RESEARCH PAPER

The grapevine VvibZIPC22 transcription factor is involved in the regulation of flavonoid biosynthesis

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Abstract

In grapevine, flavonoids constitute one of the most abundant subgroups of secondary metabolites, influencing the quality, health value, and typicity of wines. Their synthesis in many plant species is mainly regulated at the transcriptional level by modulation of flavonoid pathway genes either by single regulators or by complexes of different regulators. In particular, bZIP and MYB factors interact synergistically in the recognition of light response units present in the promoter of some genes of the pathway, thus mediating light-dependent flavonoid biosynthesis. We recently identified VvibZIPC22, a member of clade C of the grapevine bZIP family, in a quantitative trait locus (QTL) specifically associated with kaemperol content in mature berries. Here, to validate the involvement of this candidate gene in the fine regulation of flavonol biosynthesis, we characterized its function by in vitro and in vivo experiments. A role for this gene in the control of flavonol biosynthesis was indeed confirmed by its highest expression at flowering and during UV light-mediated induction, paralleled by accumulation of the flavonol synthase 1 transcript and flavonol compounds. The overexpression of VvibZIPC22 in tobacco caused a significant increase in several flavonoids in the flower, via induction of general and specific genes of the pathway. In agreement with this evidence, VvibZIPC22 was able to activate the promoters of specific genes of the flavonoid pathway, alone or together with other factors, as revealed by transient reporter assays. These findings, supported by in silico indications, allowed us to propose VvibZIPC22 as a new regulator of flavonoid biosynthesis in grapevine.

Key words: bZIP factors, co-expression analysis, flavonoids, gene expression, grapevine, luciferase assay, metabolic profiling, tobacco, transient expression.

Introduction

In grapevine (Vitis vinifera L.), as well as in many plant species, flavonoids represent one of the most abundant subgroups of phenolic compounds, whose synthesis derives from the amino acid phenylalanine through the phenylpropanoid pathway. They are composed of three main classes, anthocyanins, proanthocyanidins (also known as condensed tannins), and flavonols. They accumulate preferentially in flowers and in the peripheral layers of the berry pericarp and of the seed...
coat of grapes, and fulfil several important functions. They (i) mediate the response to biotic and abiotic stresses (temperature, UV light, nutrition, water deficit); and (ii) provide pigmentation to flowers and fruits, thereby also influencing the quality and typicity of wines. Moreover, they are associated with health-promoting effects ascribed to grape-rich diets including wine (Flamini et al., 2013; Teixeira et al., 2013).

Due to this wide range of functions, the synthesis of these compounds is finely regulated, mainly at the transcriptional level. The expression of the flavonoid biosynthetic genes is usually regulated by forming dimers (et al., 2013) and usually regulate transcription by forming specific flavonoids in the different organs and phases. Although flavonoid and especially anthocyanin biosynthesis has been extensively studied, also in grapevine, its regulation is not yet completely understood (Hichri et al., 2011a). The whole system is under the tight control of several transcription factors, which can mutually interact giving rise to regulatory specificity (Grotewold, 2006). In all species analysed to date, these factors mainly belong to the family of the R2R3-MYB domain-containing proteins (Hichri et al., 2011a; Li, 2014). However, there is evidence which indicates bZIP proteins as involved together with MYB and other factors in the transcriptional control of flavonoid biosynthesis depending on light (Hartmann et al., 2005; Czemmel et al., 2009; Stracke et al., 2010).

bZIP factors belong to a large family already characterized in many plant species, including grapevine (Liu et al., 2014). These factors are involved in the regulation of plant development (Strathmann et al., 2001), response to drought (Yoshida et al., 2010), high salinity (Seong et al., 2008), light (Hartmann et al., 2005; Stracke et al., 2010), and pathogen infection (Throw et al., 2005), as well as amino acid (Hanson et al., 2008; Dietrich et al., 2011) and phenylpropanoid biosynthesis (Heinekamp et al., 2002; Hartmann et al., 2005). So far, only a few grapevine bZIP genes have been characterized, mainly as being involved in the abscisic acid (ABA)-dependent response to abiotic stresses (Tak and Mhatre, 2013) and grape berry ripening (Nicolas et al., 2014).

Plant bZIPs bind to ACGT-containing elements (Izawa et al., 1993) and usually regulate transcription by forming dimers (Strathmann et al., 2001; Ehler et al., 2006) and interacting with non-bZIP proteins (Schutze et al., 2008), to gain flexibility in their regulatory capacity. In particular, bZIP heterodimers of the C/S1 groups of Arabidopsis have been implicated in stress response and development (Weltmeier et al., 2006) and in sugar-regulated control of amino acid metabolism during the plant response to starvation (Dietrich et al., 2011).

Flavonoid biosynthesis in grapevine is also regulated by environmental stimuli (Teixeira et al., 2013) and is particularly affected by light (Fujita et al., 2006; Matus et al., 2009; Koyama et al., 2012). Koyama et al. (2012) described the influence of light quality on the regulation of flavonoids in young berry skins: visible light primarily induces proanthocyanidin (PA) biosynthesis, whereas UV light specifically induces flavonol biosynthesis. In Arabidopsis, the light regulation of flavonol biosynthesis is directed by MYB and bZIP factors which co-operatively bind to the light regulatory units (LRUs) present in the CHS and FLS promoter sequence (Hartmann et al., 2005). A well-studied example is the Arabidopsis HY5 gene, encoding a bZIP factor which regulates numerous genes, such as AtCHS, AtFLS, and AtMYB12, during photomorphogenesis (Lee et al., 2007; Stracke et al., 2010). The presence of LRUs in the promoters of VviFLSI and VviMYB1I and their light responsiveness led Czemmel et al. (2009) to propose them as targets of a grapevine HY5 homologue.

Recently, we have identified two grapevine bZIP genes, namely the predicted VvbZIP14 and VvbZIP22 (Liu et al., 2014), within two quantitative trait locus (QTL) regions associated with the fine tuning of flavonol content in mature grapes, on chromosomes 5 and 7, respectively. Notably, these bZIPs were also expressed at different levels in grapes of ‘Syrah’×‘Pinot Noir’ progeny showing different contents of flavonols in the mature skin (Malacarne et al., 2015). The QTL on chromosome 7, containing VvbZIP22, was specifically related to kaempferol content, while the other, containing VvbZIP14, was also associated with anthocyanin content (Costantini et al., 2015). Starting from this evidence, in this study, we decided to characterization further VvbZIP22, here renamed VvbZIPC22 according to the recently proposed grapevine genome nomenclature system (Grimplet et al., 2014).

Materials and methods

Grapevine material and sampling

Inflorescences and clusters from a representative sample of V. vinifera cv. Pinot noir (clone ENTAV115) were collected at six developmental stages during the 2011 season at the Gionarion experimental field of FEM (Edmund Mach Foundation, San Michele a/Adige, Italy, 46°18’N, 11°13’E). To be more representative, different parts of the field were sampled, skins were separated from flesh and seeds directly in the field when feasible, precisely from véraison to maturity stages, and immediately frozen in liquid nitrogen. In the 2012 season, vegetative organs (young leaf, mature leaf, bud, root, internode, and tendrill), germinated seeds, green (E-L 20) and mature inflorescences (E-L 23) (divided into calyptra, stamen, and pistillum), and berries at different stages (E-L 29, 34, and 36) were sampled from the same Pinot Noir plants considered in 2011 and immediately frozen in liquid nitrogen. In the case of berries at stages E-L 34 and 36, seeds were removed during grinding. For each stage and organ of the panel, three different plants (biological replicates) were considered.

For light induction experiments, dormant hardwood cuttings of grapevine V. vinifera L. cv. Chardonnay were collected from a vineyard in Neustadt/W, Germany (49°22’9”S, 8°10’28”E). Cuttings were grown in sterile Perlite (Knauf, Sittingbourne, UK) under the following conditions: temperature, 25 °C; white light, ∼100 μmol m−2 s−1; 9 h light cycle. In this hydroponic system, the apical buds burst after 10 d and roots developed after 4 weeks. After 19 d, the rooted plants were subjected to light treatment [4% UV-B and 30% UV-A light (18 W, 6500 K)] under the following conditions: temperature, 23 °C; 9 h light cycle. In the case of treated samples, the third and the fourth leaf from the shoot tip of five plants were collected at 0, 10, 24, 48, and 72 h after the onset of light exposure. Control samples were collected at the same time points, from five other plants grown under the same conditions, but without UV light. At each time point, leaves were pooled, immediately frozen in liquid nitrogen, and stored at −80 °C till use.
All the collected samples were finely ground by liquid nitrogen using frozen metal grinding jars mounted on a Mixer Mill MM 400 (Retsch, Haan, Germany). The powder was maintained at –80 °C for 2 months to be used for both transcriptional and biochemical analyses.

Cloning of VvibZIPC22 and tobacco transformation

The cDNA from Pinot Noir berry skins at the véraison stage diluted 10-fold was used for PCR amplification with forward primer (bZIP-XhoI 5'-CTCGAGGAAATGTGCGGTGCAAC-3', XhoI recognition site underlined) and reverse primer (bZIP-KpnI 5'-GGTACCATGACATGAAACAGTGTGGT-3', KpnI recognition site underlined). The VvibZIPC22 sequence considered in this study was derived from the 12Xv1 assembly of the Pinot Noir genome (VIT_07s0005g01450, from http://genomes.cribi.unipd.it/DATA/V1, last accessed 11 December 2015) and confirmed by EST sequences (EE065899.1, EC946004.1, and EC945791.1). The complete coding DNA sequence (CDS) of VvibZIPC22 was amplified in a 12.5 µl PCR mix containing primers (200 pM each), 1 U of Phusion DNA polymerase (Thermo Fisher Scientific, Waltham, MA, USA), dNTPs (200 µM), PCR buffer (1×), and cDNA (diluted 10-fold). The PCR conditions adopted were: 95 °C for 2 min, followed by 35 cycles, each one consisting of 95 °C for 30 s, 58 °C for 30 s, and 72 °C for 45 s, followed by a final extension (7 min at 72 °C). A PCR product of 438 bp was obtained and subcloned into a pGEM-T Easy Vector (Promega, Mannheim, Germany) for sequencing. The VvibZIPC22 CDS (KK073969) was subsequently transferred into an XhoI/KpnI-digested pART17 primary cloning vector (Gleave, 1992) under the control of the Cauliflower mosaic virus (CaMV) 35S promoter to give pART7bZIPC22. In both cases, a chemical transformation was performed using One Shot® TOP10 competent Escherichia coli cells (Thermo Fisher Scientific).

pART7bZIPC22 was used both for the transient expression assays and as an intermediate to obtain pART27bZIPC22 for tobacco transformation. CaMV35S:VvibZIPC22 was therefore transferred into a NotI-digested pART27 binary vector to obtain pART27bZIPC22. Single colonies of Agrobacterium tumefaciens (strain C58C1) transformed with pART27bZIPC22 by electroporation were cultured in 50 ml of selective YEB medium and grown overnight (28 °C at 160 rpm). The cells were harvested by centrifugation at 4000 rpm for 30 min without break and resuspended in 50 ml of infiltration buffer (10 mM MES, pH 5.9 and 150 µM acetosyringone). The suspension was incubated at 28 °C and 50 rpm for 2 h. Thereafter the concentration was measured by spectroscopy (OD400 nm) and adjusted to a final concentration of OD0.01 nm of 0.3 with H2O. Nicotiana tabacum cv. Samsung leaf sections (1 cm2) were further incubated with the bacterial suspension right-side up for 30 min at room temperature. Leaf explants were transferred to a sterile wet filter paper and incubated in the dark for 2 d at room temperature. The explants were then transferred right-side down for 30 min to solid Murashige and Skoog (MS) medium plus timentin (400 µg l−1), supplemented with B5 vitamins, 3% (w/v) sucrose, 1 mg l−1 6-benzyladrene (BA), 0.1 mg l−1 α-naphthaleneacetic acid (NAA), 0.8% tissue culture agar, pH 5.6, and 100 mg l−1 kanamycin. Explants were then incubated in the dark for 2 weeks at 25 °C, and then transferred to 25 °C, 16 h light cycle. Finally, kanamycin-resistant plantlets were transferred to soil mix, acclimatized, and grown in the greenhouse. The presence of the transgene in transformed lines was established by PCR with the primers 35SCaMvPAPRT7 (5'-CAATCCCATCATCTCCGGCA-3') and bZIP-KpnI. Stamens and petal limbs at two developmental stages (open flower without pollen and mature flower) were collected from a representative sample of each transgenic line and wild-type plant and stored at –80 °C until biochemical and gene expression analyses.

Grapevine transient expression experiments

A transient expression system using Chardonnay liquid cell cultures was established as described previously (Bogs et al., 2007). Cells were bombarded with 1.6 µm gold particles using a PDS-1000/ He Biologic Particle Delivery System from Bio-Rad (München, Germany) with 4481 kPa helium pressure, a vacuum of 86 kPa, and a distance of 9.5 cm. The luciferase assay was performed as set up by Czemmel et al. (2009). Briefly, gold particles were coated with 500 ng of each respective plasmid, giving a total amount of 2 µg of plasmid per transformation for reporter/effector bombardments. Additionally, each bombardment contained a positive control of 100 ng of the Renilla luciferase plasmid pRLuc (Horstmann et al., 2004). After incubation in the dark at 27 °C for 48 h, the harvested cells were assayed for luciferase activities using the dual-luciferase reporter assay system (Promega) measured with a Lumintron LB 9507 Luminometer (Berthold Technologies, Bad Wildbad, Germany). The relative luciferase activity was calculated as the ratio between the firefly and the Renilla (control) luciferase activities after subtraction of the cell background (ground cells but not bombarded). All transfection experiments were carried out in triplicate, and each reporter experiment was repeated at least three times.

The design of the effector constructs of VviMYBP1a (AM259485) and VviMYBP2a (BAD18978), VviMYBF1 (AC81697), and VviMYC1 (ACC68685) is described in Bogs et al. (2007), Czemmel et al. (2009), and Hichri et al. (2010), respectively. The effector construct of VvibZIPC22 (pART7bZIPC22) was obtained as previously described.

The cloning of the promoter fragments of VviCHS1 (AB015872), VviCHS2 (AB066275), and VviCHS3 (AB066274) promoters by specific primers, whose sequences were as follows, with restriction sites for SacI and XhoI underlined: CHS1pF (5'-TATGAGTCATCTAACAAGCGTGTTG-3'), CHS1pR (5'-TATATCGAGGTGCTACCTGTTCTC-3'), CHS2pF (5'-TATGAGTCATCTAAGGAAAGAGG-3'), CHS2pR (5'-TATGAGTCTAGATTGACATGATAGG-3'), CHS3pF (5'-TATGAGTCATCTAAGAGGAAAGG-3'), and CHS3pR (5'-TATGAGTCTAGATTGACATGATAGG-3').

Biochemical analyses of flavonoids in grapevine and tobacco samples

Extraction of anthocyanins, flavonol aglycons, and flavan-3-ol monomers and their analysis by HPLC-DAD (diode array detection) from grapevine samples was performed as described in Mattivi et al. (2006, 2009) using a Waters 2690 HPLC system equipped with Waters 996 DAD and Empower software (Waters Corporation, Milford, MA, USA). Some modifications were applied to the extraction of flavonols from light-treated and control leaf samples. Briefly, three separate 200 mg aliquots (technical replicates) of frozen powdered extract were extracted with 1.5 ml of methanol by rotation (30 rpm for 20 min). After centrifugation (13 000 rpm for 5 min), 1.5 ml of the supernatant was transferred to a 50 ml flask. The same procedure was adopted in a second extraction with 1.5 ml of methanol. The flask with a total volume of 10 ml (3 ml of methanolic extract+2 ml methanol+5 ml of trifluoroacetic acid 2 M in water) was used for the following acid hydrolysis as in Mattivi et al. (2006).

Extraction of anthocyanins and flavonols from tobacco stamens was performed starting from three separate 150 mg aliquots (technical replicates) of frozen powder. After the addition of 1.5 ml of methanol, each sample was shaken at room temperature for 20 min. After centrifugation (14 000 rpm for 5 min), each sample was then filtered with 0.22 µm filters into LC vials and subjected to HPLC-DAD analysis.

Each anthocyanin, flavonol, and flavan-3-ol compound was identified by comparison of its retention time and UV spectra at 520, 370, and 280 nm. The concentration (µg g FW−1) of each compound was determined using the external standard method, specific for each compound. Values under the limit of detection (LOD) and of
quantification (LOQ) were given a value equal to zero in the final quantification.

Proanthocyanidins (PAs) in tobacco petals were determined using dimethylaminocinnamaldehyde (DMACA) reagent described by *Nagel and Glories* (1991). Briefly, three separate 150 mg aliquots (technical replicates) of frozen powder were extracted with 2 ml of AA-buffer (70% acetic acid, 0.1% ascorbate) by sonication in ice-water (20 kHz, amplitude 50% for 30 s) with a Sonifier SFX250 (Branson Ultrasونics, Danbury, CT, USA). After centrifugation (13 000 rpm, 4 °C), 1 ml of extract was mixed with 0.75 ml of diethyl ether to remove chlorophyll, and stored at −20 °C for 2 h. After incubation, 100 µl of the lower PA phase was mixed with 900 µl of a DMACA solution (0.1% DMACA, 1% 3N HCl) and incubated in the dark (room temperature for 60 min). Finally, the PA concentration (µg g FW⁻¹) was determined reading OD₅₄₀nm against a blank DMACA solution and using catechin (Sigma Aldrich, St. Louis, MO, USA) as standard.

**Quantitative gene expression analysis in grapevine and tobacco samples**

Total RNA was isolated from grapevine and tobacco samples with the Spectrum Plant Total RNA Kit (Sigma Aldrich) according to the manufacturer's instructions. RNA was quantified with an ND-8000 nanodrop spectrophotometer and checked for integrity with a 2001-Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). The reverse transcription-PCR and real-time RT-PCR (qRT-PCR) analyses, cDNAs were synthesized using the Superscript Vilo™ cDNA Synthesis Kit from 1.5 µg of DNase I-treated RNA (Thermo Fisher Scientific), following the manufacturer's instructions. qRT-PCR analyses were carried out using the Platinum SYBR Green qPCR SuperMix-UDG in a ViiA™ 7 thermocycler (Thermo Fisher Scientific). The 384-well plates were set up according to the sample maximization strategy proposed in *Hellemans et al.* (2007), and each reaction was run in triplicate. Reaction conditions and the protocol of analysis were the same as adopted in *Malacarne et al.* (2015). Grapevine and tobacco housekeeping genes were tested for their stability in each experiment using the GeNorm software (*Vandesompele et al.*, 2002). Normalized relative quantities (NRQs) were then calculated by dividing RQs by a normalization factor, based on the expression of the most stable reference genes (details are given in the figure legends) (*Reid et al.*, 2006). Primer sequences used for the analysis are listed in Supplementary Table S1 at *JXB* online, as designed in this work or taken from the literature.

**Bioinformatics**

For the research of plant *cis*-regulatory elements, the first 1000bp from the putative transcriptional start of the promoter sequences were screened manually and by the use of PLACE (http://www.dna.affrc.go.jp/PLACE/; *Higo et al.*, 1999).

For phylogenetic studies, the complete CDS of *VvibZIP* genes belonging to clade B and C (*Liu et al.*, 2014) and of their putative orthologues in different plant species were retrieved from public databases (http://genomes.cribi.unipd.it/DATA/V1 and http://www.ncbi.nlm.nih.gov/; last accessed 11 December 2015). Codon-based alignments of *VZIPS* CDS sequences were made using the Mace program (*Ranwez et al.*, 2011). Automatically produced alignments were loaded into the Seaview alignment editor and manually edited in amino acid mode to discard misaligned regions. The final selection of alignment columns was saved to produce (i) a 1335 positions long curated alignment of coding clade B *VvibZIP* sequences and their putative orthologues (*Supplementary Dataset S1*) and (ii) a 297 positions long curated alignment of coding clade C *VvibZIP* sequences and their putative orthologues (*Supplementary Dataset S2*). Bayesian analyses were performed employing the MPI version of Phylobayes (version 1.4) and a general specification of the CAT+GTR+G4 model. Two chains were run on each data set under each model and until 5000 cycles were sampled. Trees were built for each data set based on the two chains, discarding the first 2500 cycles as burn in, which was sufficient for ML (maximum likelihood) parameter values to maximize in all analyses. Parameter values were then sampled every cycle thereafter till the 5000th cycle.

Co-expression analyses were performed by means of the Vitis Expression Studies Platform Using COLOMBOS Compendia Instances (VESPUCCI) (*M. Moreto et al.*, 2016b), a grapevine gene expression compendium (http://vespucci.colombos.fmach.it/) based on COLOMBOS v3.0 technology (*Moreto et al.*, 2016a). The analyses by COLOMBOS were essentially based on contrast relevance and gene similarity scores as thoroughly described in Text S1 from *Engel et al.* (2011) and the updates made with the v3.0 release.

**Results**

*VvibZIP22* belongs to one group of clade C in the phylogenetic tree of *VvibZIP* factors

The recent phylogenetic analysis of the grapevine *bZIP* gene family assigned *VvibZIP22* to clade C, together with another nine *VvibZIP* genes. In particular, *VvibZIP22*, together with *VvibZIP14* and *VvibZIP37*, forms a homologous triplet with collinearity along chromosomes 5, 7, and 14 (*Liu et al.*, 2014). According to the recently suggested grapevine genome nomenclature system (*Grimplet et al.*, 2014), we renamed the VvbZIP factors indicating the clade to which they belong, with the exception of the unresolved members.

To better define the structure of clade C and to identify sequences closely related to *VvibZIP22* from other species, the CDS sequences of VvbZIP factors belonging to this clade were aligned and compared with those of bZIP factors from other important plants and crops with significant identity (>50%) to *VvibZIP22* (*Supplementary Dataset S2*). Phylogenetic analysis revealed that clade C can be divided into three main groups, designated C1, C2, and C3 (Fig. 1), similarly to clade S in Arabidopsis (*Ehler et al.*, 2006). In particular, *VvibZIP22* is part of group C1 together with *VvibZIPC14* and *VvibZIP37* and with previously characterized bZIP genes from flowering species such as *N. tabacum* (*NiBZI-3* and *NiBZI-4*; *Strathmann et al.*, 2001), *Petroselinum crispum* (*PcCPRF6*; *Rugner et al.*, 2001), Arabidopsis thaliana (*AtbZIP53*; *Dietrich et al.*, 2011), and *Solanium lycopersicum* (*SlbZIP04*; named *LebZIP2* in *Seong et al.* 2008). Moreover, other orthologues can be considered: *SiOCSI* from *Sesamum indicum*, *TcOCSI* from *Theobroma cacao*, *PrOCSI* from *Populus trichocarpa*, *PxbOCSI* from *Pyrus × bretschneideri*, *MdhZIP3* from *Malus domestica*, *GbZIP33* from *Glycine max*, *CsOCSI* and *CmOCSI* from *Cucumis sativus* and *Cucumis melo*, and *CrOCSI* from *Citrus clementina*. In our analysis, *VvbZIP07*, *VvbZIP13*, and *VvbZIP47*, which in the tree of *Liu et al.* (2014) belong to clade C, were unresolved with respect to group C1 together with *PcCPRF7* from *P. crispum* and *NiBZI-2* from *N. tabacum*. *VvbZIPC02* and *VvbZIPC39* fall into group C2 together with members of group S2 in *A. thaliana*.
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Fig. 1. Phylogenetic tree of clade C VvibZIP factors and of their putative orthologues. C1, C2, and C3 correspond to the identified groups within clade C, similarly to those identified in clade S in Arabidopsis (Ehler et al., 2006). Bootstrap values are shown next to the nodes. Gene names with orthologues. C1, C2, and C3 correspond to the identified groups within clade C, similarly to those identified in clade S in Arabidopsis (Ehler et al., 2006). Bootstrap values are shown next to the nodes. Gene names with

VvibZIP22 expression is flower specific and correlates with kaempferol and quercetin content during berry development

To determine if a significant correlation exists between VvibZIP22 expression and the content of different flavonoids, we looked at their profiles at six time points of Pinot Noir berry development (Fig. 2A). VvibZIP22 expression as well as kaempferol and quercetin aglycons showed similar profiles: they peaked at flowering, reached a minimum between pre-véraison and véraison, and increased slightly towards maturity. It is noteworthy that VvibZIP22 expression was highest at flowering, when a peak of expression was also reported for VviMYBFI (Czemmel et al., 2009) and VviFLSI (Downey et al., 2003) in developing Shiraz berries. The same analysis highlighted that delphinidin-like flavonoids had an opposite trend, with maximum accumulation in mature berries (Fig. 2A). Early and late peaks in the biosynthesis of different flavonols have already been described in the V. vinifera cv. Pinot Noir (Sternard Lemut et al., 2013). On the other hand, we did not find a significant correlation between VvibZIP22 expression and the synthesis of either total anthocyanin 3-monoglucosides or total flavan-3-ol monomers (Supplementary Fig. S1).

The expression analysis extended to different grape organs showed that VvibZIP22 is mostly expressed in the flower, as demonstrated by its level in calyptras and stamens, 13 times higher than in berries at stage E-L 36 (16° Brix) and almost three times the expression observed in the other organs (Fig. 2B).

Induction of VvibZIP22 expression by light is related to flavonol accumulation

Light-induced flavonol biosynthesis (Downey et al., 2004) is the result of an increase in VviMYBFI and VviFLSI expression mediated by light regulatory elements present in both promoters and recognized by MYB and bZIP factors (Czemmel et al., 2009). For this reason, we monitored VvibZIP22 as well as VviMYBFI and VviFLSI gene expression over a 3 d time period together with flavonol accumulation in Chardonnay plants exposed to UV light. Within 10 h of light exposure, the transcript level of VvibZIP22, VviFLSI, and VviMYBFI increased ~2.5-, 7.5-, and 22-fold in treated compared with control leaves. VvibZIP22 and VviFLSI increased by almost half at 24 h and 72 h, and VviMYBFI was barely detectable at these times (Fig. 3A). The peak of induction of the three genes at 10 h of treatment was followed by the accumulation of flavonols starting from 24 h and reaching a maximum at 72 h of light induction (Fig. 3B).

The presence of light response cis-acting elements in the VvibZIP22 promoter sequence in both Pinot Noir and Chardonnay was a further indication of its light responsiveness. Hartmann et al. (2005) showed that LRUs formed by MYB recognition elements (MREs) and ACGT-containing elements (ACES) mediate light responsiveness of the ArChIS promoter. Designated elements, such as aCS (ACGT-containing
sequence similar to ACE) and MRS (MYB recognition sequence similar to MRE), were also identified in the promoter region of VviMYBF1 and VviFLS1 (Czemmel et al., 2009). In this study, we investigated 1000 bp of the sequence of Pinot Noir and Chardonnay bZIPC22 upstream of the putative transcriptional start site: they were identical, with the exception of a polymorphism at position –303. This was not unexpected since Pinot Noir is the male parent of Chardonnay.

Moreover, we found that both promoter sequences contain an ACS element at a position (–137 bp) fairly well conserved with the ACE site in AtCHS and AtFLS, and with the ACS site in VviFLS1 and VviMYBF1. In addition, an MRS site (–583 bp), possibly recognized by MYB factors, and an IBOXCORE element (–193 bp), involved in binding of MYB factors in light-regulated genes in tomato (Rose et al., 1999), were localized upstream of the ACS site (Fig. 3B), as found in the VviFLS1 promoter (Czemmel et al., 2009).

VvibZIPC22 activates promoters of general and specific flavonoid pathway genes

In order to identify which flavonoid pathway genes could be the potential targets of VvibZIPC22, transient expression experiments were carried out by bombardment of Chardonnay liquid cell cultures and dual-luciferase assays. Five different promoters were tested: the VviCHS and VviCHI promoters, as genes controlling the general flavonoid pathway, and the VviFLS1, VviUFGT, and VviANR promoters, as genes specifically involved in the synthesis of flavonols, anthocyanins, and PAs, respectively. VvibZIPC22 was tested alone and in combination with VviMYBF1, VviMYC1–VviMYBP1, and VviMYBA2 or VviMYC1–VviMYBA2 factors to test the transactivation potential on the VviCHI and VviFLS1 promoters in the first case, on the VviCHI and VviANR promoters in the second case, and on the VviUFGT promoter in the third case (Fig. 4).

VvibZIPC22 was able to activate the promoter of VviCHS3, the isoform expressed in red-coloured berry skin during ripening (Goto-Yamamoto et al., 2002), approximately three times more than in the control without VvibZIPC22, suggesting a possible role in the general regulation of the flavonoid pathway (Fig. 4A). VvibZIPC22 also induced the promoter of VviFLS1 to an extent similar to the positive control VviMYBF1 (~3.5-fold) (Fig. 4C). Interestingly, VvibZIPC22 increased the transactivation potential of VviMYBF1 2-fold by acting synergistically on the promoters of both VviCHI (Fig. 4B) and VviFLS1 (Fig. 4C). As published before (Hichri et al., 2010), VviMYBP1 in combination with the basic helix–loop–helix (bHLH) factor VviMYC1 induced the VviANR promoter (~20-fold), but we observed that its effect
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is amplified 2-fold by the action of VvibZIPC22, suggesting this bZIP might also play a role in the regulation of PA biosynthesis (Fig. 4D). Conversely, VvibZIPC22 did not display any significant positive effect on the VviUFGT promoter either alone or in combination with the complex VviMYC1–VviMYBA2 and with VviMYBA2 (Fig. 4E).

We also noted that the promoters of VviCHS3, VviCHI, VviFLS1, and VviANR have an ACGT-containing element in a rather conserved position, between 100 bp and 150 bp upstream of the transcriptional start site. In contrast, ACGT-containing elements are present only in a more distal position in the promoter of VviCHSI and completely absent in those of VviCHS2 and VviUFGT (Supplementary Table S2).

VvibZIPC22 overexpression in tobacco leads to flavonoid biosynthesis in different flower organs

The role of VvibZIPC22 in vivo was investigated by constitutive overexpression in tobacco plants. Several transgenic lines were generated, propagated, and grown in the greenhouse together with wild-type (WT) plants used for comparison.

We then carried out the following analyses on different flower organs of three WT plants and of six different lines sampled at three development stages: (i) anthocyanin and flavonol content by HPLC-DAD; (ii) PA content by reaction with DMACA; and (iii) gene expression of VvibZIPC22 and some tobacco flavonoid pathway genes by qRT-PCR.

Overexpression of VvibZIPC22 caused line-specific abnormalities, both in the vegetative part (reduced height of the plant, internodes, and size of the leaves) and in the reproductive system (corollas showed visibly increased colour and rounded lobes, and stamens appeared reddish and reduced in size) (Fig. 5A).

As expected, the six lines displayed overexpression of VvibZIPC22 and with a rather limited variability (±25%) (Fig. 5B). As far as secondary metabolites are concerned (Fig. 5C), striking differences, compared with the WT plants, were obtained looking at the flavonol and anthocyanin content in the stamens at both stages and at the PA content in the petal limbs at the second stage. Interestingly, the increase in flavonoids was paralleled by a significant increase in the expression of several structural genes (Fig. 5D). An induction
of *NtCHS*, *NtDFR*, *NtANS*, and *NtUFGT* was observed in the stamens at the first stage together with a significant increase in the content of cyanidin 3-rutinoside (up to 4-fold more than WT plants in both stages), and of *NtFLS* together with an increase of kaempferol 3-rutinoside and quercetin 3-rutinoside in the stamens at the second stage (up to 2.6- and 2-fold more than in WT plants, respectively). In some lines, these results are in agreement with the visibly red colour of the stamens (Fig. 5A). Compared with WT petal limbs, the lines also displayed a higher expression of *NtPAL* and *NtANR1* together with a higher content of PAs at the second stage.

In silico approaches provide new preliminary insights into the VvibZIPC22 regulatory network

A specific heterodimerization network of bZIP proteins from clade S (clade C in *V. vinifera*) and clade C (clade B in *V. vinifera*) was previously described in Arabidopsis (Ehlert et al., 2006), which is conserved in tobacco (Strathmann et al., 2001) and parsley (Rugner et al., 2001). We thus looked for potential interacting partners of VvibZIPC22 belonging to clade B, by a combination of phylogenetic and co-expression analyses. Our phylogenetic reconstruction (Supplementary Fig. S2) of VvibZIP genes from clade B and closely related sequences from other crop species indicated that VvibZIPB38 and VvibZIPB09 are more related to members of Rosaceae and Solanaceae, respectively, as is the case for VvibZIPC22 which shares group C1 with members from the same species.

In silico co-expression analyses against a grapevine gene expression compendium using VvibZIPC22 (VIT_07s0005g01450) as the query gene identified a set of 210 positively co-expressed genes (correlation cut-off=0.68) in 140 contrasts, conditions where VvibZIPC22 showed the highest modulation (Supplementary Table S3). Of the 210 co-expressed
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genes, 18 coded for transcription factors including the two bZIP factors, VvibZIPB38 (VIT_14s0030g02200, \( r = 0.68 \)) and VvibZIPC14 (VIT_05s0077g01140, \( r = 0.69 \)), belonging to clade B and clade C (group C1), respectively. Taken together, the two approaches suggest VvibZIPB38 as a putative heterodimerization partner of VvibZIPC22, and VvibZIPC14 and VvibZIPC22 as mutual interacting partners with VvibZIP38.

In addition, the co-expression analysis provided further preliminary insights into the VvibZIPC22 gene regulatory network. The list of co-expressed genes was enriched in genes belonging to carbohydrate and amino acid metabolism, to ABA and ethylene signalling, and flower development. In particular, it is worth noting that among the genes with the highest correlation to the VvibZIPC22 profile, eight code for starch and sucrose metabolism enzymes (one coding for a trehalose phosphatase, one for a trehalase synthase, two for trehalase 6-phosphate synthases, one for an \( \alpha \)-amylase precursor, one for a \( \beta \)-galactosidase, one for a glycoseyl transferase, and one for a \( \beta \)-fructofuranosidase, \( r \) from 0.79–0.87), one for a ferulate-5-hydroxylase, one for an amine oxidase, and one for an amine oxidase, involved in the upper part of the phenylpropanoid pathway and in flavonoid biosynthesis, respectively, and one for an amine oxidase, an oxidoreductase that can also participate in phenylalanine metabolism (Fig. 6; Supplementary Table S3).

Discussion

bZIP factors belongs to a large and diverse gene family identified in several plant genomes. Recently, the bZIP family has
also been characterized in grapevine in terms of identification of members and phylogenesis (Liu et al., 2014). Plant bZIPs are involved in the regulation of many different plant processes, including the transcriptional control of amino acid (Hanson et al., 2008; Dietrich et al., 2011) and phenylpropanoid biosynthesis (Heinekamp et al., 2002; Hartmann et al., 2005).

During development, grape flowers and berries accumulate different flavonoids at the organ- and stage-specific level as a consequence of the timely expression of genes necessary for their synthesis. Although flavonoid biosynthesis has been investigated in depth also in grapevine, its regulation is not completely elucidated (Hichri et al., 2011a).

R2R3-MYB factors are known to be the key regulators of this transcriptional network; however, there are examples supporting the involvement of bZIP proteins, acting synergistically with MYB and other factors, in the control of light-induced flavonoid biosynthesis (Hartmann et al., 2005). The most studied is ELONGATED HYPOCOTYL5 (HY5), a bZIP associated with AtCHS and PFG1/MYB12 activation and accumulation of flavonoids upon light treatment (Stracke et al., 2010).

Interestingly, we have recently identified a gene encoding a bZIP factor, namely VvbZIP22, located on chromosome 7 within a QTL region specific for kaemperol content in mature berries (Malacarne et al., 2015), and therefore a good candidate as a regulator of the flavonol pathway. Starting from this evidence, in this study we further characterized the involvement of VvbZIP22, here renamed VvibZIP22C, in the fine regulation of flavonol biosynthesis.

By phylogenetic reconstruction, we showed that VvibZIP22, together with VvbZIPC14 and VvbZIP38, forms a group of the tree and is the potential orthologue of previously characterized bZIPs from flowering species such as N. tabacum (NtBZI-3 and NtBZI-4), P. crispum (PcCPRF6), A. thaliana (AtbZIP53), and S. lycopersicum (SlbZIP04, also known as LebZIP2). These genes are associated with flower development (Strathmann et al., 2001), are induced by salt and light treatments (Seong et al., 2008; Li et al., 2015), mediate the sugar-dependent level of specific amino acids during starvation (Dietrich et al., 2011), and are involved in the regulation of phenylpropanoid pathway genes by binding to specific cis-regulatory elements (Rugner et al., 2001; Heinekamp et al., 2002).

VvibZIP22 expression was particularly high in the flower, the same organ in which kaempferol and quercetin accumulate the most, and its profile during Pinot Noir berry development resembled that of VviMYBF1 (Czemmel et al., 2009) and VviFLSI (Downey et al., 2003), encoding the key regulators of flavonol biosynthesis in grapes. This result was a further indication that VvibZIP22C is involved in the fine regulation of flavonol biosynthesis.

Flavonols, among flavonoids, are known to be modulated by UV light and function as UV protectants (Teixeira et al.,...
The light-induced flavonol biosynthesis is guided by the expression of \textit{VviMYB51} and \textit{VviFLS1}, genes containing light regulatory cis-acting elements in their promoters, recognized by MYB and bZIP factors (Czemmel et al., 2009). Based on this knowledge, we examined whether VvibZIPC22 is also involved in UV light-induced flavonol biosynthesis. Indeed, VvibZIPC22 appeared to be a light-inducible gene, being up-regulated within 10 h of UV light treatment of Chardonnay leaves, together with \textit{VviMYB51} and \textit{VviFLS1}. A similar result was also previously observed for \textit{VviMYB51} and \textit{VviFLS1} in Chardonnay cell cultures (Czemmel et al., 2009). The induction of these genes was accompanied by flavonol accumulation starting at 24 h and peaking at 72 h after treatment. The analysis of the promoter region of VvibZIPC22 indicated that its light-induced expression might be due to the presence of MRS and ACS light regulatory cis-acting elements, the same as found in the promoter sequence of \textit{VviMYB51} and \textit{VviFLS1} (Czemmel et al., 2009) and conferring light responsiveness to \textit{AtCHS} and \textit{AtFLS} (Hartmann et al., 2005). Whether VvibZIPC22 light induction is mediated by the grapevine HY5 orthologue and by MYB factors, however, remains to be elucidated.

Overexpression of VvibZIPC22 in tobacco and transient promoter assays in Chardonnay cell cultures were then adopted as complementary approaches to investigate the role VvibZIPC22 further in vivo, demonstrate a causal link between VvibZIPC22 and flavonol biosynthesis, and gain insights into its putative regulatory network. By these experiments, we provided evidence that VvibZIPC22 impacts the entire flavonoid pathway. Indeed, the tobacco overexpressing lines displayed a significantly higher content of the flavonols kaempferol and quercetin 3-rutinoside, the anthocyanin cyanidin 3-rutinoside, and the PAs in flowers, at the organ- and stage-specific level. However, due to the type of assay, we cannot completely exclude that the monitored increase in PAs corresponded to a change of the mean degree of polymerization (mDP), rather than a net increase in the absolute amount of PAs. These biochemical changes were accompanied by a significant induction of several flavonoid structural genes. Noteworthy is the ectopic expression of VvibZIPC22 that caused in most lines a visibly stronger red colouring of the stems and corolla but not of the vegetative organs, as observed in tobacco flowers over-expressing \textit{VviMYB5a} and \textit{VviMYB5b} (Deluc et al., 2006, 2008). Conversely, the ectopic expression of \textit{V1MYBA1-2} triggered \textit{de novo} production and storage of anthocyanins in all vegetative organs of the grapevine transgenic lines, attesting to a major effect of the gene on anthocyanin biosynthesis (Cutanda-Pérez et al., 2009). As proposed in the model derived from the functional characterization of \textit{VviMYB5b} (Hichri et al., 2011b), we can speculate that similarly VvibZIPC22 interacts with endogenous partners in the flower, where the anthocyanin pathway is already active (corolla), or induces \textit{de novo} anthocyanin biosynthesis by interacting with other endogenous partners of the organs where the pathway is not yet active (stamens).

The transient promoter assays confirmed that VvibZIPC22 acts on different points of the flavonoid biosynthetic pathway, working either alone or in combination with other factors. VvibZIPC22 was capable of directly activating the flavonol-specific gene \textit{VviFLS1} and the flavan-3-ol-specific gene \textit{VviANR}, as well as the genes of the general flavonoid pathway, \textit{VviCHS3} and \textit{VviCHL}, by putatively binding to an ACGT-containing element identified in a conserved position in their promoters. Interestingly, VvibZIPC22 also increased the transactivation potential of known regulators of the general and specific flavonoid pathway genes, with the exception of the \textit{VviMYBA2–MYC1} complex which is specific for the regulation of the anthocyanin gene \textit{VviUFGT} (Hichri et al., 2010). This result, contrasting with findings in tobacco, might be explained by a limit in the transactivation capacity of the \textit{VviUFGT} promoter in this assay. Here, the availability of specific regulators is much higher than in reality and therefore there is no possibility of further activation by other factors such as VvibZIPC22.

Heterodimerization is a typical mechanism for regulating bZIP factor activity, and the formation of specific bZIP pairs is essential to exert their function (Llorca et al., 2014). In particular, in Arabidopsis (EhIert et al., 2006), bZIPs from clade S (clade C in \textit{V vinifera}, to which VvibZIPC22 belongs) form specific heterodimers with bZIPs from clade C (clade B in \textit{V vinifera}). This heterodimeric complex is also conserved in tobacco (Strathmann et al., 2001) and parsley (Rugner et al., 2001). Accordingly, the putative heterodimerization partners of VvibZIPC22 were searched for within the bZIPs of clade B (Liu et al., 2014). From phylogenetic and co-expression results, we obtained an indication of VvibZIPB38 as a putative heterodimerization partner of VvibZIPC22, to be experimentally validated.

The co-expression analysis also gave further insights into the VvibZIPC22 putative regulatory network. Remarkably, 40% of the 210 genes strongly co-expressed with VvibZIPC22 fell within the QTL regions recently associated with the fine regulation of anthocyanin and flavonol content in mature berries (Costantini et al., 2015; Malacarne et al., 2015). This percentage represents a significant enrichment over a random distribution of these genes along the genome, supporting their involvement in flavonoid metabolism. In addition, we noted that several genes with the highest correlation to the VvibZIPC22 profile coded for enzymes related to glucose metabolism. Interestingly, in Arabidopsis the expression of bZIP genes from class S1 was shown to be regulated by sugars and therefore to participate in a sugar-dependent control of target genes (Hanson et al., 2008). Overall, the results of our study point to a role for VvibZIPC22 in the regulation of different branches of the flavonoid pathway. In our hypothetical model, UV light and perhaps the glucose level promote the transcription of VvibZIPC22, which then regulates the expression of several genes of the flavonoid pathway, either alone or in combination with other factors (Fig. 7).

**Conclusions**

We have functionally characterized VvibZIPC22, a grapevine bZIP factor belonging to clade C in the phylogenetic reconstruction of the grapevine bZIP gene family. Our results indicated...
that, upon UV light stimulus, VvibZIPC22 regulates the content of different flavonoids, acting on promoters of general and specific genes of the pathway, alone or in combination with other factors. Altogether, our data provide new insights into the transcriptional control of the grapevine flavonoid pathway.

**Supplementary data**

Supplementary data are available at *JXB* online.

Table S1. Primers used in qRT-PCR analyses of grapevine and tobacco samples.

Table S2. Putative ACGT-containing elements in the promoters of the grapevine flavonoid pathway genes tested in transient reporter assays.

Table S3. Results of co-expression analysis obtained with VvibZIPC22 as input query.

Figure S1. Profiles of VvibZIPC22 relative expression and of anthocyanin 3-monoglucosides and flavan-3-ol monomers during Pinot Noir berry development.

Figure S2. Phylogenetic tree of clade B VvibZIP factors and of their putative orthologues.

Dataset S1. Codon-based alignments of CDS sequences of clade B VvibZIP genes and of their putative orthologues.

Dataset S2. Codon-based alignments of CDS sequences of clade C VvibZIP genes and of their putative orthologues.

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