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Molecular characterization of berry skin color reversion on grape somatic variants

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Abstract.

BACKGROUND: During grapevine domestication somatic variation has been used as a source of diversity for clonal selection.

OBJECTIVE: This work provides additional information on the molecular mechanisms responsible for berry skin color reversion on a subset of somatic variants for berry skin color never investigated before.

METHODS: The berry color locus and its surrounding genomic region were genetically characterized through a layer-specific approach, which has already been proven to be a successful method to decipher the molecular mechanisms responsible for color reversions on somatic variants.

RESULTS: Deletions of different extent and positions were detected among less-pigmented/unpigmented variants derived from a pigmented wild-type. These deletions affected only the inner cell layer in the less pigmented variants and both cell layers in the unpigmented variants. Regarding the pigmented variants derived from an unpigmented wild-type, only one group was distinguished by the *Gret1* retrotransposon partial excision from the *VvMybA1* promoter. Moreover, within this latter group, *VvMybA2* showed an important role regarding the phenotypic variation, through the recovery of the functional G allele.

CONCLUSIONS: This investigation focused on the berry color locus using somatic variants for berry skin color, promotes a better understanding of the evolutionary events behind their origin and variability, opening an opportunity for their use in the genetic improvement of varieties.

Keywords: Berry skin color, grapevine, L1 and L2 cell layers, MYB genes, somatic mutation, SSR markers

1. Introduction

Domestication of the wild *Vitis vinifera* ssp. *sylvestris* to *Vitis vinifera* ssp. *sativa* species was a complex, long-term, and ongoing evolutionary process that involved multiple loci [1]. This process dramatically changed the grape biology in order to ensure higher sugar content, greater yield, regular production and adaptation to different environments [2].

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Today, most of the cultivated grapevine varieties are the result of continuous selection and vegetative propagation over centuries, spontaneously generated by sexual reproduction and via somatic mutations. These events led to grape varieties which have enough similar vegetative and reproductive features to allow them to freely interbreed [3]. Clones are usually selected for a particular character or combination of characters which, when propagated by appropriate means, retain those characters and are genetically identical, i.e., true-to-type [4].

Although vegetative propagation has been used as a conservative strategy to maintain and propagate clones, somatic mutations may occur naturally when a spontaneous morpho-physiological mutation appears on a shoot, leading to clonal polymorphism [3, 5]. When a mutation appears in a single cell of a shoot apical meristem layer, the following mitotic divisions lead to the emergence of a mutated section [6], giving rise to a periclinal chimera. These structures corresponds to specific types of genetic mosaicism in which one or two entire meristematic cell layers are genetically different from the others, and are developmentally independent from the adjacent layers [7]. Usually they are stable and do not threaten plant fitness, and are maintained by vegetative propagation. However, occasionally these cellular rearrangements in the periclinal chimera lead to homogenization of the genotype to the whole plant [5] in a two-step process. First, a somatic mutation emerges in one cell of the shoot apical meristem and then is propagated by cell division to an entire cell layer, generating a stable chimeric structure. Subsequently, the replacement of one cell layer by another can lead to a homogenization of the genotype of the whole plant, representing the second step of the process [8]. Therefore, divergent genotypes and, to some extent divergent phenotypes, can appear due to these molecular and cellular mechanisms, representing a valuable source of heritable variation that has been widely exploited by winemakers [5, 9]. There are many examples of spontaneous variant traits in grapevines, including berry skin and flesh color or flavor, size and compactness of bunches, canopy growth, date of ripening and productivity [3, 5].

At the genetic level, the most well-documented polymorphisms leading to qualitative variation within varieties are those that affect berry skin color. Currently, several pigmented varieties have certified clones with different skin color shades, such as Aramon, Grenache, Pinot or Terret. Moreover, unpigmented skin berry varieties such as Savagnin, Chardonnay and Chasselas also comprise pigmented clones [5].

The first comparative study of molecular polymorphisms arising during vegetative propagation on the whole genome scale was done on spontaneously generated grape clones. Despite the small number of SNP (single nucleotide polymorphism) and indel events observed, mobile elements were identified as being involved in most of the polymorphisms detected [10].

Overall, the grapevine somatic variants for berry skin color can be divided in two main groups: A) unpigmented and less-pigmented variants from a pigmented ancestor, namely white and grey/pink-skinned variants derived from a black-skinned ancestor; B) pigmented variants from an unpigmented wild-type, namely red/pink-skinned variants derived from a white-skinned wild-type [11–13].

Research on grape berry color shows that the color locus is a cluster of four *Myb* and *Myb*-like genes, spread along a 200 kb-region located on chromosome 2, which comprises *VvMybA1* and *VvMybA2*, two adjacent transcription factors that regulate the transcription of *VvUFGT* (UDP-glucose:flavonoid 3-O-glucosyltransferase), a key point in the anthocyanin pathway. In white-skinned cultivars, the absence of anthocyanins has been related with the insertion of the *Gret1*, a 10,422 bp long retrotransposon, in the promoter region of *VvMybA1* gene combined with two non-conservative mutations in the coding sequence of *VvMybA2*, a non-synonymous point mutation that leads to an amino acid substitution (change of arginine residue at position 44 in the red allele [G] altered to leucine in the white allele [T]) and a 2-bp deletion (CA) altering the reading frame, that results in a smaller protein [14–17]. An additional point mutation of *VvMybA2* gene (C22) related to berry pigmentation was described by Carrasco et al. [18] in wild grapevine accessions with low anthocyanin content. These mutations in both genes lead to the loss of transcription factors expression, which consequently prevents anthocyanin biosynthesis.

Researchers have recently focused on the somatic variation affecting berry skin color in different varieties, using a layer-specific approach to identify the molecular mechanisms responsible for the polymorphisms occurring

at berry color locus and to understand the evolutionary events behind their origin [5, 13, 19]. Vezzulli et al. [19] observed a homozygous-like region in Pinot Gris and Pinot Blanc clones, suggesting that the mutation impairing the color locus was directly related with the presence of deletions, extending for at least 4.2 Mb for the L2 cell layer of Pinot Gris and ranging from 100 to 179 kb for both cell layers of Pinot Blanc. Based on these results, they concluded that Pinot Noir gave rise to Pinot Blanc and Pinot Gris independently, suggesting a parallel evolutionary model. Considering the same family of somatic variants, Pelsy et al. [5] investigated a collection of Pinot Noir, Pinot Gris and Pinot Blanc clones and proposed a model integrating both mutation and cell layer rearrangements to explain the mechanism of clone diversification. On the other hand, Migliaro et al. [13] identified polymorphisms that enable the distinction of less pigmented/unpigmented and pigmented mutants from the corresponding wild-type genotype, concluding that phenotypic variation is due to deletion events that can result in either gain- or loss-of-gene function. Deletions of different extent and position were detected among less pigmented/unpigmented variants derived from a pigmented wild-type. Regarding the pigmented variants derived from an unpigmented wild-type, the same authors observed that the main mechanism for color gain was the partial *Gret1* retrotransposon excision from the *VvMybA1* promoter, leaving only the solo-3'LTR region.

The present study was focused on the genetic characterization of the berry skin color locus and its surrounding genomic region, aiming to uncover the molecular mechanisms responsible for the skin color phenotype reversion on a set of somatic variants for berry skin color, analyzed here for the first time. The results obtained represent an important addition to the current knowledge about the structural dynamics along the distal arm of chromosome 2 and the evolutionary events behind the origin of somatic variants for berry skin color.

2. Materials and methods

2.1. Plant material

The molecular mechanisms of color variation in grapevine somatic variants for berry skin color were analyzed in 26 accessions belonging to 10 *V. vinifera* spp. *sativa* varieties, including the Pinot family (Pinot Noir, Pinot Gris and Pinot Blanc) which was used as a reference (Table 1). The different accessions were provided by the Institut National de la Recherche Agronomique – INRA, Centre of Grapevine Genetic Resources (Montpellier, France); Universidade de Trás-os-Montes e Alto Douro – UTAD (Vila Real, Portugal) and the Instituto Nacional de Investigação Agrária e Veterinária – INIAV, Coleção Ampelográfica Nacional (Dois Portos, Portugal). All accessions were divided in two groups (A and B) according to their botanical origin (Table 1).

2.2. Genomic DNA extraction

To implement the layer-specific approach, two genomic DNA samples for each accession were isolated: one from 100 to 200 mg of young leaf (L1+L2 derived tissue), corresponding to stage E (2 to 3 leaves unfolded) of the Baggioolini scale, and the other from 200 to 300 mg of woody shoot pith (L2 derived tissue) at stage A (winter bud) of the same scale, for a total of 52 tissue-specific DNA samples. The leaf and woody shoot pith materials were ground in a TissueLyser II (Qiagen, Hilden, Germany) and DNA extraction was performed using the DNeasy 96 Plant Kit (Qiagen, Hilden, Germany), according to the manufacturer's protocol. In each analysis, the genetic difference between the L1+L2 (leaf) and L2 (woody shoot pith) corresponds to the make-up of L1 cell layer. DNA concentration was determined by UV spectrometer (Nanodrop® ND-1000, ThermoFisher Scientific, USA). DNA quality was also checked with a 1% (w/v) agarose gel electrophoresis using 1x TBE buffer followed by ethidium bromide staining. Necessary dilutions were performed (approximately 10 ng/μL) and kept at 4° C for further use. Subsequent molecular analyses were performed independently on both DNA samples from each accession.

Table 1
List of the studied somatic variants for berry skin color divided in two groups (A and B), respective berry skin color, repository and accession number

Cultivar	Berry skin color ¹	Repository ²	Accession number
<i>Group A</i>			
Aramon	B	INRA/CGGR	22Mtp1
Aramon Gris	G	INRA/CGGR	23Mtp2
Aramon Blanc	W	INRA/CGGR	24Mtp1
Aspiran Noir	B	INRA/CGGR	1Mtp3
Aspiran Gris	G	INRA/CGGR	3Mtp2
Aspiran Blanc	W	INRA/CGGR	2Mtp1
Grolleau	B	INRA/CGGR	297Mtp11
Grolleau Gris	G	INRA/CGGR	305Mtp2
Grolleau Blanc	W	INRA/CGGR	356Mtp2
Pinot Noir	B	UTAD	F2.2.3
Pinot Gris	G	UTAD	F2.13.2
Pinot Blanc	W	UTAD	F2.13.1
Pique-poul Noir	B	INRA/CGGR	11Mtp8
Pique-poul Gris	G	INRA/CGGR	12Mtp5
Pique-poul Blanc	W	INRA/CGGR	13Mtp7
Terret Noir	B	INRA/CGGR	14Mtp3
Terret Gris	G	INRA/CGGR	15Mtp2
Terret Blanc	W	INRA/CGGR	16Mtp3
<i>Group B</i>			
Folgasao Roxo	R	INIAV/CAN	52709
Folgasão	W	INIAV/CAN	52708
Malvasia Cândida Roxo	R	INIAV/CAN	50911
Malvasia Cândida	W	INIAV/CAN	50810
Malvasia Fina Roxo	R	UTAD	F2.3.1
Malvasia Fina	W	UTAD	F2.13.5
Mourisco	B	INIAV/CAN	52002
Mourisco Roxo	R	INIAV/CAN	52001

¹ B – Black; G – Grey; R – Red; W – White. ²INRA – Institut National de la Recherche Agronomique, CGGR – Centre of Grapevine Genetic Resources (Montpellier, France); UTAD - Universidade de Trás-os-Montes e Alto Douro (Vila Real, Portugal); INIAV - Instituto Nacional de Investigação Agrária e Veterinária, CAN – Coleção Ampelográfica Nacional (Dois Portos, Portugal).

2.3. Single sequence repeat (SSR) analyses

To determine whether the berry color variants used in the study were true-to-type, i.e. if all berry skin color variants were actually bud sports of the original variety (wild type) genotype, a set of 10 markers (VVS2, VVMD5, VVMD7, VVMD27, VrZAG62, VrZAG79, VMC4f3, VVMD28, VVMD32 and VVIv67), which included the Organisation Internationale de la Vigne et du Vin (OIV) core set established by the European Project GENRES#81 for grapevine cultivars identification, was used. Two multiplex PCRs were carried out with the OIV SSR core set as reported by Castro et al. [20], the first one involving VVS2, VVMD5 and VVMD7 (set A), and the second VVMD27, ssrVrZAG62 and ssrVrZAG79 (set B). Individual reactions were performed with the remaining four

primer pairs (VMC4f3, VVMD28, VVMD32 and VVIv67) according to the same authors. The SSR allelic profile of each accession was compared with its published reference profile (Table 2).

Another set of 10 markers (SC8.0146.010, SC8.0146.026, VVNTM1, VVNTM2, VVNTM3, VVNTM4, VVNMT5, VVNTM6, VVIU20, VMC7G3) surrounding the berry color locus and distributed along the distal arm of chromosome 2 was used to investigate polymorphisms in this region (Table 2). PCR reactions were performed as described by Vezzulli et al. [19]. Capillary electrophoresis was carried out in an ABI 3130xl Genetic Analyzer (Life Technologies, Foster City, CA, USA) and the fragments were sized with Peak Scanner V1.0 software (PE Applied Biosystems, CA, USA) using the GeneScan 500 LIZ size standard (Life Technologies) as an internal ladder.

2.4. *VvMybA1* and *VvMybA2* gene structure

The *VvMybA1* and *VvMybA2* gene polymorphisms (the detection of functional and non-functional alleles) were also investigated.

Regarding characterization of the *VvMybA1* locus, the presence or absence of *Gret1* retroelement in the *VvMybA1* promoter region was detected using the primers *a* (5'-AAAAAGGGGGGCAATGTAGGGACCC-3') and *d3* (5'-CCTGCAGCTTTTTCG GCATCT-3') as described by Lijavetzky et al. [14]. PCR amplifications and detection of amplified fragments were performed as reported in Lijavetzky et al. [14]. In order to detect putative functional alleles, the primers used were *F2* (5'-GGACGTTAAAAAATGGTTGCACGTG-3') described by Azuma et al. [21] and *R1* (5'-GAACCTCCTTTTGAAGTGGTGACT-3') by Lijavetzky et al. [14]. PCR amplifications and detection were performed as reported in Azuma et al. [21] and Carrasco et al. [18], respectively.

For the *VvMybA2* gene, the single nucleotide polymorphism (SNP) related to berry color *VvMybA2R44* [11] was investigated by a SNaPshot assay. *VvMybA2* amplification, SNP genotyping by ABI PRISM SNaPshot Multiplex kit (Life Technologies Corporation, Carlsbad, California) and SNP detection on ABI PRISM 310 Genetic Analyzer Sequencer were performed, as reported in Carrasco et al. [18].

3. Results

3.1. Varietal identification

Ten SSR loci were used to ascertain the genetic identity of the somatic variants and their relative ancestor to confirm their trueness-to-type. This fingerprinting system revealed that the pigmented and unpigmented variants of each variety are closely related, showing the same genetic profile, which confirms that all the berry skin color variants analyzed were bud sports of their respective wild-type genotype. Cultivar identification was achieved by comparing each SSR profile with the European *Vitis* database (<http://www.eu-vitis.de/>) or public profile (Table S1).

3.2. *VvMybA1* and *VvMybA2* allelic composition

To characterize the structural dynamics of the berry color locus two *Myb*-related genes, *VvMybA1* and *VvMybA2*, were analyzed, since they are known to be functionally involved in berry pigmentation.

Regarding these two *Myb*-related genes, their allelic composition allowed inferring some of the color differences observed among the studied variants. For all the wild-type pigmented ancestors of the group A, the *VvMybA1* promoter was heterozygous, containing both the *Gret1* (non-functional *VvMybA1*^{ITA}) and non-*Gret1* (functional *VvMybA1*^{AFL}) alleles in the two cell layers, as well as in the L1+L2 cell layer of the red/grey-skinned derivative variants (Table 3). In turn, only the *Gret1* (non-functional *VvMybA1*^{ITA}) allele, present in

Table 2
Primer information of the microsatellite markers used to all the studied loci

SSR Locus	LG	Use	5'-3' Forward primer sequence	5'-3' Reverse primer sequence	Reference
VVS2 ^{1,2}	11	True-to-type confirmation	CAGCCCGTAAATGTATCCATC	AAATTCAAAAATTCTAATTCAACTGG	Thomas and Scott (1993); This et al. (2004)
VVMD5 ^{1,2}	16	True-to-type confirmation	CTAGAGCTACGCCAATCCAA	TATACCAAAAATCATATTCTCTAAA	Bowers et al. (1996, 1999); This et al. (2004)
VVMD7 ^{1,2}	7	True-to-type confirmation	AGAGTTGCGGAGAACAGGAT	CGAACCTTTCACACGCTTGAT	Bowers et al. (1996, 1999); This et al. (2004)
VVMD27 ^{1,2}	5	True-to-type confirmation	GTACCAGATCTGAATACATCCGTAAAGT	ACGGGTATAGAGCAACCGGTGT	Bowers et al. (1999); This et al. (2004)
VrZAG62 ^{1,2}	7	True-to-type confirmation	GGTGAAATGGGCACCGAACACACGC	CCATGTCTCTCTCTCAGCTTCTCAGC	Sefc (1999); This et al. (2004)
VrZag79 ^{1,2}	5	True-to-type confirmation	AGATTGTGGAGGAGGGAACAAACCG	TGCCCCCATTTTCAAACTCCCTTCC	Sefc (1999); This et al. (2004)
VMC4f3	12	True-to-type confirmation	AAAGCACTATGGTGGGTGTAAA	TAACCATACATGCATCAAGGA	Di Gaspero et al. (2000)
VVMD282	3	True-to-type confirmation	AACAATTCAATGAAAAGAGAGAGAGAGA	TCATCAATTTCTGATCTCTATTGTCTG	Bowers et al. (1999)
VVMD32 ²	4	True-to-type confirmation	TATGATTTTATAGGGGGGTGAGG	GGAAAGATGGGATGACTCGC	Bowers et al. (1999)
VVlv67	15	True-to-type confirmation	TATACTTCTCATAGGGGTTTCC	TTGGAGTCCCATCAAAATTCATCT	Merdinoglu et al. (2005)
SC8.0146.010	2	Delition delimitation	-	-	Adam-Blondon A.F., personal communication
SC8.0146.026	2	Delition delimitation	-	-	Adam-Blondon A.F., personal communication
VVNTM1	2	Delition delimitation	CCAGCCACTATTGTGTAAC	TGCACCGTATCAAGATCATGTTC	Fournier-Level et al. (2009)
VVNTM2	2	Delition delimitation	TACCTGCTAACAAATGCATTATG	TAITTTGGTTTTTCTAATAAGA	Fournier-Level et al. (2009)
VVNTM3	2	Delition delimitation	TGCTGACCTGAATCAITTTTCTACT	GATGTTCTGGAGAGATGCTTATC	Fournier-Level et al. (2009)
VVNTM4	2	Delition delimitation	TTTGCACTGACTGCTTGGTGTAT	CCCATTGCTAAACCCCTACTCCT	Fournier-Level et al. (2009)
VVNTM5	2	Delition delimitation	AGGAGGAATCCACATCAAAAAGA	TGATTCAAAGGAATAAATAACCATCA	Fournier-Level et al. (2009)
VVNTM6	2	Delition delimitation	CCTTCTTGGACACCCATACAAA	TTCCCTATCAACAAACTTGAGG	Fournier-Level et al. (2009)
VVIU20	2	Delition delimitation	ACAACCTTAATGCTTCTACCAA	TCACCATGGAGATTTTCTGTAG	Fournier-Level et al. (2009)
VMC7G3	2	Delition delimitation	TTACTAGTGTGCTGCTGCTCCA	TGCTTCTCTCTTTTCAACTTTCA	Merdinoglu et al. (2005) Pellerone et al. (2001)

¹OIV core set, ²VIVC database.

homozygosity, was detected in the L2 of red/grey-skinned variants and in both cell layers of unpigmented variants (Table 3).

The *VvMybA1* gene did not allow discrimination of most variants of group B where the wild-type ancestor corresponds to a white-skinned variant, since they share the same allelic composition for this gene (Table 4). The partial excision of the *Gret1* retrotransposon was only observed in the Mourisco variety of group B (Table 4). The presence of the functional G allele at *VvMybA2R44* position was detected in some less-pigmented light red variants (Malvasia Fina Roxo, Folgasão Roxo and Malvasia Cândida Roxo) derived from a white-skinned ancestor (Table 4).

3.3. Genetics behind black-to-grey and/or white skin color reversions

As previously mentioned, all the pigmented ancestor accessions of group A were heterozygous, containing the *Gret1* (*VvMybA1*^{ITA}) and non-*Gret1* (*VvMybA1*^{AFL}) alleles of *VvMybA1* in both cell layers (Table 3). Moreover, the clonal differences between the pigmented and the derived unpigmented variants was shown to be caused by deletions of different extents, resulting in the removal of the functional colored allele of both *VvMybA* genes, leading to the white phenotype, as was also observed by Migliaro et al. [13]. In cultivars derived from asexual reproduction, such as the ones analyzed here, the emergence of the homozygous-like pattern compared with the ancestor (usually heterozygous) can only be assigned to a deletion (grey-shaded region in Table 3). The homozygous-like term corresponds to the genetic hemizygous term (which refers to a null allele, called *VvMybA1*^{PNB}), which will be used henceforth.

The black-skinned Pinot Noir was homozygous for two microsatellite markers (VVNTM6 and VVNTM4) and heterozygous for eight, as well as for the *Gret1* retrotransposon insertion and *VvMybA2* SNP, in either L1+L2- or L2- cell layers. Pinot Gris differed from Pinot Noir in having only one allele in the region between the VVNTM1 and VMC7G3 SSR markers, as well as only the *Gret1* allele in L2-derived tissues. In turn, Pinot Blanc was hemizygous in a smaller region, between SC08_010 and VVNTM5, both on L1+L2-, and L2-derived tissues (Table 3). A similar situation was observed for Pique-poul and Aramon varieties, where both less-pigmented and unpigmented variants probably derived from the wild-type pigmented ancestor in an independent way due to the different deletion extent and position. Pique-poul Blanc differ from Pique-poul Noir since it has only one allele in the region between the SC8_010 and VVNTM6 SSR markers, as well as only the *Gret1* allele (*VvMybA1*^{ITA}) in both L1+L2- and L2-derived tissues. On the other hand, the hemizygous profile of Aramon Blanc was between the VVNTM2 and VVNTM6 SSR markers. Regarding the less-pigmented variants of both varieties, they only showed a difference from their wild-type pigmented ancestor in the L2 cell layer. Pique-poul Gris revealed the presence of only one allele in the region encompassed by the VVNTM1 and VVNTM6 markers, also affecting the non-*Gret1* allele. Aramon Gris, similar to the less pigmented variants already described, can only be distinguished from its wild-type cultivar in L2-derived tissues, between the VVNTM2 and VVNTM3 SSR markers (Table 3).

For Aspiran Gris and Aspiran Blanc, one allele was observed from VVNTM1 to VVNTM6 SSR markers in the L2 cell layer of Aspiran Gris and in the L1+L2-derived tissues and the L2 cell layer of Aspiran Blanc. The same deletion pattern was also detected for Terret Gris and Terret Blanc (Table 3). Finally, in Grolleau Gris, as in the two previous cases (Aspiran and Terret), the homozygous status was restricted to the L2 cell layer, this being the main difference when compared with its wild-type. This deletion extended, at least, from the *VvMybA2* marker region to VVNTM6 SSR marker. Due to the homozygosity of the black-skinned ancestor Grolleau upstream of the *VvMybA2* marker it was not possible to have an accurate resolution of the deletion on the 5' border. The same deletion pattern observed for Grolleau Gris was found in Grolleau Blanc, but in this color variant it was also present in L1+L2-derived tissues. Besides that, the homozygosity of the black-skinned ancestor for Grolleau and Terret in VVNTM4 SSR marker also prevents a more precise delimitation of the deletion on the 3' border (Table 3).

Table 3
Genetic profile of berry color locus and its surrounding genomic region on black-to-grey and/or white revertant somatic variants and its ancestors.

Cultivar	Layer	Berry skin color ¹	Molecular marker and genomic coordinate on chromosome 2 (in Mb)													
			SC8_010	SC8_026	VVNTM1	VVNTM2	VvMybA2R44	VvMybA1	VVNTM3	VVNTM5	VVNTM6	VVNTM4	VVIU20	VMC7G3		
			12674	12970	14149	14151	14181	14248	14288	14325	14330	14384	16539	18270		
Aramon	L1+L2	B	ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	he	he	he	he	
	L2		ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	he	he	he	he	
Aramon Gris	L1+L2	G	ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	he	he	he	he	
	L2		ho	he	he	he	T	Gret1	ho	he	he	he	he	he	he	
Aramon Blanc	L1+L2	W	he	he	he	he	T	Gret1	ho	ho	ho	he	he	he	he	
	L2		he	he	he	he	T	Gret1	ho	ho	ho	he	he	he	he	
Aspiran Noir	L1+L2	B	ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
	L2		ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
Aspiran Gris	L1+L2	G	ho	he	he	he	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
	L2		ho	he	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	
Aspiran Blanc	L1+L2	W	ho	he	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	
	L2		ho	he	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	
Grolleau	L1+L2	B	ho	ho	ho	ho	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
	L2		ho	ho	ho	ho	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
Grolleau Gris	L1+L2	G	ho	ho	ho	ho	T/G	Gret1/Non-Gret1	ho	he	he	ho	he	he	he	
	L2		ho	ho	ho	ho	T	Gret1	ho	ho	ho	ho	he	he	he	
Grolleau Blanc	L1+L2	W	ho	ho	ho	ho	T	Gret1	ho	ho	ho	ho	he	he	he	
	L2		ho	ho	ho	ho	T	Gret1	ho	ho	ho	ho	he	he	he	
Pinot Noir	L1+L2	B	he	ho	he	he	T/G	Gret1/Non-Gret1	he	he	ho	ho	he	he	he	
	L2		he	ho	he	he	T/G	Gret1/Non-Gret1	he	he	ho	ho	he	he	he	
Pinot Gris	L1+L2	G	he	ho	he	he	T/G	Gret1/Non-Gret1	he	he	ho	ho	he	he	he	
	L2		he	ho	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	
Pinot Blanc	L1+L2	W	he	ho	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	
	L2		he	ho	he	he	T	Gret1	ho	ho	ho	ho	he	he	he	

Table 4
Genetic profile of berry color locus and its surrounding genomic region on white-to-red revertant somatic variants and its ancestors.

Cultivar	Layer	Berry skin color ¹	Molecular marker and genomic coordinate on chromosome 2 (in Mb)															
			SC8.010	SC8.026	VVNTM1	VVNTM2	VvMybA2R44	VvMybA1	VVNTM3	VVNTM5	VVNTM6	VVNTM4	VVIU20	VMC7G3				
			12674	12970	14149	14151	14181	14248	14288	14325	14330	14384	16539	18270				
Folgasão Roxo	L1+L2	R	he	he	ho	ho	T/G	Gret1/?	ho	ho	ho	ho	ho	ho				
	L2		he	he	ho	ho	T/G	Gret1/?	ho	ho	ho	ho	ho	ho				
Folgasão	L1+L2	W	he	he	ho	ho	T	Gret1	ho	ho	ho	ho	ho	ho				
	L2		he	he	ho	ho	T	Gret1	ho	ho	ho	ho	ho	ho				
Malvasia Cândida Roxo	L1+L2	R	he	he	ho	ho	T/G	Gret1/?	ho	he	ho	ho	ho	ho				
	L2		he	he	ho	ho	T/G	Gret1/?	ho	he	ho	ho	ho	ho				
Malvasia Cândida	L1+L2	W	he	he	ho	ho	T	Gret1	ho	he	ho	ho	ho	ho				
	L2		he	he	ho	ho	T	Gret1	ho	he	ho	ho	ho	ho				
Malvasia Fina Roxo	L1+L2	R	ho	ho	ho	ho	T/G	Gret1/?	ho	ho	ho	ho	ho	ho				
	L2		ho	ho	ho	ho	T/G	Gret1/?	ho	ho	ho	ho	ho	ho				
Malvasia Fina	L1+L2	W	ho	ho	ho	ho	T	Gret1	ho	ho	ho	ho	ho	ho				
	L2		ho	ho	ho	ho	T	Gret1	ho	ho	ho	ho	ho	ho				
Mourisco Tinto	L1+L2	B	he	he	ho	ho	T/G	Non-Gret1	he	he	ho	ho	he	he				
	L2		he	he	ho	ho	T/G	Non-Gret1	ho	he	ho	ho	he	he				
Mourisco Roxo	L1+L2	R	he	he	ho	ho	T/G	Non-Gret1	he	he	ho	ho	he	he				
	L2		he	he	ho	ho	T/G	Non-Gret1	ho	he	ho	ho	he	he				

¹R – Red; B – Black; W – White. The grey background indicates the putatively homozygous regions. he – heterozygous; ho – homozygous; Gret1 – non-functional allele; ? – undetected functional allele; Non-Gret1 – functional allele.

3.4. Genetics behind white-to-red skin color reversion

The molecular characterization of group B (pigmented variants derived from an unpigmented wild-type) followed the same layer-specific approach described above for group A. No genetic differences between layers were found within each variety, comparing the wild-type ancestral and its corresponding variants based on the markers used in this study (Table 4).

Although showing the same genetic pattern in L1+L2- and L2- derived tissues, the red and black-skinned Mourisco Roxo and Mourisco Tinto cultivars were both homozygous for a putative functional allele, described as a double insertion of 111- and 44-bp within the promoter region of *VvMybA1*, which should correspond to *VvMybA1*^{SUB} allele. The remaining variants of group B were putatively homozygous for the *Gret1* insertion (*VvMybA1*^{ITA} allele) in the entire shoot apical meristem.

Regarding the results obtained for *VvMybA2*, all cultivars showed to be homozygous for the non-functional T allele, except Folgasão Roxo, Malvasia Cândida Roxo and Malvasia Fina Roxo, which carried the functional G allele (Table 4).

With the exception of Mourisco, in the remaining varieties of group B the markers used were homozygous and monomorphic along an extensive genomic region on the distal arm of chromosome 2. While in Malvasia Fina this putatively homozygous and monomorphic region is present along the entire distal arm of chromosome 2, in Folgasão and Malvasia Cândida this homozygosity is of a smaller length, restricted to the region between VVNTM1 and VMC4f3 and between VVNTM1 and VVNTM3 SSR markers, respectively (Table 4).

4. Discussion

4.1. True-to-type confirmation

One of the most challenging and unresolved problems for viticulture and germplasm management worldwide concerns the fact that grapevine is a species with alternative nomenclatures, with several cases of synonymies and homonymies. For this reason, it is particularly relevant to confirm that somatic variants correspond to true-to-type of the wild-type ancestor variety at the molecular level. Based on the results obtained by SSR genotyping, possible homonymy cases, as previously observed by Ferreira et al. [22] for cultivars of the varieties Alvarelhão, Bastardo, Carrega and Touriga, were discarded.

4.2. Berry skin color reversions associated to somatic variation

The genetic analysis of group A varieties plainly showed that the deletion length encompassing the non-*Gret1* allele (*VvMybA1*^{AFL}) of *VvMybA1* is highly variable among different varieties and also within the same variety, resulting in different color phenotypes. Thus, this mutational event on the distal arm of chromosome 2 is likely to happen in a repeated and independent way. All white-skinned variants of group A can be differentiated from the respective wild-type counterpart. In the white-skinned variants, both layers are affected by the deletion and, consequently, the hemizygous state is detectable in the entire shoot apical meristem. For the less-pigmented variants from group A, a layer-specific molecular approach is required to discriminate them from the pigmented ancestor, because the same hemizygous state described for the white-skinned variants was only detected in the L2 cell layer. Thus, all less pigmented variants showed a chimerical status, while in all unpigmented ones the deletion was conserved, involving the entire shoot apical meristem.

Although no differences in the deletion extension of grey- and white-skinned variants could be observed in Aramon, Aspiran, Groulleau and Terret varieties with the layer-specific molecular approach adopted, these data do not exclude the hypothesis that the white-skinned variant evolved from the grey-skinned one, or that

these variants evolved independently from their pigmented wild-type ancestor. Thus, it could be helpful to use an integrative analysis considering the available historical information. Migliaro et al. [13] concluded that Tempranillo Gris probably evolved from Tempranillo due to an independent mutational event, considering the historical documentation about the direct selection of Tempranillo Blanco from Tempranillo. The authors inferred that the loss of the functional allele associated with the berry skin color was followed by L1 colonization and displacement to the L2 cell layer by the mutant cell, avoiding the grey berry chimerical status. Therefore, the same process could have occurred for Aramon, Aspiran, Groulleau and Terret varieties where no differences in the deletion extension of grey- and white-skinned mutants could be detected. However, the sequential evolutionary model, described by Walker et al. [15] for Cabernet Sauvignon cannot be excluded since the grey-skinned berries could represent a periclinal chimera of the black-skinned wild-type ancestor. In turn, the white-skinned berries could be the bud sport of the grey-skinned through a process that includes a deletion in the berry color locus in the L2 cells, while the white berry phenotype results from a cellular rearrangement (displacement) in the grey-skinned berries where the L2 cell layer (unpigmented) replaces the L1 cells (pigmented). Furthermore, although considered a rare event due to the stability of the anticlinal cell divisions the opposite phenomenon, L1 cell invasion of the inner layer (a process called replacement), can also be considered [9].

The main mechanism found for color gain in cultivars that belong to group B is the partial excision of *Gret1*, called *VvMybA1^{RUO}* allele, which leaves the last part of the retrotransposon (solo-3'LTR region) at *VvMybA1* promotor as described by Migliaro et al. [13]. However, a similar mechanism was only observed in the Mourisco variety. Considering the results obtained for the Mourisco variety, both cultivars (Mourisco Roxo and Mourisco Tinto) should have derived from an unpigmented wild-type ancestor, homozygous for the *Gret1* insertion (*VvMybA1^{ITA}*). At least the L2 layer must have been affected by the subsequent partial excision of the *Gret1*, although it is not possible to verify if the mutation resulting in the partial excision of *Gret1* affected the L1 layer as well, since this layer-specific approach was not applied on tissues containing only L1 cells. Thus, the skin color differences observed among these two somatic variants are probably due to specific interferences on the anthocyanin biosynthetic pathway regulated by different regulatory transcription factors, since they both showed the same allelic composition for *VvMybA1* and *VvMybA2* genes.

The same wild-type genotype could be affected by different mutations resulting in color gain, as previously reported for two variants of Italia. The recovery of *VvMybA1* expression on the less-pigmented Ruby Okuyama cultivar was caused by intra-LTR recombination within *Gret1* retrotransposon, while for the more intense pigmented Benitaka cultivar, color gain was caused by homologous recombination between *VvMybA1* and *VvMybA3*, called *VvMybA1^{BEN}* allele, which was identified as a novel functional allele that restored *VvMybA1* transcripts [16].

Although the *Gret1* insertion at the promoter region of *VvMybA1* has been considered as the main factor determining grape skin color variation and responsible for the recent and extremely rapid diffusion of the white phenotype regarding the grapevine domestication history, the *VvMybA2* polymorphisms also seem to play a relevant role in the *VvMybA* diversification process [23], as observed for some less-pigmented variants of group B, namely Folgasão Roxo, Malvasia Cândida Roxo and Malvasia Fina Roxo.

As previously mentioned, except for the Mourisco variety, in the remaining varieties of group B, the analyzed molecular markers were homozygous and monomorphic along an extensive genomic region of chromosome 2. These results are in accordance with the homozygosity found by Migliaro et al. [13]. Thus, all these findings agree with the history of grape evolution and domestication. Although domestication and breeding are usually associated with a decrease of grape diversity, its influence, on a genome-wide scale, seems to be weak due to the different changes in morphology observed since grape domestication, including a wide range of berry colors. Myles et al. [23] identified a 5-Mb region on chromosome 2 encompassing the MYB transcription factor genes having observed a positive selection for white grapes around this locus, which is in agreement with the intense breeding for lighter berry color and the rapid dissemination of the MYB mutations responsible for reduced pigmentation previously referred to by Fournier-Level et al. [17].

In summary, this layer-specific approach was informative regarding the discrimination of skin color somatic variants with a pigmented ancestor, particularly the VVNTM1 and VVNTM5 SSR markers. Moreover, VVNTM3 and VVNTM4 SSR markers were the less informative markers since they were shown to be homozygous for almost all the somatic variants for berry skin color analyzed. Additionally, for somatic variants derived from an unpigmented ancestor this approach was shown to be not very informative regarding the discrimination of the variants, due to the extensive homozygosity observed either between variants or between layers of each variant.

4.3. Evolutionary mechanisms behind the origin of skin color somatic variants

Based on layer localization and the differences in the size of the deletion on the distal arm of chromosome 2, it is suggested that the less-pigmented and unpigmented somatic variants analyzed in this study can be ascribed from a pigmented ancestor (group A) in accordance with two theoretical models: 1) the Sequential model, which is the first evolutionary model proposed to explain the somatic color mutants development within a variety, also reported as ‘Cabernet Sauvignon-model’ by Walker et al. [11]; according to this model, the black-skinned wild-type ancestor gives rise to a less-pigmented variant which in turn gives rise to the unpigmented variant (Fig. 1A); and 2) the Parallel model, first proposed for Pinot family by Vezzulli et al. [19], where the black-skinned berry ancestor gives rise to the less-pigmented (Pinot Gris) and the unpigmented (Pinot Blanc) variants independently (Fig. 1B). Based on the results obtained, three varieties seem to follow the Sequential model, namely; Aspiran, Grolleau and Terret. The remaining varieties (Aramon, Pinot and Pique-poul) seem to be in agreement with the Parallel model.

Regarding the experimental data obtained for group B, where the wild-type ancestor is a white-skinned cultivar giving rise to a less-pigmented variant, a Revertant model (Fig. 1C) is proposed. As mentioned above, the main mechanism described as being responsible for color gain involves the partial excision of *Gret1* retrotransposon from the *VvMybA1* promoter. However, this mechanism can only be applied to the Mourisco variety, in both pigmented variants. Our data suggests a novel mechanism for the genetic make-up of less-pigmented variants evolving from an unpigmented ancestor. For the cases of the less-pigmented variants (Folgasão Roxo, Malvasia Cândida Roxo and Malvasia Fina Roxo) color gain seems to result from the recovery of the functional G allele on *VvMybA2R44* position. This event might be playing a fundamental role, probably by restoring *VvMybA2* transcripts (Fig. 1C).

5. Conclusions

These findings represent a significant breakthrough regarding the mechanisms behind the formation of somatic variants for berry skin color, namely unpigmented or pigmented-related grape cultivars with different color shades, since such molecular information has not been previously available for somatic variants derived from the pigmented ancestors Aramon, Aspiran, Grolleau, Pique-poul and Terret, nor for the unpigmented ancestors, Folgasão, Malvasia Cândida, Malvasia Fina and Mourisco.

These data support the conclusion that, besides *VvMybA1* and *VvMybA2* playing an important role regarding the phenotypic variation observed among the berry skin color somatic variants due to black-to-grey and/or white reversions, the molecular mechanism leading to skin color variation results from different deletion patterns encompassing the berry color locus. Moreover, the experimental data regarding the white-to-red reversion suggest that *VvMybA2* gene might play a fundamental role for color gain through the recovery of functional G allele on *VvMybA2R44* position.

This approach was also shown to be particularly informative regarding the discrimination of somatic variants for berry skin color with a pigmented ancestor, therefore it could be used to solve problems of cultivar identification and newly appearing somatic variants for berry skin color.

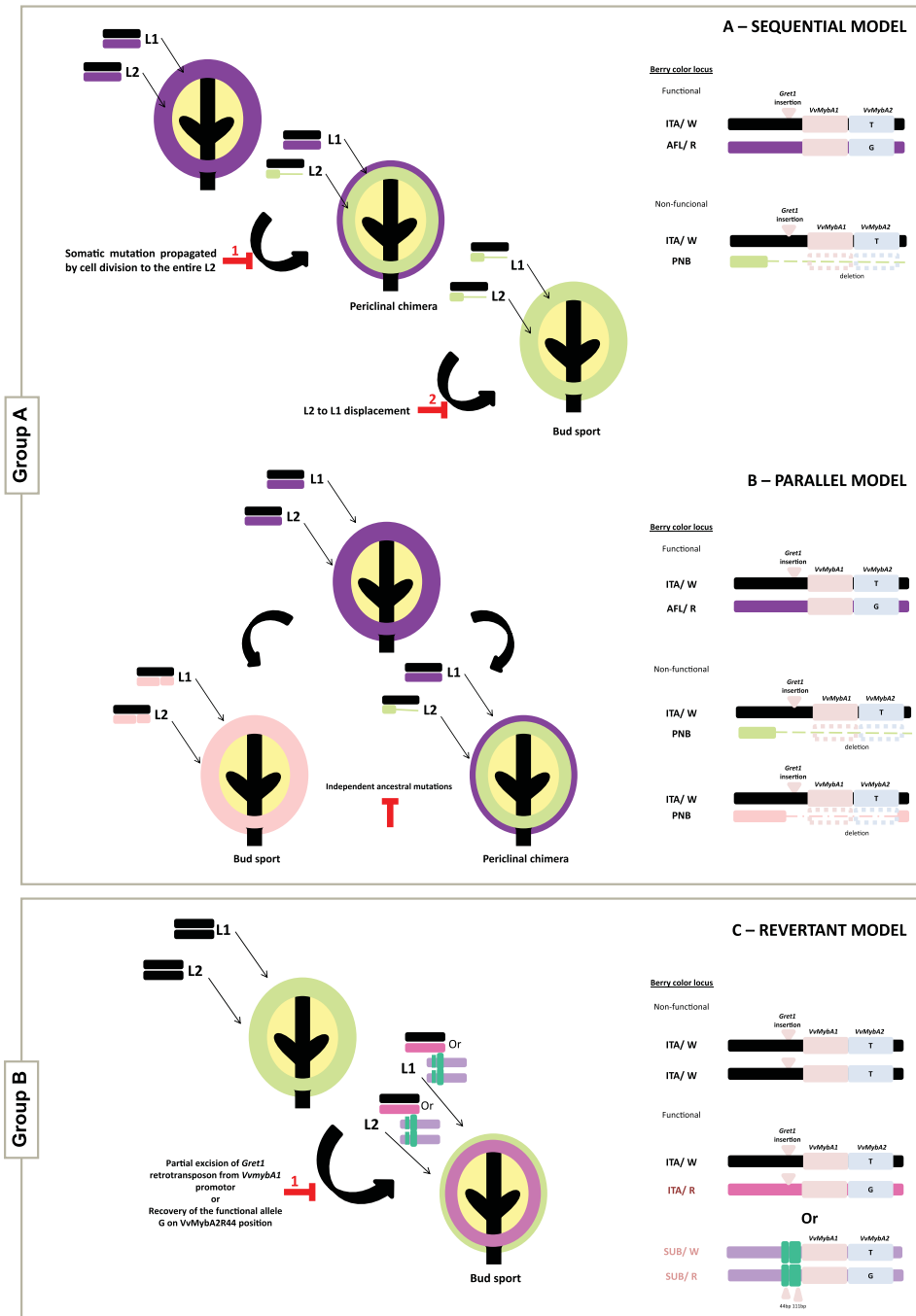


Fig. 1. Schematic representation of the evolutionary models describing the structural dynamics at the berry color locus in the L1 and L2 cell layers. Variants of the Group A can originate either through the Sequential model (A) or the Parallel model (B). The origin of variants from the Group B can be explained by the Revertant model (C), from one out of two different mechanisms, the partial excision of the Gret1 retrotransposon from the *VvMybA1* promoter or the recovery of the functional allele of *VvMybA2*. On the right side are represented the structures of the different alleles and genotypes found in the somatic variants for berry skin color studied.

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Conflict of Interest

The authors have no conflict of interest to report.

Supplementary material

The supplementary material is available in the electronic version of this article: <http://dx.doi.org/10.3233/JBR-170289>.

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