04-FM, ctg2373 would carry the haplotype and only slightly detectable in fully expanded leaves. It was also present in all berry tissues. The other sequence VvF3'H-2, which translated into a truncated flavonoid 3'-hydroxylase due to a 23-bp frameshift deletion, was apparently not expressed and is likely to be a pseudogene.

Transcripts of VvF3'5'H-1 were present in all tissues. High levels of expression were detected in pigmented apical leaflets, flowers, seeds and berries but not in green fully expanded leaves and roots. Differential expression could be detected between green berry skin and red berry skin, the expression being higher in pigmented berry skin. The expression pattern of VvF3'5'H-1 mimicked the expression profile of Chs2, Chs3 and the gene coding for flavanone hydroxylase (F3'H). The gene VvF3'5'H-2 expressed transcripts of different size that could be distinguished on agarose gel (VvF3'5'H-2-short, VvF3'5'H-2-medium and VvF3'5'H-2-long). The two shorter bands were invariably present in all tissues, even though at a different level of expression. The longest band was strongly expressed only in red berry skin, whilst it was slightly detectable in seeds and not detectable at all in other tissues.

Based on the data available to us, we propose that the chromosomal region spanning the VvF3'5'H gene cluster includes four copies of VvF3'5'H-1 and five to seven copies of VvF3'5'H-2, all sharing a high sequence similarity. We were able to present a preliminary physical map of this cluster of genes by a combination of computational assembly of fingerprinted BACs and by PCR-based alignment of BAC clones. The region has likely originated from both tandem duplications that generated nearby gene copies (i.e. paralogs within BACs 28B15 and 99I17 in the ctg313) and from segmental intrachromosomal duplications involving a larger block of DNA (i.e. blocks of paralogs duplicated into ctg313 and ctg1320). Segmental duplications are an important source of gene evolution [25] and they have been estimated to cover 1.7–2.0% of the mouse genome [26] and 4% of the human genome [27]. In the rat genome, duplicated genes are largely confined into <1 Mb intrachromosomal duplications rather than widely interspersed through the genome [28]. A similar model of genome dynamics could also have shaped plant genomes as it is emerging for model species [29].

**Expression analysis of VvF3'H and VvF3'5'H along with major structural genes of the flavonoid pathway in different grape tissues**

The genes encoding both flavonoid 3'-hydroxylase and flavonoid 3',5'-hydroxylases are widely expressed in grapevine organs (Figure 8). Transcripts of gene VvF3'H-1 were detected in all tissues tested. VvF3'H-1 was strongly expressed in green apical leaflets and flowers but it was only slightly detectable in fully expanded leaves. The expression of most flavonoid genes, including those coding for flavonoid hydroxylases, was more intensively activated in flowers and seeds than in other organs. Seeds expressed all isofoms of the flavonoid hydroxylase genes identified in this work, whilst all but VvF3'5'H-2-long were expressed in flowers. Among berry tissues, flesh poorly expressed the genes of the flavonoid pathway, except for BAN, VvF3'H-1, VvF3'5'H-1, VvF3'5'H-2-long and VvF3'5'H-2-medium. In the epicarp of berries collected at different developmental stages, Fls1 and BAN were more expressed at the pre-véraison stage; Chs3, F3'H and VvF3'5'H-1 were induced in red skin compared to green skin; UFGT, VvF3'H-1 and VvF3'5'H-2-long were strongly expressed in red skin but were undetectable in green skin.

The flavonoid pathway does not exclusively lead to the synthesis of anthocyanins and several flavonoid hydroxylase isogenes were therefore expressed also in non-pigmented tissues. Berry flesh and seeds accumulate non-anthocyanin flavonoids such as proanthocyanidins and catechins. Their synthesis is catalysed downstream of the flavonoid hydroxylases by the genes DFR, LDOX and BAN. Gene BAN was strongly expressed in seeds, berry flesh and berry skin at pre-véraison, whilst was almost undetectable in post-véraison berry skin at the stage when the rate of accumulation of proanthocyanins and catechins is declining (Figure 9). Sunlight exposed tissues expressed the gene coding for flavonol synthase (Fls1). Flavonol synthase converts the products of flavonoid
hydroxylases into the UV-protecting flavonols. Inner tissues (seeds and berry flesh) and underground organs (rootlets) did not exhibit detectable levels of Fls1.

Expression profiling of flavonoid hydroxylase genes in skin of ripening berries

The time-course of expression of flavonoid hydroxylase genes in the red-skinned cultivar ‘Merlot’ revealed that gene VvF3’H-1 is expressed throughout ripening. High transcript levels were detected from the first sampling date at the stage when catechins are rapidly synthesised and peak at 5.6 mg g\(^{-1}\) of berry skin (Figure 9). VvF3’H-1 expression decreased 8–10 weeks after blooming, at the ripening stage when most genes committed to the flavonoid pathway have been reported to level off [1]. VvF3’H-1 transcripts increased soon after, at the onset of véraison (August 12\(^{th}\)), which were at that point in time 2-fold higher in red berries than in green berries collected from the same bunch. VvF3’H-1 expression lasted till harvest, with a secondary peak of expression one week before harvest, concurrently at the peak of anthocyanin content.

The gene VvF3’5’H-1 was weakly expressed during the earliest ripening stages. At the onset of véraison, transcripts of VvF3’5’H-1 were poorly detectable in green berries, while they increased 50-fold in berries that had already turned red within the same bunch. The peak of VvF3’5’H-1 expression corresponded to August 24\(^{th}\), at the stage when all berries had turned red. As described for VvF3’H-1, a weaker peak of expression was found one week before harvest. Expression of VvF3’5’H-2 was checked by semi-quantitative PCR because the different isoforms of this gene were distinguishable by size (Figure 10). Transcripts of the isoforms VvF3’5’H-2-long and VvF3’5’H-2-medium did appear at the onset of véraison and lasted till full ripening, whilst the isogene VvF3’5’H-2-short was constitutively expressed throughout the ripening.

Accumulation, composition of anthocyanins and colour evolution in berry skin during the progress of ripening

Biometric and analytical parameters have been used to follow the progress of ripening (Figure 9). The onset of ripening began on August 3\(^{rd}\) when sugars started to accumulate and acidity commenced to decline. The onset of véraison (August 12\(^{th}\)) was considered as the time when 10% of the berries had turned red (\(H^* = 102.98, C^* = 18.27\)). Bunches went through this phase of colour transition until August 24\(^{th}\) when all berries had turned red. As described for VvF3’H-1, a weaker peak of expression was found one week before harvest. Expression of VvF3’5’H-2 was checked by semi-quantitative PCR because the different isoforms of this gene were distinguishable by size (Figure 10). Transcripts of the isoforms VvF3’5’H-2-long and VvF3’5’H-2-medium did appear at the onset of véraison and lasted till full ripening, whilst the isogene VvF3’5’H-2-short was constitutively expressed throughout the ripening.

At the earliest stage of the véraison, cyanidin-based anthocyanins (cyanidin and peonidin) were roughly as abundant as delphinidin-based anthocyanins (delphinidin, malvidin and petunidin) (42% versus 58%). Then, on August 24\(^{th}\) the percentage of delphinidin-based anthocyanin (79%) peaked simultaneously with the highest level of expression of VvF3’5’H-1. The fraction of tri-substituted anthocyanins slightly decreased in the following weeks (77–78%), then remaining substantially unchanged until harvest (75%). Berry skin colour moderately varied from \(H^* = 331.55, C^* = 3.83\) to \(H^* = 306, C^* = 2.59\) from September 2\(^{nd}\) to September 28\(^{th}\). Anthocyanin composition in berry skin changed over the period of
Transcript profiling of \textit{VvF3'H} and \textit{VvF3'5'H-1} in ripening berries. Ripening curve of \textit{V. vinifera} 'Merlot' was based on analytical parameters (berry weight, soluble solids, titratable acidity expressed as tartaric acid equivalents, skin total phenols expressed as (+)-gallic acid, skin catechins expressed as (+)-catechin, skin anthocyanins) at the following sampling dates: 1. July 15th, 2. July 29th, 3. August 3rd, 4. August 9th, 5. August 12th, 6. August 24th, 7. September 3rd, 8. September 13th, 9. September 20th, 10. September 28th. Concomitant gene expression of \textit{VvF3'H} and \textit{VvF3'5'H-1} in berry skin was assessed by quantitative RT-PCR and expressed as the ratio between the \textit{C}_T of the gene under study and the \textit{C}_T of the actin gene. Bars represent ± s.e.
called red-skinned cultivars. Skin colour from brick red to dark blue among the so
complexity was not unexpected for species like grape
plants that are known to accumulate flavonoids, such as
leaves, flowers, roots, seeds and, particularly, in skin of
berries in the grapevine genome and are organised in com-
plex gene clusters that most likely resulted from gene
duplication and intra-chromosomal segmental duplica-
tion. These genes are expressed in coordination with other
genes which emerged from fine structural analysis at phys-
ical level, has first been shown for plant hydroxylases. This
phenolic compounds in the red-skinned variety 'Merlot'. Apical leaflets of 1 cm² surface
area, fully expanded leaves (>10 cm²) and adventitious
rootlets were collected from canes grown in hydroponic
culture for two months. Flower buttons were collected
one week before blooming, seeds and flesh were extracted
from berries collected two weeks after véraison. Transcript
and metabolite profiling in berry skin was performed on
samples of 40 berries collected at ten ripening stages from
pre-véraison (July 15th, 2004) till harvest (September 28th,
2004) from plots of twelve vines replicated four times.

**Cloning of flavonoid 3'- and 3',5'-hydroxylase genes with
the help of degenerate primers**

Degenerate primers were designed on two specific and
conserved regions of plant F3'H and F3',5'H genes (Table
1, Figure 1). PCRs were performed in a 40 µl volume con-
taining 200 µM of each dNTP, 2.5 µM of each primer, 1 U
of HotMaster Taq polymerase (Eppendorf) with 2.5 mM
MgCl₂ buffer, and 40 ng of template DNA. The denatura-
tion step was 95°C for 1 min, followed by 40 cycles of
92°C for 45 s, 60°C for F3'H primers and 56°C for
F3',5'H primers for 50 s, 65°C for 90 s. Amplified bands
were extracted from agarose gel with Nucleospin Extract
(Macherey-Nagel) and cloned into pGEM-T vector
(Amersham Biosciences).

**Sequence analysis**

Nucleotide and amino acid sequences were compared
with GenBank accessions, with EST singletons and tenta-
tive consensus sequences (TCs) held at the TIGR grape
genome database. Conceptual translation was obtained with
DNAclub software. Amino acid entries of 670 plant cyto-
chrome P450 monoxygenases were downloaded from
GenBank as of November 30th, 2004. Cluster analysis of
plant cyt P450s and the candidate grape flavonoid
hydroxylases was done with the Neighbor-joining
method.

**Genetic mapping**

Nested primers were designed to amplify an exon of
grapevine flavonoid 3'- and 3',5'-hydroxylase genes (F3,
F3-1, F35-1, F35-2, see Table 1 and 2). Additional primer
pairs (F3int, F35-1int and F35-2int) were designed in the
exon sequences flanking an intron for detecting SNPs use-

**Plant material**

Genomic DNA was extracted from young leaves [30].
‘Cabernet Sauvignon’ was used for cloning flavonoid 3'-
and 3',5'-hydroxylase genes. Forty six full-sibs of the cross
‘Cabernet Sauvignon’ × *Vitis* ‘breeding line 20/3’ were
used for genetic mapping. Gene expression was analysed
in the red variety ‘Merlot’. Apical leaflets of 1 cm² surface
area, fully expanded leaves (>10 cm²) and adventitious
rootlets were collected from canes grown in hydroponic
culture for two months. Flower buttons were collected
one week before blooming, seeds and flesh were extracted
from berries collected two weeks after véraison.
ful for mapping in untranslated regions (Table 1 and 2). Flavonoid hydroxylase gene fragments containing the intron were amplified in a 10 µl volume containing 200 µM of dNTPs, 0.4 µM of each primer, 0.5 U of Taq polymerase, 2.5 mM MgCl₂, and 20 ng of template DNA. The PCR profile was 35 cycles of 92°C for 50s, annealing temperature as reported in Table 2 for 50s, 65°C for 1 min. Gene fragments were separated on 4% glycerol and 0.5 × MDE polyacrylamide gel (BioWhittaker Molecular Applications) for SSCP analysis, run at room temperature for 16 hours at 6 W and stained with silver nitrate. The segregation of F3int, F35-1int, F35-2int and 335 SSR markers was followed in the ‘Cabernet Sauvignon’ × *Vitis* ‘breeding line 20/3’ full-sib family [G. Di Gaspéro, unpublished data]. Linkage groups of markers segregating from the ‘Cabernet Sauvignon’ parent were identified using Carthagegene with a LOD of 5.0 [31]. Marker order was calculated using the command Build 3.

**Local physical maps**

A 6X BAC library of ‘Cabernet Sauvignon’ was screened by PCR with the F3, F35-1 and F35-2 primer pairs (Table 2) using a 3D pooling strategy [23]. Then, positive individual clones were pre-cultured in 2X LB medium for 20 hours and 2 µL of cell suspension were inoculated into the same medium and grown for 16 hours. Cultured clones were directly used as PCR template. Positive BACs were confirmed using the corresponding F3int, F35-1int and F35-2int primer pairs (Table 2). PCR products were then run into a SSCP gel (as above) for identifying the haplotype of each BAC clone. A set of 30,151 BAC fingerprints of the ‘Cabernet Sauvignon’ BAC clones [S. Paillard, unpublished data] were assembled into contigs using FPC v8.0 [24]. The first assembly was made at a cut-off value of 10⁻⁵⁰ and with a tolerance of 0.4 bp. After several steps of dQing and merging, 2,035 BAC contigs were obtained [S. Paillard, unpublished data]. BAC clones that were positive at the PCR screening were localized over the contigs. Clone order was determined by running the consensus band (CB) algorithm on each individual positive contig. Contig size was estimated by converting CB units into band (CB) algorithm on each individual positive contig. PCR with the F3, F35-1 and F35-2 (Table 2) were used for expression analysis of *VvF3'H-1, VvF3'S'H-1* and *VvF3'S'H-2*, respectively. Each cDNA sample was analysed at two different dilutions and with two replicates per dilution. Gene transcripts were quantified comparing the threshold cycle *(Cₚ)* with that of the constitutive actin gene [32]. Semi-quantitative PCR was performed on different tissues upon cDNA normalisation based on the expression of the actin gene and visualised on agarose gel stained with ethidium bromide. Primers pairs for the genes of the flavonoid pathway were retrieved from the literature: primers for *Chs1, Chs2, Chs3, DFR, LDOX, UFGT* were from [33], primers for *Fls1* were from [34]. Primers for *F3H* and *BAN* were newly designed on the original gene sequences [35,36] to tag 150–200 bp gene fragments.

**Transcript profiling**

Total RNA was extracted from berry skin and flesh, leaves, flower bunches, rootlets and seeds following the procedure described in [31]. Total RNA was treated with 0.5 U/μg RQ1 DNase (Promega) in presence of an RNAs inhibitor (RNAGuard, Amersham Biosciences). First strand cDNA was synthesised using 2 µg of RNA, 0.5 µM (dT)₁₈ primer and 50 U of M-MLV Reverse Transcriptase (Promega). Quantitative real-time PCR was carried out on a DNA Engine Opticon2 (MJ Research) using SYBR Green Jump Start Taq ReadyMix (Sigma). The primer pairs F3-1, F35-1 and F35-2 (Table 2) were used for expression analysis of *VvF3'H-1, VvF3'S'H-1* and *VvF3'S'H-2*, respectively. Total RNA was extracted from berry skin and flesh, leaves, flower bunches, rootlets and seeds following the procedure described in [37]. The calibration curve was obtained with (+)-gallic acid (Sigma). Total flavan-3-ols (catechins) were quantified as reported in [38]. The calibration curve was constructed with (+)-catechin (Extrasynthase). Absorbance was measured by a UV/VIS LAMBDA 2 spectrometer (Perkin-Elmer). Anthocyanins for the HPLC analysis were extracted for 4 hours from 200 mg of berry skin with 2 mL of methanol, then centrifuged at 5,000 rpm for 15 min and filtered with a 0.2 µm PTFE filter (Chemtek Analytica). Total phenols were determined following the procedure of [39]. The calibration curve was obtained with (+)-gallic acid (Sigma). Total flavan-3-ols (catechins) were quantified as reported in [38]. The calibration curve was constructed with (+)-catechin (Extrasynthase). Absorbance was measured by a UV/VIS LAMBDA 2 spectrometer (Perkin-Elmer Series 4) using a C18 Purospher RP-18 (5 µm, 250 mm × 4 mm) column (Merck) protected by a C18 guard column, according to the procedure reported...
Anthocyanins were detected by a LC-95 spectrophotometric detector (Perkin Elmer) operating at 520 nm. Anthocyanin content was expressed as mg/L of malvidin 3-glucoside upon the construction of a standard curve. The composition of monoglucoside anthocyanins was used for calculating the ratio of tri-/di-substituted derivatives. Berry colour was measured with an X-Rite 948 Chromameter (X-Rite) and averaged over 160 berries at each sampling date. Colorimetric specification was referenced to the CIELab scale. Hue angle (H°) and chroma (C°) were calculated according to [40].

Authors’ contributions
SDC participated in the design of the study, carried out gene cloning, expression profiling, field experiments and analyses of metabolites, participated in the interpretation of the results. GDG conceived the study, coordinated the experiments, carried out genetic mapping, interpreted the results and drafted the manuscript. RM managed and screened the BAC library. AN carried out and interpreted experiments, carried out genetic mapping, interpreted the results and drafted the manuscript. EP critically revised the manuscript. AN carried out and interpreted the expression analysis. RT revised the manuscript. SPK and DS critically revised the manuscript. JH and RT revised the manuscript. All authors read and approved the final manuscript.

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