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**Title: Combinatorial Regulation of Stilbene Synthase Genes by WRKY and MYB Transcription Factors  
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**Running head:** WRKYs and MYBs control stilbene synthesis in grape

**Corresponding author:** Alessandro Vannozzi; Department of Agronomy, Food, Natural resources, Animals,  
and Environment (DAFNAE), University of Padova, 35020 Legnaro, Italy; Tel: +393920094888; Email address:  
[alessandro.vannozzi@unipd.it](mailto:alessandro.vannozzi@unipd.it)

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**Title: Combinatorial Regulation of Stilbene Synthase Genes by WRKY and MYB Transcription Factors in Grapevine (*Vitis vinifera* L.)**

**Running head:** WRKYs and MYBs control stilbene synthesis in grape

Alessandro Vannozzi<sup>1\*+</sup>; Darren Chern Jan Wong<sup>2+</sup>; Janine Höll<sup>3</sup>; Ibrahim Hmam<sup>1#</sup>; José Tomás Matus<sup>4</sup>; Jochen Bogs<sup>3</sup>; Tobias Ziegler<sup>3</sup>; Ian Dry<sup>5</sup>; Gianni Barcaccia<sup>1</sup>; Margherita Lucchin<sup>1</sup>

<sup>1</sup>Department of Agronomy, Food, Natural resources, Animals, and Environment (DAFNAE), University of Padova, 35020 Legnaro, Italy.

<sup>2</sup>Ecology and Evolution, Research School of Biology, Australian National University Acton, ACT 2601, Australia.

<sup>3</sup>Centre for Organismal Studies Heidelberg, University of Heidelberg, 69120 Heidelberg, Germany.

<sup>4</sup>Centre for Research in Agricultural Genomics (CRAG) CSIC-IRTA-UAB-UB, Barcelona 08034, Spain.

<sup>5</sup>CSIRO Agriculture & Food, Urrbrae, SA 5064, Australia.

\*: Corresponding Author; Email: [alessandro.vannozzi@unipd.it](mailto:alessandro.vannozzi@unipd.it)

+: Equal contribution

#: current address: Department of Pomology, Faculty of Agriculture, Cairo University, 12613 Giza, Egypt

Abbreviations: AI, aliphatic index; bHLH, basic helix-loop-helix; CHS, chalcone synthase; CRE, *cis*-regulatory element; EF1, elongation factor 1; ERF, ethylene responsive factor; FC, fold change; FDR, false discovery rate; GCN, gene coexpression network; GRAVY, grand average of hydropathicity; GRsPaV, grapevine rupestris stem pitting-associated virus; GUS,  $\beta$ -glucuronidase; HsF, heat shock transcription factor; II, protein instability index; JA, jasmonic acid; LSD, least significant difference; LUC, luciferase; MR, mutual ranking; MUSCLE, multiple sequence comparison by log-expectation; MW, molecular weight; NGS, next generation sequencing; PCC, Pearson correlation coefficient; pI, isoelectric point; PKS, polyketide synthase; qPCR, quantitative PCR; RNA-seq, RNA sequencing; SA, salicylic acid; SE, standard error; SRA, sequence read archive; STS, stilbene synthase; TF, transcription factor; TFBS, transcription factor binding site; TFDB, transcription factor database; TPS, terpene synthase; TSS, transcription starting site; UV-C, ultraviolet C; UTR, untranslated region.

## Abstract

Stilbene synthase (STS) is the key enzyme leading to the biosynthesis of resveratrol. Recently we reported two R2R3-MYB transcription factors (TFs) that regulate the stilbene biosynthetic pathway in grapevine: *VviMYB14* and *VviMYB15*. These genes strongly co-express with *STS*s under a range of stress and developmental conditions, in agreement with the specific activation of *STS* promoters by these TFs. Genome-wide gene co-expression analysis using two separate transcriptome compendia based on microarray and RNA-Seq data revealed that WRKY TFs were the top TF family correlated with *STS* genes. On the basis of correlation frequency, four WRKY genes, namely *VviWRKY03*, *VviWRKY24*, *VviWRKY43* and *VviWRKY53*, were further shortlisted and functionally validated. Expression analyses under both unstressed and stressed conditions, together with promoter-luciferase reporter assays, suggested different hierarchies for these TFs in the regulation of the stilbene biosynthetic pathway. In particular, *VviWRKY24* seems to act as a singular effector in the activation of the *VviSTS29* promoter, while *VviWRKY03* acts through a combinatorial effect with *VviMYB14*, suggesting these two regulators may interact at the protein level as previously reported in other species.

**Keywords:** Gene co-expression, Network, Resveratrol, *Vitis vinifera*

## Introduction

In the last decade, the availability of an accurate grapevine (*Vitis vinifera* L.) genome assembly (Jaillon et al. 2007), together with the release of a detailed annotation, namely 12X.v2 (Vitulo et al. 2014) (recently updated with the release of VCost.v3; Canaguier et al. 2017) has been accompanied with a remarkable rise in genomic and transcriptomic data available for this species. As a matter of fact, grapevine represents one of the most representative examples of how next generation sequencing technology (NGS) massively impacts plant genomics and plant molecular biology (Fabres et al. 2017). Recently, network analyses have contributed to an increased understanding of the regulatory mechanisms that control grape berry development and composition (Ali et al 2010). In this sense, and based on the notion that genes involved in similar or related processes may exhibit similar expression patterns over a range of experimental conditions, an increasing number of studies have used gene co-expression networks to find common pathways and putative targets for transcription factors related to berry development and secondary metabolism (reviewed by Wong and Matus 2017).

The value of exploiting omics data in *Vitis* species relies on the fact that, wild and cultivate grapevines produce a vast array of chemical compounds many of which are related to wine quality and have been implicated in promoting human health (Wong and Matus 2017). Among these, stilbenes, a class of phenolic secondary metabolites characterized by the presence of a 1,2-diphenylethylene backbone, have been increasingly studied over the last decade because of their nutraceutical properties, with considerable potential in drug research (e.g. anticancerinogenesis; Ali et al. 2010, Pangen et al. 2014, Weiskirchen and Weiskirchen 2016) and also with important roles in the protection of plants against pests, pathogens and abiotic stresses (Chong et al. 2009, Jeandet et al. 2010).

Together with flavonoids, stilbenes belong to the plant polyketide class representing a major group of phenylpropanoids derived from the extension of the activated form of coumaric acid with three acetyl moieties. Apart from the *Vitaceae*, current literature indicates that stilbenes are produced by a polyphyletic group of species limited to approximately 50 plant families including dicotyledons, monocotyledons, conifers, liverworts, and ferns (Pangen et al. 2014, Weiskirchen and Weiskirchen 2016). Despite the multiplicity of stilbene units found in different plant species (Shen et al. 2009, Rivière et al. 2012), most of them (including those in grapevine), are derivatives of the basic unit trans-resveratrol (3,5,4'-trihydroxy-*trans*-stilbene). Resveratrol has been the subject of numerous research studies since it was postulated to have a role in the so-called 'French paradox', which refers to the observation that French people have a relatively low incidence of coronary heart disease, despite having a diet rich in saturated fats. In fact, hundreds of subsequent studies have reported that this compound can prevent or slow down a variety of diseases, including cancer, diabetes, as well as extend the

lifespan of various organisms (Pengen et al. 2014, Weiskirchen and Weiskirchen 2016, Tzai et al. 2017). The biosynthesis of resveratrol is achieved through a small branch of the general phenylpropanoid pathway and can be considered as a competitive extension of the flavonoid branch (Vannozzi et al. 2012). Stilbene synthase (STS) is the key enzyme leading to the production of resveratrol and belongs to the chalcone synthase (CHS) superfamily of type III polyketide synthases (PKSs; Chong et al. 2009). In grapevine, an analysis of the *STS* multigenic family based on both the PN40024 and PN ENTAV 115 genomes (Jaillon et al. 2007, Velasco et al. 2007) led to the identification of 48 putative *STS* gene sequences, with at least 33 encoding full length STS proteins (Vannozzi et al. 2012).

Two R2R3-type V-myb myeloblastosis viral oncogene homolog (MYB) transcription factors (TFs) have been shown to regulate stilbene biosynthesis in the grapevine (Höhl et al. 2013). These R2R3-MYB-type TFs, designated as *MYB14* and *MYB15*, are found to be strongly co-expressed with certain *STS* genes in different grapevine organs in response to biotic and abiotic stress including downy mildew (*Plasmopara viticola*) infection, mechanical wounding and exposure to UV-C irradiation (Höhl et al. 2013). The expression of *MYB14* and *MYB15*, is also correlated with the accumulation of *trans*-piceid in developing grape berries (Höhl et al. 2013). Furthermore, grapevine cell cultures transiently expressing *STS promoter::luciferase* reporter constructs showed considerable induction of the activities of the promoters of *STS29* and *STS41*, whenever co-transfected with *MYB14* and *MYB15*. The involvement of these TFs in the regulation of stilbene biosynthesis *in planta* was demonstrated using transgenic grapevine hairy roots overexpressing *MYB15*, which showed an increased accumulation of *trans*-piceid, associated with an up-regulation of *STS29* and *STS41* transcription levels. Furthermore, Fang et al. (2014) demonstrated that the variation in expression of *MYB14* correlated with the variation in resveratrol content in two grapevine cultivars (*Vitis monticola* x *Vitis riparia* - high resveratrol producer and *V. vinifera* - low resveratrol producer) and, using a one-hybrid yeast assay, showed that MYB14 directly interacts with the *STS* promoter *in vitro*. This study also demonstrated that a transient overexpression of *MYB14* could activate *STS* expression in grapevine leaves and that its overexpression in transgenic *Arabidopsis* could activate GUS expression driven by a *STS* promoter. Wong et al. (2016) showed that these two TF genes shared close similarity (in sequence and expression) with *MYB13*, suggesting that, in addition to MYB14 and MYB15, MYB13 may be also involved in the transcriptional regulation of at least some *STS* genes in grapevine. Recently, a composite network for STS regulation was constructed with the aim of illustrating different approaches of data integration for network analysis in grapevine (Wong and Matus. 2017). Using publicly available berry-specific RNA-Seq data, the authors overlapped gene co-expression networks with the presence of promoter *cis*-binding elements (CRE), microRNAs and long non-coding RNAs. As a result, a systems-level STS

regulatory network was inferred from the context of berry development and ripening. However, this network still needs to be demonstrated to operate *in planta*.

The present study was aimed at extending the current knowledge pertaining to the regulation of stilbene biosynthesis in grapevine, by identifying and characterizing TFs, other than R2R3-MYBs, that are potentially involved in the regulation of the stilbene biosynthesis. To do this we performed a large-scale co-expression analysis identifying novel candidate TFs belonging to different gene families. Amongst these was the WRKY TF family, which represented one of the most enriched families in terms of correlation frequencies with *STS* genes. Based on network connectivity properties, we selected *WRKY03*, *WRKY24*, *WRKY43* and *WRKY53* for further examination. Expression analyses of grapevine tissues under different stress and unstressed conditions, together with functional reporter gene assays suggest different roles for these TFs in the regulation of the *STS* pathway.

## Results and Discussion

### An integrated co-expression network confirms and identifies potential regulators of *STS* expression

The flavonoid biosynthetic pathway is considered one of the best systems available for studying the regulation of gene expression in plants (Davies and Schwinn 2003) and also in grapevine (*V. vinifera* L.), where it represents one of the most studied crops in this regard. Although many TFs involved in the regulation of specific flavonoid structural genes in grapes have been identified, there is evidence to suggest that novel regulators remain to be characterized (Wong and Matus 2017). To investigate this, we first constructed two independent global gene co-expression networks (GCNs) based on mutual ranking (MR) using datasets produced with Next Generation Sequencing (NGS) (21 experiments, 235 conditions averaged from 654 RNA-Seq assays; Supplementary Table S1) and microarrays (23 experiments, 359 conditions averaged from 914 arrays; Supplementary Table S1). The choice of the MR index over Pearson correlation coefficient (PCC) as the preferred co-expression measure in this study is supported by previous studies showing that the latter is more robust to outliers and has higher predictive power in gene function inference compared to correlation-based metrics (Obayashi et al. 2009).

For the construction of a robust *STS* GCN, we first determined the optimal threshold of neighborhood size,  $k$  aimed at maximizing the number of genes included while keeping potential false-positives inclusions to a minimum. We began by investigating the relationship of different  $k$  thresholds (e.g.  $k$  of 100 to 1000) on the distribution of PCC values for each member of the grapevine *STS* family described in Vannozzi et al. (2012) in the microarray and NGS *STS* GCNs separately (See materials and methods). At a  $k$  of 300, a widely-adopted limit for establishing a practical size of co-expressed genes lists for functional validation (Obayashi and Kinoshita, 2010, Aoki et al. 2016), the median PCC value in the observed microarray *STS* and random

microarray GCN was 0.57 and 0.29, respectively (Supplementary Figure 1A). The same  $k$  in the observed *STS* and random NGS GCN showed a median PCC of 0.67 and 0.01, respectively (Supplementary Figure 1B). In the microarray GCN, overlaps in PCC distribution between the observed *STS* and random GCN was minimal when  $k \leq 300$ . However, establishment of an appropriate  $k$  is not as straightforward for the *STS* NGS GCN. The observed PCC distribution in the latter was generally high (PCC between 0.64 and 0.66) even at high  $k$  ranges (e.g.  $k > 500$ ) while the PCC distribution in random GCNs were often close to 0 across all  $k$  ranges, complicating the selection of an appropriate  $k$  with this approach. Several studies have shown that RNA-seq derived PCC GCN can have skewed PCC density distributions (in both negative and/or positive directions) and that the PCC measure is often sensitive to differences in sample sizes (Giorgi et al. 2013, Huang et al. 2017, Wisecaver et al. 2017). For these reasons, the rank of correlations (e.g. MR) are often preferred in recent co-expression studies. As an additional measure to guide the choice of choosing the appropriate  $k$  threshold, we established the statistical significance of MR values in the global microarray and NGS GCN, by analyzing the distribution of MR values over 1,000 permutations of the respective dataset. The analysis revealed that  $MR \leq 383$  and  $\leq 417$  are significant at  $P < 0.01$ , for the global microarray and NGS GCN, respectively (Supplementary Figure 1C-D). These results are comparable with earlier studies across a wide range of plant GCNs constructed with correlation-ranked metrics at stringent thresholds (e.g.  $P < 0.01$ ) (Wong et al. 2014, Wong et al. 2013, Mutwil et al. 2011). Nonetheless, weaker MR (e.g.  $MR < 1000$ ) observed at  $P$  between 0.01 and 0.05 in both GCNs (Supplementary Figure 1C-D) may still be statistically reliable and biologically meaningful (Obayashi and Kinoshita, 2010, Aoki et al. 2016). By combining clues garnered from both statistical approaches, we determined that a top  $k$  threshold of 200 would be suited for the construction of a robust *STS* GCN from both platforms. This is supported by the fact that nearly all observed MR values are significant at  $P < 0.01$  (with the exception of 123 of 9000 co-expression gene pairs in the NGS *STS* GCN; Supplementary Table S2) and a minimal overlap of PCC distribution (across quartiles) between the observed and permuted GCN (Supplementary Figure 1C-D), in both platforms. Furthermore, a  $k$  of 200 is within reasonable limits for designing functional studies (Obayashi and Kinoshita, 2010, Aoki et al. 2016). Similar thresholds have also been used in prioritizing candidate genes in recent grape functional studies (e.g. top100 for microarray co-expressions in Wong et al. 2016, and top300 for RNA-seq co-expressions in Loyola et al. 2016 and Sun et al. 2018). By merging the two independent *STS* co-expression modules obtained from both platforms ( $k$  of 200) into a combined GCN (Supplementary Table S2), we hypothesized that an integrated *STS* GCN would hold more biologically meaningful co-expression relationships as GCNs constructed from different platforms have the potential to highlight additional functional categories and co-expression relationships (Giorgi et al. 2013). To



assess TF-STS co-expression, we kept only those TF accessions corresponding to predicted grapevine TFs based on the Plant transcription factor database (Plant TFDB; Jin et al. 2017), encompassing 1,256 grapevine TFs distributed among 58 families. In addition, as dense connections between *STS* genes are generally observed in the *STS* GCN (Supplementary Table S2), we hypothesized that *bona fide* regulatory genes involved in *STS* regulation should be frequently co-expressed with multiple *STS* members. As such, we only considered those TFs showing a node degree higher than or equals to 5, i.e. those TFs correlated with at least 5 *STS* genes (10% of total grapevine STS). A list of all top200 STS-related TFs identified independently by the degree is reported in Supplementary table S3. Initial inspection of the exclusive filtered STS-TF CGN showed a strong correlation between 42 *STS*s and 31 TFs, connected by 569 edges (Fig. 1).

Shared edges between RNA-Seq and microarray networks accounted for 75.8% of the total number of edges and 63.8 % and 63.5% of the total microarray and RNA-Seq specific edges, respectively. Within the whole STS-TF GCN, we identified TFs belonging to 8 different families. The most highly represented family comprised the WRKY TFs, with 10 genes (30% of all TFs in the GCN), followed by Ethylene Responsive Factors (7 genes; 23.3%), MYBs (6 genes; 20%), NACs (4 genes; 13.3%), GRASs (2; 6.6%), C2H2, HSF and bHLH (1 gene; 3.3%). Table 1 provides a list of all the TFs co-expressed with *STS* genes and represented in the GCN.

Considering the contribution of edges from both datasets, nodes belonging to the shared-interaction module showed a higher number of interactions with *STS*s (Fig. 1) compared to nodes belonging exclusively to the microarray or RNA-Seq modules. This is the case of *MYB14* (VIT\_07s0005g03340), the TF with the highest number of interactions, showing up to 72 edges. *MYB15* is another direct regulator of *STS* transcription (Höll et al. 2013) that appeared within the top200 *STS*-co-expressed genes. Differing from its paralog (both genes belong to R2R3-MYB Subgroup 2), *MYB15* was represented only within the microarray-interaction module, and the number of edges was much lower compared to *MYB14*. This reveals a potential limitation of the GCN approach as the network output can be influenced by the type of experimental conditions used in GCN construction (Usadel et al. 2009) or that the majority of experimental conditions used to generate the RNA-Seq and microarray datasets do not constitute tissues and treatments in which MYB15 has a functional role.

APETALA2/ethylene response factor (*AP2/ERF*) transcription factor TFs were also highly represented in the GCN including *VvERF098* (VIT\_07s0005g03220), *VvERF112* (VIT\_01s0150g00120), *VvERF113* (VIT\_07s0031g01980) and *VvERF114* (VIT\_18s0072g00260), all belonging to the group X ERF subfamily (Licausi et al. 2010). The members of this ERF subfamily are involved in the plant response to abiotic stresses, including drought and salinity (Fujimoto 2000) and they are expected to be involved in gene regulation under stress conditions involving both ethylene-dependent and -independent pathways (Mizoi et al. 2012). As a matter

of fact, grapevine *STS* genes have been previously demonstrated to be induced in grapevine leaves following treatment with ethephon, an ethylene releasing compound (Belhadj et al. 2008, Becatti et al. 2014), suggesting the signalling pathway related to this hormone could be involved in the activation of the plant stress-response related to stilbene accumulation. This observation, together with the presence of other TFs such as AP2/ERFs within the GCN, leaves open the possibility that the stilbene biosynthetic pathway could be regulated by many different TFs and hormone signalling pathways. Indeed, recent surveys show various AP2/ERF TF binding sites (TFBS) are present within the 1kb promoter region of many grapevine *STS* promoters (Wong and Matus, 2017).

The GCN analysis also identified genes involved in plant stress responses that represent interesting candidates for further analyses. For example, the second largest node in term of degree within the shared-interaction module was VIT\_08s0007g08750, a gene that encodes the heat shock transcription factor (HsF), *VviHSFA3a*, one of 19 *HsF* genes predicted within the grapevine genome (Hu et al. 2016). Plant HsF proteins function as TFs regulating the expression of heat shock proteins and other general stress related genes such as the non-chaperone encoding genes *GOLS1* (galactinol synthase 1) or *APX2* (ascorbate peroxidase 2) (Scharf et al. 2012). *VviHSFA3a* belongs to the grapevine *HsF* subgroup A, and was recently demonstrated to be highly responsive to ethylene treatments in *V. pseudoreticulata* (*VpHsf3a*; Hu et al. 2016). This is an interesting observation considering the high number of ethylene responsive factors (ERFs) genes that we identified in the co-expression analysis and the fact that also grape *STSs* are also known to be induced by ethylene (Belhadj et al., 2008). HsFs play pivotal roles in adaptation to heat stress and other stress stimuli including cold, salt, drought, and oxidative stresses (Hu et al., 2016). Many of these abiotic stresses also cause the induction *STS* genes (Vannozzi et al. 2012, Corso et al. 2015).

Another gene of interest identified in the shared-interaction module is VIT\_11s0016g02070 which encodes a basic helix-loop-helix (bHLH) protein. The bHLH proteins are a superfamily of TFs that are important regulatory components of many transcriptional complexes, controlling processes such as regulation of flavonoid biosynthesis, floral organogenesis, hormone and light signaling responses and epidermal cell fate determination such as trichome, root hair and stomata formation (Toledo-Ortiz 2003, Sema 2007). It has been demonstrated in various species including maize (*Zea mays*), petunia (*Petunia hybrida*), snapdragon (*Antirrhinum majus*), Arabidopsis (*Arabidopsis thaliana*) and grapevine that MYC-like bHLHs generally interact with R2R3-MYB and WD40 proteins to regulate structural genes involved in flavonoid metabolism (reviewed in Chezem and Clay 2016). In grapevine, VviMYC1, one of the predicted 115 bHLHs based on the Plant TFdb (Jin et al. 2017) has been demonstrated to regulate the anthocyanin and proanthocyanidin pathways by interacting with different flavonoid R2R3-MYB activators (Hichri et al. 2010, Matus et al. 2017) and repressors (Cavallini et al. 2015).

Although our previous research suggested that regulation of stilbene biosynthesis in grapevine by R2R3-MYB TFs was bHLH-independent (Höller et al. 2013), we detected a high level of co-expression between *VIT\_11s0016g02070*, *VvSTSs* and *VviMYB14/15* (both in term of PCC and degree). Further work needs to be undertaken to determine the potential role of *VIT\_11s0016g02070* in the direct or indirect regulation of stilbene biosynthesis.

### **Potential dual regulation of *STS* by MYB and WRKY transcription factors**

The strong co-expression relationships present in the integrated TF-STS GCN is also exemplified by a highly modular GCN between these TFs (Fig. 2). The inferred TF-TF GCN contained 31 genes connected by 191 edges and was organized in three densely connected modules consisting of 13 (module 1), 11 (module 2), and 7 (module 3) genes. The analysis confirmed known and putative *STS* regulators such as *MYB14*, *MYB15*, and *MYB13*, all partitioned to Module 1, with *MYB14* having the highest node degree connecting 9 members module 1, but also to others (i.e. 6 members of module 2, and 3 members of module 3). Meanwhile, three of the four WRKY TFs partitioned to module 2 such as *WRKY02* (15 TFs/genomes), *WRKY23* (12 TFs/genomes), and *WRKY43* (11 TFs/genomes) were also among the top 10 TFs sharing the highest connectivity in the network. The presence of multiple modules with overlapping connectivity also suggests the presence of several regulatory networks that may function in controlling both unique and/or overlapping sets of *STSs*. The high level of connectivity common with *MYB14* and several WRKYs lead us to hypothesize a strong likelihood of combinatorial and synergistic regulation of grapevine *STS* genes by members of these two TF families.

### **Phylogenetic analysis of WRKY genes highly co-regulated with *STS* genes**

Based on our GCN analysis, genes encoding WRKY (12 genes; 10 with degree > 5) is the top TF family connecting *STS* genes irrespective of the platforms used in the network construction (Fig. 1), suggesting that regulation of *VvSTS* genes could be orchestrated by WRKY TFs. This hypothesis is reinforced by the observation that WRKY cis-regulatory elements (CREs) are found in the promoters of many *VvSTS* members (Wong and Matus, 2017).

*WRKY* genes are classified into three main groups (I, II and III) based on the number of WRKY domains and the pattern/position of their zinc finger motifs (Eugelm et al. 2000). Group I WRKYs typically contain two WRKY domains whereas group II and group III members contain a single WRKY domain. Group II WRKYs can be further sub-divided into five subgroups: IIa, IIb, IIc, IId, and IIe (Zhang et al. 2005). Based on phylogenetic analysis of the grapevine WRKY family classification performed by Wang et al. (2014) the WRKY genes found

to be co-expressed with *STS* belong mainly to the WRKY subgroup II (Fig. 3A, Table 1), with four genes belonging to group IIb (*VviWRKY02*, *VviWRKY39*, *VviWRKY29*, and *VviWRKY53*), three genes belonging to group IIc (*VviWRKY03*, *VviWRKY43*, and *VviWRKY11*), and two genes belonging to group IIe (*VviWRKY4* and *VviWRKY30*). Only one co-expressed WRKY TF was found to belong to group I (*VviWRKY24*).

Amongst candidate WRKY TFs, we focused our investigation on four genes, namely *VviWRKY03*, *VviWRKY24*, *VviWRKY43* and *VviWRKY53*. *VviWRKY03* and *VviWRKY24* were the top two candidates frequently correlated with *STS* gene expression in the RNA-seq compendia while *VviWRKY43* and *VviWRKY53* were among the top three inferred from the microarray compendia (see Fig. 1). These four candidates are also uniquely positioned across all three modules in the integrated TF-TF GCN (Fig. 2): *VviWRKY03* in module 1 along with the known *STS* regulators (*VviMYB14* and *VviMYB5*), *VviWRKY53* in module 2, and *VviWRKY24* and *VviWRKY43* in module 3, the module containing the highest number of WRKY TFs. To provide additional support for this selection, phylogenetic analyses of deduced protein sequences of *VviWRKY43* and *VviWRKY03* show them to be most closely related to AtWRKY75, sharing 48% and 53% amino acid identity, respectively. Both genes have similar genomic organizations, containing one single intron (phase 2), and encoding for small proteins (189 aa and 182 aa, respectively). *VviWRKY53*, whose closest homolog in the Arabidopsis genome is AtWRKY72 (36% amino acid identity), possess five “phase 0” introns and encode for a much larger protein (605 aa) (Fig. 3C).

Finally, *VviWRKY24* is the orthologue of AtWRKY33, encoding a 552 aa protein. Sequence analysis of these *VviWRKY*s using PSORT program (Nakai et al. 1999) confirmed the presence of putative nuclear localization signals in all the TFs considered (RKPR for both *VviWRKY03* and *VviWRKY43* at position 83; PTKKKVE for *VviWRKY24* at position 261, PAKRCRV for *VviWRKY53* at position 240).

Figure 3B illustrates the results of a phylogenetic analysis based on the predicted translation product of *WRKY* genes identified to be co-expressed with *STS* genes in our GCN, together with WRKY TFs which have already been functionally characterized in grapevine or in other plant species. Interestingly, 27 WRKYs enclosed in the phylogenetic analysis are involved in the regulation of secondary metabolism including many branches of the phenylpropanoid pathway, whereas the remaining proteins have been related to the plant response to both biotic and abiotic stresses. The comparison of candidate grape WRKYs involved in *STS* transcription with those characterized in other species (Fig. 3B) indicated both *VviWRKY03* and *VviWRKY43* as closely related to *Captis japonica* CjWRKY01, *Solanum tuberosum* StWRKY1 TFs and also to the grape *VviWRKY52* (previously named WRKY1 by Marchive et al. 2007, 2013). CjWRKY01 has been shown to have a role in the transcriptional regulation of several structural genes involved in the biosynthesis of the alkaloid barberine (Kato et al. 2007) whereas the tomato StWRKY1 TF is involved in the regulation of hydroxycinnamic acid amid

(HCAA) biosynthetic genes and in the cell wall straightening upon *Plasmopara infestans* invasion (Yogendra et al. 2015). Overexpression of *VviWRKY1* (*VviWRKY52*) in tobacco has been shown to improve resistance to pathogenic fungi such as *Phytophthora* and to oomycetes such as *Peronospora tabacina* (Marchive et al. 2007), whereas in grapevine it was associated with the transcriptional regulation of three genes putatively involved in the Jasmonic acid signalling pathway and in the reduced susceptibility to downy mildew (*Plasmopara viticola*) infection (Marchive et al. 2013).

*VviWRKY24* is a putative orthologue of the Arabidopsis *AtWRKY33* gene, known to be involved in many processes including heat and NaCl tolerance, redox homeostasis, resistance to *Botrytis cinerea* and *Pseudomonas syringae*, SA signaling, ethylene-JA-mediated cross-communication and camalexin biosynthesis (Birkenbihl et al. 2012). In grapevine *VviWRKY24* (known as *WRKY33* in the nomenclature introduced by Merz et al. 2015), is associated with an increased resistance to *P. viticola* infection in the susceptible cultivar (cv.) ‘Shiraz’ and seems to be functionally related to defense. Finally, *VviWRKY53* is closely associated with *SlWRKY73*, which was found to transiently trans-activate a tomato terpene synthase (TPS) gene in *Nicotiana benthamiana* leaves (Spyropoulou et al. 2014).

#### **Expression of selected *WRKY* genes correlates with *MYB14/15* and *STS* transcripts under biotic and abiotic stress**

We further explored the tight relationship observed between *STS* and their candidate regulators using the datasets from which the GCNs were constructed. We extrapolated the expression patterns of *STSs*, *MYBs* and *WRKYs* from a subset of biotic stressed samples (Fig. 4), showing that most *STS* genes are induced upon a range of different biotic stresses including infection with the necrotrophic fungus *Botrytis cinerea*, the biotrophic powdery mildew (*Erysiphe necator*) and oomycetes such as downy mildew (*P. viticola*). Of particular interest was the transcriptional regulation of a subgroup of *STS* genes that form a small cluster on chromosome 10 (*STS1-6*). These genes showed a much lower induction in response to downy mildew infection in comparison to most of the other members of the *STS* gene family (which cluster in a 500 Kb region on chromosome 16). This supports the previous report of Vannozzi et al. (2012), who showed that members of this small cluster of *STS* genes on chromosome 10 genes were less responsive to *P. viticola* infection than *STS* genes on chromosome 16. However, Fig. 4 clearly shows that both sets of *STS* genes are equally responsive to other biotic stresses. No induction of *STSs* was observed in response to infection with GRSPaV (Grapevine rupestris stem pitting-associated virus) (Fig. 4). This is in agreement with the work of Gambino et al. (2012) who observed that genes

involved in stress and pathogen responses are downregulated in the presence of co-evolved viruses such as GRSPaV.

Looking at the transcriptional response of the selected R2R3-MYB and WRKY TFs, we observed a good correlation with *STS* transcription (Fig. 4). *MYB14/15* were induced whenever *STS*s were induced, as previously reported in Höll et al. (2013). Similarly, transcription of several *WRKY* genes also showed a high level of correlation with *STS* genes. This was particularly evident for *WRKY03*, *-43* and *-24*. The induction of *WRKY24* in response to biotic stress previously reported by Merz et al. (2015), who observed a strong up-regulation in leaves infected with downy mildew in the resistant cv. ‘Regent’. A less clear relationship was observed between transcription of *STS* family members and *MYB13* or *WRKY53*. For example, both TF genes were down-regulated in powdery mildew-infected leaves of some grapevine accessions while the former is also down-regulated in the late stages of *Botrytis*-infected berries (*B. cinerea* S2, and S3; Blanco-Ulate et al. 2015). As suggested previously, MYB13 may be responsible for *STS* regulation under basal (non-stressed) conditions or to developmental transitions.

Under abiotic stress conditions, the correlation between transcription of *STS*s and the candidate TF regulators was less evident compared to biotic stress (Figure 5). Nonetheless, a clear induction was found under drought stress during the ripening of white berries, late véraison heat stress of berries, and UV-C irradiation in berry skins, and in response to hormone (i.e. Gibberellic acid) treatments in flowers.

To further validate the correlations observed in the GCNs and heatmaps, we analysed the transcript levels of a subset of WRKY candidates (i.e. *WRKY03*, *WRKY43*, *WRKY53*), *MYB14* and *MYB15* TFs, and three highly responsive *STS*s (*STS29*, *STS41* and *STS48*), in cv. ‘Pinot Noir’ leaves exposed to wounding and UV-C treatments by qPCR (Fig. 6). As previously reported in Höll et al. (2013) since the grapevine *STS* family is composed of 48 closely related genes (Vannozzi et al. 2012, Parage et al. 2012), it was not possible to design sequence specific-primers for the detection of only one *STS* isoform. Therefore, primers STS41-F/R detect isoforms *STS41* & 45, while the primers STS29-F/R detect isoforms *VvSTS25*, 27 & 29. The results of the qPCR analysis confirmed our GCN analysis, showing a marked co-induction between the *STS*.

In response to mechanical wounding the transcript level of *VvSTS29*, *-41* and *-48* gradually increased over a 48-hour period, reaching a peak at 48h. *STS29* was the *STS* member showing the highest induction in terms of normalized transcript level, showing a fold change (FC) of 2400 times higher respect to the unwounded leaf (T0) followed by *STS41* (FC  $\approx$  1400) and *STS48* (FC  $\approx$  750). Looking at the expression of *VvSTS* candidate TF regulators, both WRKY and R2R3-MYB TFs were induced under wound stress. Similar to what was observed by Höll et al. (2013), *MYB14* and *MYB15* were both induced upon stress reaching their peaks at 48 h. *MYB14*

reached higher values compared to *VviMYB15* although, looking at the fold change respect to the 0h time point, both TFs reach similar values at their peak ( $FC \approx 60-80$ ). *WRKY03* showed the highest and most significant induction reaching a peak at 48 h corresponding to a FC of 495. *WRKY43* and *WRKY53* showed a lower but progressive increase over the 48h time course, peaking at the last stage ( $FC= 61$  and  $115$ , respectively). In the UV-C treatment, gene expression was plotted as a log2 fold change between UV-C treated and untreated leaf discs (Fig. 6). In the UV-C treatment, gene expression was plotted as a log2 fold change between treated (UV-C) and untreated (i.e. wounded) samples at the same time point (Fig. 6). Thus, it must be noted that the lower fold change values observed for UV-C treated samples shouldn't be ascribed to lower responsiveness of candidate genes to the irradiation treatment per se, but to the fact that the untreated samples already showed a very high expression for these genes. *STS29* and *STS48* reached their peaks of induction at 6h ( $FC \approx 39$  and  $12$  respectively), followed by a slight decrease at 24 h. *STS41* showed a gradual increase reaching its peak at 24h ( $FC \approx 41$ ). *WRKY03* reached the maximum induction at 3h ( $FC \approx 3$ ), maintained this level until 6h ( $FC=2.81$ ) and then decreased at 24h ( $FC=1.94$ ). *WRKY43* reached a first peak of fold change value at 3h ( $FC=6.93$ ), followed by a slight decrease at 6 h with a fold change of over 4, then a higher peak at 24h ( $FC=18.49$ ). A similar trend was observed for *WRKY53*, which reached a first peak of expression at 3h, followed by a slight decrease at 6h, and by a second higher peak at 24h ( $FC \approx 5$ ). *MYB14* showed a progressive increase in expression, reaching its peak at 24h ( $FC \approx 5$ ), whereas *MYB15* reached the maximum expression 3 h after the imposition of the stress, with a fold change of approximately 14, and maintained this level in the following hours. We also evaluated the effects of the wound and UVC stress treatments in a shorter time frame, i.e. within the first 10 hours after the stress imposition (Supplementary figure S2) conforming the induction of *VvSTSs*, *VviMYBs* and candidate *VviWRKY* TFs is coherent with what reported in Fig. 6 also at earlier time-points.

### **Singular and combinatorial roles of WRKY and R2R3-MYB transcription factors in STS regulation in grapevine**

To assess whether WRKY TFs are able to regulate the transcription of *STS* genes in grapevine cells, we performed transient gene reporter assays using the *VviSTS29* gene promoter. We selected this gene because it belongs to the highly responsive stilbene synthase subgroup B and showed a high correlation with candidate *WRKYs*. The dual luciferase reporter assay has been used previously to functionally validate the role of many other transcriptional regulators of the flavonoid pathway, including *VviMYB14* and *VviMYB15* (Höll et al. 2013).

A ~1.2 Kb promoter fragment of *VviSTS29* gene isolated previously (Höll et al. 2013) was comprehensively screened for canonical MYB (i.e. type I – CNGTTR, II – TNGTTR, and IIG/AC-element – CCWAMC; where N=A/C/G/T, R=A/G, W=A/T, M=A/C) and WRKY (i.e. TTGACY; where Y=C/T) TFBS. A total of two, six, and two type I, II, and IIG MYB binding sites respectively and two WRKY TFBS were identified (Fig. 7A; Supplementary Table S4). Interestingly, the type IIG/AC-element TFBS was situated in close proximity ( $< \pm 50$  bp) with the two WRKY binding sites. Many studies have now established that functional combinatorial relationships between multiple TFBS are widespread across plant promoters, and this property play a key role in determining the transcriptional dynamics of organ-, tissue-, and/or stress-specific gene expression in plants (Vandepoele et al. 2006, Maruyama et al. 2012, Wong et al. 2017). Distance constraint ( $< \pm 100$  bp) between multiple TFBS is also essential for their functionality (Vandepoele et al. 2006) and are strong indicators of interacting TFs (Yu et al. 2006a, 2006b). Therefore, the co-occurrence of multiple MYB and WRKY TFBS and a strong distant constraint between them ( $< \pm 100$  bp) observed for *STS29* promoter provides support for both singular and combinatorial control of *STS29* by MYB and WRKY TFs that may be potentially mediated by MYB and WRKY TF interaction.

To test for WRKY and MYB activation of the *VviSTS29* promoter, transient expression assays were conducted on cv. ‘Chardonnay’ berry suspension cell cultures using a dual reporter luciferase system as previously described (Höll et al. 2013). The *VviSTS29* promoter sequence was fused to the *Firefly LUCIFERASE (LUC)* gene and co-transfected in cells with candidate TFs. Candidate TFs were cloned into thepART7 vector (Gleave 1992) under the control a 35S promoter and transfected in cells as single or combined TFs (Fig. 7B-C). Chardonnay cell suspensions transiently expressing the *proSTS29:LUC* luciferase reporter construct showed significant increases in the *STS29* promoter activity of approximately 4-fold when co-transformed with *VviMYB14* and 5-fold when co-transformed with *VviMYB15*, in line with previous results (Höll et al. 2013). Of the candidate WRKY TFs tested, a statistically significant induction of *VviSTS29* promoter activity was only observed in cells co-transfected with *VviWRKY24*, which led to a 4-fold increase in the luciferase activity. This induction of the *VviSTS29* promoter by *VviWRKY24* is comparable to the activation observed with *VviMYB14* and *VviMYB15*.

None of the other candidate WRKY TFs analyzed were found to produce a statistically significant effect on the *VviSTS29* promoter activity. This includes *VviWRKY52*, which was not co-expressed with any *STS* genes in the combined GCN, and thus serves as a null candidate without evident roles in directly regulating *STS* expression including *STS29*. Despite this, we observed several interactions when WRKY and R2R3-MYB TFs were co-transfected (Fig. 7). A statistically significant increase in the *VviSTS29p* luciferase activity was observed when



VviMYB14 was transfected with VviWRKY03 leading to two-fold increase when compared to VviMYB14 alone, corresponding to an 8-fold increase compared to the control. This observation suggests a combinatorial effect of VviWRKY03 and VviMYB14 in the regulation of the pathway, or at least of this particular VviSTS member, which may be specific to the regulatory networks implicated in module 1 (Fig. 2). This observation also raises the question whether these MYB and WRKY proteins could interact. Similar results were observed in *Petunia hybrida*, where the WRKY transcription factor PH3 interacts with a MYB-bHLH-WD40 complex (MBW), constituted by PhPH4, PhAN1, and PhAN11 encoding for a MYB, a bHLH and a WD40, respectively, and activates downstream genes in multiple distinct pathways involved in flower pigmentation and seed development (Verweij et al. 2016). Stilbene biosynthesis in grapevine is spatially and developmentally regulated and additionally induced by many abiotic and biotic environmental cues, which needs a complex regulatory network. The combinatorial regulation of *VviSTS29p* by VviMYB14 and VviWRKY03 and its induction by VviWRKY24 could be part of this network leading to fine adjustments of stilbene biosynthesis in respect to changing developmental and environmental conditions.

#### Concluding remarks

A systems-oriented study encompassing genome-wide gene co-expression (GCN) analysis, integrated GCN, phylogenetics, and DNA-binding motif analysis, was performed with the ultimate goal of identifying novel transcriptional regulators of the grapevine stilbene biosynthetic pathway. In this study, the use of the integrated TF–STS network has provided an added advantage of revealing additional co-expression relationships between transcription factors and STS genes that may have not been detected had only a single platform been used for GCN analysis (Wong et al. 2016, Wong and Matus, 2017). In Arabidopsis, formal assessments have shown that RNA-Seq GCN can be accurate, satisfying both biological and robust network topology properties, while revealing novel functional gene neighborhoods missed in microarray-based GCN (Giorgi et al. 2013).

The integrated TF–STS network analysis indicated a number of TFs belonging to different families, including WRKYs, MYBs and an ERF, that are putatively involved in the regulation of the grapevine STS multigenic family. Amongst the best candidate regulatory genes identified by this analysis was *VviMYB14*, belonging to the R2R3-MYB family, for which a role in the transcriptional regulation of at least two STSs has already been documented (Höll et al. 2013). This observation further validates the choice and validity of GCN analysis in gene function prediction used in this and other studies (Usadel et al. 2009). Amongst the candidate TFs inferred from the combined grapevine STS and TF GCN, we focused on members of the WRKY family which collectively showed the highest correlation with STS gene expression under a stringent connectivity threshold as well as the

well-documented roles of this TF family in the regulation of stress related genes in plants (reviewed in Jiang et al. 2017). Most of the WRKY TFs found as co-expressed in our GCN analysis were found to be potential orthologues of genes already characterized in grapevine or in other plant species, involved in biotic and abiotic stress responses, in signalling pathways related to the response to exogenous stimuli, and in the biosynthesis of different families of secondary metabolites.

Detailed analysis was carried out on four *WRKY* genes (*WRKY3*, *WRKY24*, *WRKY43* and *WRKY53* according to the nomenclature proposed by Wang et al. 2014) based on their level of PCC correlation and on the number of interactions they showed with *STSs*. Generally, the observations obtained by the meta-analysis of two large gene expression compendia used in the GCN construction and the quantitative PCR analyses performed on stressed leaves (wounded and UV-C treated) confirmed that, together with *VviMYB14* and *VviMYB15*, these *WRKY* TFs were induced whenever *STSs* and *R2R3-MYBs* were induced. This observation further reinforces their coordinated regulation, especially under stress, and strongly suggests a role in the regulation of the stilbene biosynthetic pathway.

Functional validation of candidate WRKYs indicated both a singular and combinatorial role for several members. In particular, *VvWRKY24*, an orthologue of Arabidopsis *WRKY33*, was found to have a direct effect on the promoter activity of *VvSTS29*, independent of *VvMYB14* and *VvMYB15*. The fact that stilbenes act as phytoalexins in grapevine and the phylogenetic relatedness between *VviWRKY24* and *AtWRKY33*, suggests some similarities with the regulation of the PTI (pattern triggered immunity) response in Arabidopsis (Jiang et al. 2017). Interestingly, *VviWRKY3*, which had no effect on *STS29* promoter activity on its own, appeared to act synergistically with *VviMYB14* to increase STS promoter activity. This observation, together with the presence of the type IIG/AC-element TFBS in close proximity ( $< \pm 50$  bp) with two WRKY binding sites within the *VvSTS29* promoter region supports the hypothesis of a protein-protein interaction between the MYB and WRKY TFs. Although a direct interaction between these two TF proteins is unlikely, it could be mediated by other “bridge” proteins such as WD40s and bHLHs as already observed in *Petunia* (Verweij et al. 2016). Validation of this hypothesis will require yeast-2-hybrid (Y2H) assays to investigate interactions between *VviWRKY3* and *VvMYB14* and to screen *prey* libraries with the aim of identifying potential “bridge” proteins.

In addition to the new insights into the regulatory roles of *VviWRKY24* and *VviWRKY3* in the regulation of the STS pathway, this study has also identified a large collection of other candidate TFs for future gene characterization studies. Validation of these candidates will require a combination of many different approaches including expression profiling experiments associated with chromatin immunoprecipitation, yeast-2-hybrid

assays to investigate protein-protein interactions and *in planta* functional assays to validate roles of these regulators in the grapevine stilbene pathway.

## **Materials and methods**

### **Compilation of transcriptome datasets and gene expression analysis**

Two separate transcriptome compendia were constructed: one based on microarray datasets (29K NimbleGen Grape Whole-genome platform) and another one with next-generation sequencing (RNA-Seq) datasets. Details regarding each dataset are available in Supplementary Table S1. For compiled microarray datasets, raw intensity data were summarized with *oligo* (Carvalho et al. 2010) using the robust multi-array average method in R (<http://www.r-project.org>). The final microarray dataset consists of RMA-normalized values across 356 conditions with biological replicates being averaged when present. For RNA-Seq datasets, raw paired-end or single-end fastq reads were first trimmed and quality filtered using Trimmomatic v0.36 (Bolger et al. 2014) with the following parameters; LEADING, TRAILING, SLIDINGWINDOW, MINLEN, and AVGQUAL of 20, 20, 4:20, 40, and 20, respectively. Trimmed reads were then aligned to the 12X v1 grapevine reference genome (Jaillon et al. 2007), count summarized, and FPKM transcript abundance estimated using HISAT2 v2.0.5 (Kim et al. 2015), featureCounts (Liao et al. 2014), and edgeR (Robinson et al. 2009), respectively using default settings. The final RNA-Seq expression dataset consists of expression estimates ( $\log_2 \text{FPKM} + 1$ ) across 236 conditions with biological replicates being averaged when present. Re-analysis of differential gene expression was performed using *limma* (Ritchie et al., 2015) and *DESeq2* (Love et al., 2014) for microarray and RNA-seq datasets, respectively. False discovery rate (FDR) < 0.05 and an absolute  $\log_2 \text{FC} > 0.5$  defines significant differential gene expression between contrasts (treatment/control) evaluated (See Supplementary table S1).

### **Gene co-expression network construction and statistical significance of reciprocal ranks**

Construction of a mutual rank (MR; Obayashi et al. 2009) gene co-expression network (GCN) for the microarray and RNA-Seq transcriptome compendia was performed separately as previously reported (Wong et al. 2017 in R. The MR score for any two genes (i.e. gene A and gene B), is determined according to:  $\text{MR}_{(AB)} = \sqrt{(\text{Rank}_{(A \rightarrow B)} \times \text{Rank}_{(B \rightarrow A)})}$ , whereby  $\text{Rank}_{(A \rightarrow B)}$  corresponds to the assigned ranking of gene B in a PCC -ordered (descending) list of gene A co-expressed genes, and *vice versa* for  $\text{Rank}_{(B \rightarrow A)}$ . The final order of each genes' co-expressed genes list is sorted by ascending MR scores, with smaller scores indicating strong and robust co-expression (Obayashi et al. 2009). The optimal size of each STS gene co-expression neighborhood, *k* to be considered for

the construction of the complete *STS* ‘guide’ co-expression modules was determined using two approaches. The first involves the inspection of the observed (complete *STS* GCN) and representative null PCC distribution at various *k* intervals of 100 to 1,000 (stepwise of 100). The null PCC distribution was first obtained by random sampling of 1,000 genes (Vandepoele et al. 2006). This sampling procedure is then repeated 100 times to obtain a representative null distribution. The second involves the establishment of statistical significant MR following the approach of Mutwil et al. (2011) which involves the analysis of MR distribution over 1,000 permutations of the original gene expression dataset. Both analyses were performed separately for respective microarray and RNA-Seq transcriptome compendia. Visualization of network modules was achieved using Cytoscape v3.3 (Shannon et al. 2003). Final aggregation of the modules in the two compendia was performed by merging node (degree) and edge (PCC) attributes to produce a final integrated network. Highly interconnected and modular structures in the integrated *STS*-correlated *TF-TF* subnetwork was identified with *GLay* (Su et al. 2010) implemented in Cytoscape.

#### **Phylogeny, structural and protein analysis of candidate genes**

Multiple sequence alignment (MSA) of 13 candidate WRKY genes, including *VviWRKY03* (VIT\_01s0010g03930), *VviWRKY24* (VIT\_08s0058g00690), *VviWRKY43* (VIT\_14s0068g01770) and *VviWRKY53* (VIT\_17s0000g05810) and the other forty-two WRKY TFs already characterized in other plant species, was inferred by using MUSCLE (<http://www.ebi.ac.uk/Tools/msa/muscle/>). Phylogenetic analyses were performed with MEGA software using the Neighbor-joining (NJ) algorithm and 1000 bootstrap iterations. The accessions of WRKY proteins considered in the analysis are as follows: *AaWRKY01* (FJ390842), *AtWRKY01* (NP\_178565.1), *AtWRKY12* (AF404857), *AtWRKY16* (NM\_180802.2), *AtWRKY18* (NM\_119329.4), *AtWRKY22* (NM\_116355.3), *AtWRKY23* (AY052647), *AtWRKY29* (NM\_118486.6), *AtWRKY33* (AK226301), *AtWRKY40* (NM\_106732.4), *AtWRKY44* (NM\_129282), *AtWRKY46* (NM\_130204.3), *AtWRKY50* (NM\_122518.3), *AtWRKY51* (NM\_125877.4), *AtWRKY52* (NM\_001344604.1), *AtWRKY60* (NM\_001335968.1), *CjWRKY01* (AB267401), *CrWRKY01* (HQ646368), *GaWRKY01* (AY507929), *HbWRKY01* (JF742559), *HbWRKY41* (GU372969), *MtSTP* (HM622066), *MtWRKY100577* (EU526033), *MtWRKY100630* (EU526034), *MtWRKY108715* (EU526035), *MtWRKY109669* (EU526036), *NbWRKY08* (AB445392), *OsWRKY13* (EF143611), *OsWRKY45* (AK066255), *OsWRKY53* (AB190436), *OsWRKY74* (XP\_015651250.1), *OsWRKY76* (AK068337), *OsWRKY89* (AY781112), *PgWRKY01* (KR060074), *PqWRKY01* (JF508376), *PsWRKY01* (JQ775582), *PtrWRKY73* (Potri.013G153400.1), *SlWRKY73* (NM\_001247873), *SpWRKY01* (AK320342), *TcWRKY01* (JQ250831), *VviWRKY01* (AY585679), *VviWRKY02* (AY596466). Length of protein sequences,

molecular weight (MW), theoretical isoelectric point (pI), protein instability index (II), aliphatic index (AI), and grand average of hydropathicity (GRAVY) of *VviWRKY03*, -24, -43, and -53 were calculated using ProtParam Expasy tool (<http://web.expasy.org/protparam>; Gasteiger et al. 2005).

### **Mechanical wounding and UV-C stress treatments**

Leaf discs (10 mm diameter) were punched from healthy leaves detached from glasshouse-grown *V. vinifera* cv. ‘Pinot Noir’ vines. Discs were pooled from leaves of the same stage of development, based on leaf size and nodal position, collected from a minimum number of three different potted vines. The punching of discs was considered as a wounding treatment *per se*. Five discs randomly chosen from the pool were sampled at 0, 1, 3, 6, 24 and 48 h after wounding, incubated upside down on moist 3MM filter paper in large Petri dishes in the dark at 22 °C until harvest, immediately frozen in liquid nitrogen and stored at -80 °C until RNA extraction. Control discs (0 h) were collected from an unwounded leaf immediately following detaching from a healthy vine. The UV-C treatment was carried out as described previously (Vannozzi et al. 2012), with discs irradiated with a 254 nm light source (0.36J cm<sup>-3</sup>) at a distance of 10 cm for 10 min. Efficiency of the elicitation treatments under different experimental conditions was determined histochemically by evaluating the intensity of auto-fluorescence of discs mounted with the underside up in a lactic acid, glycerol and water mixture (1:1:1, v/v/v) on glass slides under long-wave UV light (365 nm). The intensity of the observed blue fluorescence was correlated with the quantity of resveratrol quantified in samples. Control discs (not elicited) were exposed to normal light conditions. After treatment, all samples were incubated in the dark at 22 °C. Five discs were randomly chosen from control and UV-C treatments at 0, 1, 3, 6, and 24 h, immediately frozen in liquid nitrogen, and stored at -80 °C until RNA extraction for expression analyses.

### **Quantitative RT-PCR expression analysis of Group-B STSs, CHSs, MYB14/15 and selected WRKY genes in grapevine**

Expression analyses were carried out by quantitative RT-PCR using the “Fast SYBR® Green Master Mix” and the StepOne™ Plus Real-Time PCR System (Applied Biosystems). The samples were analyzed in three technical replicates. Each 10 µl reaction contained 5 µl SYBR Green Master Mix, 0.6 µl of each primer, 1 µl cDNA and 2.8 µl H<sub>2</sub>O. The thermal cycling conditions used were 95 °C for 10 min followed by 40 cycles of: 95 °C for 15 s, 60 °C for 1 min., and 72 °C for 30 s, followed by a melt cycle with 1 °C increments from 55 to 96 °C. The selection of reference genes to normalize the cDNA represents a critical step in any quantitative RT-

PCR analysis. After testing its suitability, elongation factor EF1 (GenBank Accession no. AF176496) was selected for normalization of all samples analyzed. The expression of each target gene was calculated relative to the expression of elongation factor in each cDNA using StepOne™ Software version 2.1 (Applied Biosystems) to calculate normalized expression values (Yuan et al. 2006), observe melt profiles, extrapolate the concentration and measure primer pairs efficiencies. All oligonucleotide primer sequences are listed in Supplementary Table S5.

#### ***in silico* cis-regulatory element screening of *VvSTS29* promoter**

The cloned *VvSTS29* promoter fragment (1.2 Kb fragment upstream of TSS) was scanned for the main R2R3-MYB (e.g. type I, CNGTTR; type II, TNGTTR; and type IIG/AC, CCWAMC) (Prouse and Campbell 2012 and WRKY (i.e. core W-box, TTGACY) TF binding sites (Eulgem et al. 2000), using regular expression functions in R for exact pattern match (with no mismatch allowed) along both + and – strands.

#### **Cloning of *VviWRKY03*, *VviWRKY43*, and *VvWRKY53* for dual reporter luciferase assays**

The complete coding sequences of *VviWRKY3*, *VviWRKY43* and *VviWRKY53* were amplified from cv. ‘Pinot Noir’ cDNA obtained from UV-C irradiated grapevine leaves using proofreading Taq polymerase. Sequence specific primers (Supplementary Table S5) designed to the putative 5’ and 3’ UTRs of target genes were designed using Geneious R8 software on grapevine sequences downloaded from the grape Genome browser (<http://genomes.cribi.unipd.it/grape>). The generated PCR fragments were purified from agarose gels, cloned directly in pENTR/D TOPO gateway vector (ThermoFisher Scientific) and transferred into GW-pART7 vector to produce pART7-*VvWRKY3*, pART7-*VvWRKY43* and pART7-*VvWRKY53* constructs, where WRKY factors were under control of a 35S promoter. The vector pART7<sup>30</sup> was previously modified into a gateway compatible destination vector by using the Gateway® Vector Conversion System (ThermoFisher Scientific®, according to the manufacture protocol; Poschet G., unpublished data). Cloning of pART7-*VviWRKY24* (former *VviWRKY33*) was described by Merz et al. (2015) Reporter constructs carrying firefly luciferase and the promoter sequences of *VvSTS29* gene, together with pART-MYB14 and pART-MYB15 constructs were previously described in Höll et al. (2013).

#### **Transient transfection experiments and dual luciferase assay**

Transient promoter-reporter gene assays were performed using cell suspension cultures obtained from *V. vinifera* cv Chardonnay and Pinot noir petiole callus culture as previously described (Bogs et al. 2007, Walker et al.

2007). The Dual Luciferase assay was modified according to Czernmel et al. (2009). The Renilla luciferase pRLuc was utilized as an internal control in each transfection experiment (Horstmann et al. 2004). Transfection experiments were conducted using single or combined effectors. All transfection experiments were repeated 3 to 5 times, with three technical replicates per experiment. Promoter activity was measured as a fold change compared to control. Mean values of firefly and *Renilla* luciferase ratios are reported as relative luciferase activity with error bars indicating SE.

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#### **Disclosure**

The authors have no conflicts of interest to declare.

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851      **Table 1.** List of all the TFs co-expressed with *VvSTS* genes (degree  $\geq 5$ ) represented in the GCN.

ID	Vitis Name	Family/subfamily	Degree	Function	Reference
<i>Shared interaction module</i>					
VIT_07s0005g03340	<i>VviMYB14</i>	R2R3-MYB / S2	74	STS regulation	Höll et al., 2013
VIT_08s0007g08750	<i>VviHsfB3a</i>	HSF / A	51	unknown	Hu et al., 2016
VIT_01s0010g03930	<i>VvWRKY3</i>	WRKY / IIc	41	unknown	Wang et al., 2014
VIT_01s0026g01730	<i>VvWRKY2</i>	WRKY / IIb	39	unknown	Wang et al., 2014
VIT_11s0016g02070	-	bHLH	38	unknown	
VIT_14s0068g01770	<i>VvWRKY43</i>	WRKY / IIc	36	unknown	Wang et al., 2014
VIT_08s0058g00690	<i>VvWRKY24</i>	WRKY / I	29	unknown	Wang et al., 2014
VIT_06s0004g00020	<i>VvNAC44</i>	NAC / S3	32	unknown	Wang et al., 2013
VIT_07s0005g03220	<i>VvERF098</i>	ERF/AP2 / IX	29	unknown	Licausi et al., 2010
VIT_12s0055g00340	<i>VvWRKY39</i>	WRKY / IIb	22	unknown	Wang et al., 2014
VIT_10s0116g01200	<i>VvWRKY29</i>	WRKY / IIb	14	unknown	Wang et al., 2014
VIT_00s1352g00010	<i>VviMYB148</i>	R2R3-MYB / S14	11	unknown	Wong et al., 2016
VIT_19s0027g00860	<i>VvNAC31</i>	NAC / S6	7	unknown	Wang et al., 2014
VIT_19s0085g00050	<i>VviMYB139</i>	R2R3-MYB / S3	5	unknown	Wong et al., 2016
<i>RNaseq interaction module</i>					
VIT_10s0003g01600	<i>VvWRKY30</i>	WRKY / IIe	34	unknown	Wang et al., 2014
VIT_17s0000g05810	<i>VvWRKY53</i>	WRKY / IIb	26	unknown	Wang et al., 2014
VIT_05s0077g00500	<i>VviMYB108A</i>	R2R3-MYB / S20	11	unknown	Wong et al., 2016
VIT_06s0061g00780	<i>C2H2</i>	C2H2	11	unknown	
VIT_09s0002g01190	<i>VviSHR3</i>	GRAS	10	unknown	Grimplet et al., 2016
VIT_15s0021g01600	-	ERF/AP2	10	unknown	Licausi et al., 2010
VIT_06s0004g04990	<i>VviLISCL2</i>	GRAS	7	unknown	Grimplet et al., 2016
VIT_15s0021g01610	-	ERF/AP2	6	unknown	Licausi et al., 2010
VIT_01s0150g00120	<i>VvERF112</i>	ERF/AP2 - X	5	unknown	Licausi et al., 2010
<i>Microarray interaction module</i>					
VIT_18s0072g00260	<i>VvERF114</i>	ERF/AP2 - X	29	unknown	Licausi et al., 2010
VIT_12s0028g00860	<i>VvNAC36</i>	NAC / S6	26	unknown	Wang et al., 2014
VIT_07s0031g01980	<i>VvERF113</i>	ERF/AP2 - X	22	unknown	Licausi et al., 2010
VIT_05s0049g01020	<i>VviMYB15</i>	R2R3-MYB / S2	15	STS regulation	Höll et al., 2013
VIT_04s0069g00970	<i>VvWRKY11</i>	WRKY / IIc	9	unknown	Wang et al., 2014
VIT_17s0000g00200	-	ERF/AP2	9	unknown	Licausi et al., 2010
VIT_02s0025g00420	<i>VvWRKY4</i>	WRKY / IIe	6	unknown	Wang et al., 2014
VIT_05s0049g01010	<i>VviMYB13</i>	R2R3-MYB / S2	5	STS regulation	Wong et al. 2016

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## 855   **Figures**

856   **Figure 1.** Integrated gene co-expression network of grapevine stilbene synthase (*STS*) and transcription factors  
857   (*TFs*), obtained by integrating two subnetworks generated from microarray and RNA-Seq data, respectively.  
858   Each network was acquired by selecting the top200 ranking genes for each *STS* gene and filtering only those  
859   accession encoding for transcription factors based on Plant TFdb. The size of the TF nodes is proportional to the  
860   number of edges (i.e. the number of *STS* a particular TF is correlated with). The thickness of edges is  
861   proportional to the Pearson Correlation Coefficient (PCC). Nodes showing degree (number of edges) lower than  
862   5 were excluded. Different TF families are represented by different colors.

863   **Figure 2.** Integrated *STS*-correlated *TF-TF* gene co-expression network. The TFs prioritized in the *STS*-TF  
864   top200 MR GCN were considered for highlighting their reciprocal relationships in term of co-expression. The  
865   size of the TF nodes is proportional to the number of edges, i.e. the number of *VvSTS* members that particular TF  
866   is correlated with. The thickness of edges is proportional to the Pearson Correlation Coefficient (PCC).

867   **Figure 3.** (A) Phylogenetic relationships of WRKY proteins from grapevine and Arabidopsis. The deduced  
868   proteins were aligned with MUSCLE and phylogenetic analyses were performed with MEGA software using the  
869   Neighbor-joining (NJ) algorithm and 1000 bootstrap iterations. WRKYs identified in the CGN analysis are  
870   indicated in red; (B) Phylogenetic tree obtained by multiple sequence alignment (MSA) of candidate WRKY  
871   genes together with other forty-two WRKY TFs already characterized in grapevine and in other plant species;  
872   (C) Length of protein sequences, molecular weight (MW), theoretical isoelectric point (pI), protein instability  
873   index (II), aliphatic index (AI), and grand average of hydropathicity (GRAVY) of candidate WRKYs were  
874   calculated using Protparam Expasy tool.

875   **Figure 4.** Gene expression heat map of stress responsiveness of the complete grapevine *STS* gene family,  
876   together with *VviMYB13/14/15* and the candidate *WRKY* genes *VviWRKY3*, *VviWRKY24*, *VviWRKY43* and  
877   *VviWRKY53*. A subset of samples representing biotic stress conditions was extrapolated from transcriptome  
878   compendia. Significantly up-regulated and down-regulated genes are indicated with varying intensities. False  
879   discovery rate (FDR) < 0.05 and an absolute log2FC > 0.5 defines significant differential gene expression  
880   between contrasts (treatment/control) evaluated (marked with \*)

881   **Figure 5.** Heat map showing the expression of the whole *STS* gene family and of *VviMYB13/14/15*, *VviWRKY3*,  
882   *VviWRKY24*, *VviWRKY43* and *VviWRKY53* genes in a subset of samples extrapolated from transcriptome  
883   compendia representing abiotic stresses. Significantly up-regulated and down-regulated genes are indicated with

varying intensities. False discovery rate (FDR) < 0.05 and an absolute log<sub>2</sub>FC > 0.5 defines significant differential gene expression between contrasts (treatment versus control, marked with \*).

**Figure 6.** Quantitative RT-PCR analysis of grapevine *STSs*, *R2R3-MYB* and candidate *VvWRKY* transcript accumulation in response to mechanical wounding and UV-C irradiation. Transcript levels were normalized to the expression of elongation factor *EF1-α* and plotted as log<sub>2</sub> (fold change). Fold change for wounded discs was calculated relative to the untreated sample (0 h), whereas fold change for UV-C-treated discs was obtained by calculating the ratio between treated (UV-C) and untreated (i.e., wounded discs) samples at the same time point. The experiment was repeated twice with similar results. Data show the results of one of these experiments. Bars indicate SE of three technical replicates.

**Figure 7.** Transient expression in *V. vinifera* cv Chardonnay suspension cell culture following particle bombardment. Specific promoters linked to a *Firefly* luciferase gene were co-bombarded into cells with pART7-TF constructs. The *Firefly* luciferase activity was normalized with the *Renilla reniformis* luciferase activity, under control of a 35S promoter. For normalization of all data to the background activity of the *STS29* promoter, the empty pART7 vector was co-transformed with the *STS29* promoter::luciferase construct and used as negative control. A) Canonical MYB and WRKY TF binding sites (TFBS) in the cv “Shiraz” *VviSTS29* 1.2 Kb region used for luciferase assays. A total of two, six, and two type I, II, and IIG MYB binding sites respectively and two WRKY TFBS were identified; B) Dual reporter luciferase assays performed with singular MYB or WRKY effectors; C) Dual reporter luciferase assays performed with combined effectors (MYBs + WRKYs). Results were obtained from 3 to 5 independent experiments and mean averaged. The method currently being used to discriminate among the means is Fisher's least significant difference (LSD) procedure.

## Supplementary material

**Supplementary figure S1.** Distribution of Pearson correlation coefficients (PCC) in the observed and random (A) microarray and (B) NGS *STS* gene co-expression network at different top *k* thresholds. Observed PCC distribution in (A) and (B) is represented as purple and red boxplots, respectively. Random PCC distribution in (A) and (B) is represented as green and cyan boxplots, respectively. Outlier PCC values in (A) and (B) are removed for clarity. Distribution of statistically significant mutual ranks in the global (C) microarray (D) NGS gene co-expression network at different significance level intervals ( $0.05 > P\text{-value} > 0.001$ ). Scores are expressed as  $-\log_{10}(P\text{-value})$ .

913 **Supplementary Figure S2.** Quantitative RT-PCR analysis of grapevine *STSs*, *R2R3-MYB* and candidate  
914 *VvWRKY* transcript accumulation in response to mechanical wounding and UV-C irradiation within the first 10 h  
915 after treatment. Transcript levels were normalized to the expression of elongation factor *EFL-a* and plotted as  
916  $\log_2$  (fold change). Fold change for wounded discs was calculated relative to the untreated sample (0 h), whereas  
917 fold change for UV-C-treated discs was obtained by calculating the ratio between treated (UV-C) and untreated  
918 (i.e., wounded discs) samples at the same time point. Data show the results of one of these experiments. Bars  
919 indicate SE of three technical replicates.

920 **Supplementary Table S1.** Table summarizing the metadata used in the GCN construction indicating SRA ID,  
921 publication year, title, references, and number of assays.

922 **Supplementary Table S2.** List of the top 200 MR-ranked genes for each *VvSTS* member in both microarray and  
923 RNA-Seq compendia.

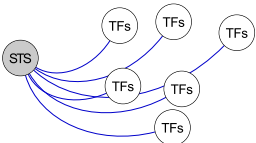
924 **Supplementary Table S3.** List of all the TFs co-expressed with *VvSTS* genes. Accessions highlighted in red  
925 represent TF with degree < 5 not reported in the main text (Table 1, Figure 1, Figure 2).

926 **Supplementary Table S4.** Distribution of canonical MYB and WRKY TF binding sites (TFBS) in the promoter  
927 fragment of *VviSTS29*.

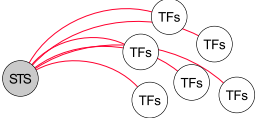
928 **Supplementary Table S5.** List of oligonucleotides used in this study.

929

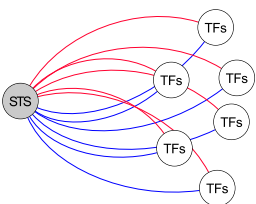
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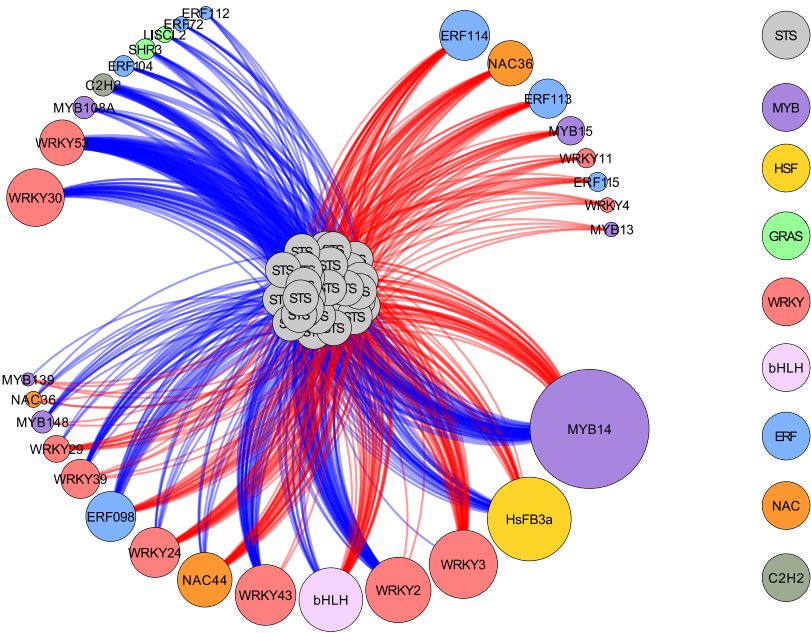
RNA-seq interactions



Microarray interactions



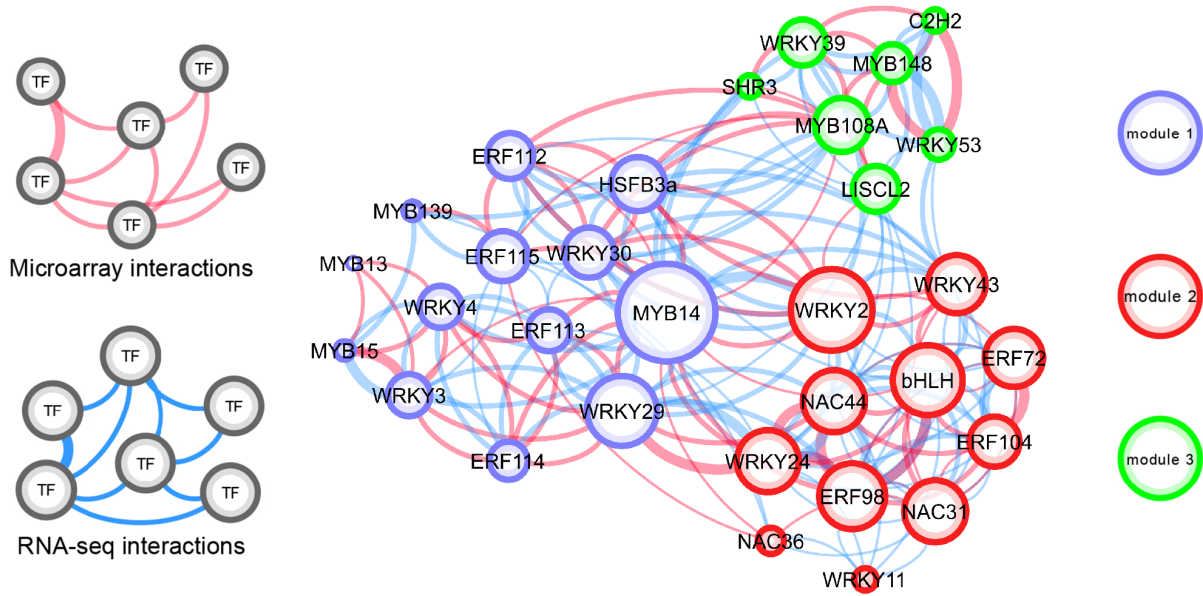
Shared interactions



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