The plant WRKY gene family represents an ancient and complex class of zinc-finger transcription factors (TFs) that are involved in the regulation of various physiological processes, such as development and senescence, and in plant response to many biotic and abiotic stresses. Despite the growing number of studies on the genomic organisation of WRKY gene family in different species, little information is available about this family in grapevine (*Vitis vinifera* L.). In the present study, a total number of 59 putative grapevine WRKY transcription factors (*VvWRKYs*) were identified based on the analysis of various genomic and proteomic grapevine databases. According to their structural and phylogenetic features, the identified grapevine WRKY transcription factors were classified into three main groups. In order to shed light into their regulatory roles in growth and development as well as in response to biotic and abiotic stress in grapevine, the *VvWRKYs* expression profiles were examined in publicly available microarray data. Bioinformatics analysis of these data revealed distinct temporal and spatial expression patterns of *VvWRKYs* in various tissues, organs and developmental stages, as well as in response to biotic and abiotic stresses. To also extend our analysis to situations not covered by the arrays and to validate our results, the expression profiles of selected *VvWRKYs* in response to drought stress, *Erysiphe necator* (powdery mildew) infection, and hormone treatments (salicylic acid and ethylene), were investigated by quantitative real-time reverse transcription PCR (qRT-PCR).

The present study provides a foundation for further comparative genomics and functional studies of this important class of transcriptional regulators in grapevine.

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**INTRODUCTION**

Plants have evolved unique strategies to adapt to and cope with biotic and abiotic stresses due to their inability to escape predation or adverse environmental conditions. Plant response to these stresses is mediated by a complex signal transduction network that results in changes in biochemistry, physiology and morphology.\(^1\)\(^-\)\(^4\) These adaptive responses are associated with distinct changes in gene expression, mediated by the action of transcription factors.\(^5\)\(^-\)\(^7\)

WRKY proteins represent an important class of transcriptional regulators in higher plants and are found throughout the whole green lineage (green algae and land plants).\(^6\) The majority of studies on WRKY transcription factors (TFs) showed that numerous members of this multigene family are involved in the response to biotic stresses and are central components of many aspects of the innate plant immune system. Many loss- and gain-of-function studies in *Arabidopsis thaliana*, which are extensively described in Rushton et al.,\(^8\) demonstrated that these regulatory proteins act in a complex defense response network as both positive and negative regulators of the two partly interconnected branches of plant innate immunity: the microbe/pathogen-triggered immunity and the effector-triggered immunity. Pathogen-triggered immunity and effector-triggered immunity activate both local and systemic defence responses, which are mainly modulated by the signalling pathways related to the jasmonic acid (JA),\(^9\)\(^-\)\(^10\) generally associated to necroptrophic pathogens, chopping insects attack and salicylic acid (SA), more related to defenses triggered by biotrophic pathogenes. JA and SA signalling often act antagonistically, but synergisms between these two phytohormones have also been observed.\(^11\)

In *Arabidopsis*, many WRKY TFs appear to be involved in fine regulation of the balance between SA- and JA-dependent defense pathways. For example, AtWRKY70, a common regulatory component of SA- and JA-dependent defence signalling, mediates the cross-talk between these antagonistic pathways and is a positive regulator of R-gene-mediated resistance and systemic defense responses.\(^12\)\(^,\)\(^13\)

In 2008, Lai et al.,\(^14\) demonstrated that two closely related WRKY TFs, AtWRKY3 and AtWRKY4, play positive role in plant resistance toward necroptrophic pathogens, as Atwrky4, Atwrky3 and Atwrky3 wrky4 mutants showed increased susceptibility toward the fungus *B. cinerea*, whereas the AtWRKY4 overexpression enanched the plant susceptibility to the biotrophic bacterium *Psudomonas syringae*. Many other WRKY TFs were found to act as negative regulators of defence signalling, including AtWRKY7, \(-38\), \(-48\) and \(-62.15\)\(^-\)\(^18\)

The role and complexity of WRKY TFs in plant defense signalling have been demonstrated not only in *Arabidopsis*, but also in other plants, such as rice (*Oryza sativa*),\(^19\)\(^,\)\(^20\) barley (*Hordeum vulgare*)\(^21\) and tobacco (*Nicotiana attenuata*).\(^22\) In rice, a recent study revealed that the pair of allelic genes OsWRKY45-1 and OsWRKY45-2, which encode proteins with a 10-amino acid difference, play opposite roles in rice resistance against bacterial pathogens.\(^19\) Despite both alleles positively regulate resistance to the rice fungal pathogen *Magnaporthe grisea*, they differentially regulate resistance to *Xanthomonas oryzae* since the former appears to be a negative regulator of the defense response, whereas the latter acts as a positive regulator. Again, the opposite roles of the two allelic genes...
seem to be due to their mediation of different JA and SA signalling defense pathways.

Altogether research on the role of WRKY TFs in plant response to abiotic stresses has lagged behind that on biotic stresses, a growing number of evidences are unrevealing the important role of WRKYs in the regulation of abiotic stress-induced pathways. Many examples illustrates that WRKY TFs form part of signalling processes associated with transcriptional reprogramming when plant encounters high salt, heat, osmotic stress, high CO$_2$ levels, high ozone concentrations, cold or drought.\cite{19, 23–25} Moreover, recently, scientists have focussed on additional roles of WRKY factors in other important plant processes, such as seed dormancy and germination,\cite{24, 26, 27, 28, 29} and leaf senescence.

One of the defining features of WRKY TFs is their DNA binding region, also called WRKY domain, that consists of about 60 amino acids,\cite{34, 35} characterized by the highly conserved WRKY signature at the N-terminus adjacent to an atypical, either C$_x$$\Delta$C$_{4–5}$C$_{22–23}$HXH or C$_x$C$_{4–5}$C$_{22–23}$HXC, zinc-finger motif at the C-terminus.\cite{36} WRKY genes have been classified into three major groups based on the number of WRKY domains present.\cite{35} Group I members are characterized by two WRKY domains containing a C$_2$H$_2$ zinc-finger motif. Group II WRKY genes contain only one WRKY domain, characterized by a C$_2$H$_2$ zinc-finger motif. This group has been further divided into five subgroups (II-a, b, c, d and e, respectively). Group III consists of a small number of genes characterized by a single WRKY domain with a C$_2$HC zinc-finger motif. In a few WRKY proteins, the WRKY amino-acid sequences have been replaced by WRKY, WSKY, WKRY, WVKY or WKKY.\cite{36} The conservation of the WRKY domain is mirrored by a remarkable conservation of its cognate binding site, the W box (TTGACC/T).\cite{34, 35} Both bioinformatic-based and functional studies of plant promoters have found clusters of W boxes in stress-inducible promoters.\cite{37} In some cases, the multiple W boxes appear to have a synergistic effect on transcription, as observed in parsley (Petroselinum crispum)\cite{5, 58}

Grapevine (Vitis vinifera L.) is a major economic crop species with world production approaching 70 million tonnes and a harvest area of over seven million hectares. It is cultivated for table grapes, juice, raisins and wine production.\cite{39} Over the course of the last few decades, there has been a great increase in the economic competitiveness of grape production. The competitive challenge is associated with the need to maintain quality and productivity, and the need to adapt production to new geographical areas. Climate changes, the advent of new diseases and increasing market demand have led to a remarkable increase in studies related to the physiology and pathology studies of this species.

Only six individual WRKYs have been characterized in grapevine thus far. Among them, VvWRKY1 appears to be involved in plant response to downy mildew (Plasmopara viticola) infection through a JA-mediated transcriptional reprogramming.\cite{40} VvWRKY2 plays a role in regulating lignification, possibly in response to biotic or abiotic stresses,\cite{41} and VvWRKY11 has an important role in the response to water stress.\cite{42} The other three VvWRKYs, VpWRKY1, VpWRKY2 and VpWRKY3, were cloned from a wild Chinese species, Vitis pseuderocaula W. T. Wang ‘Baie-35-1’, and all of them have been shown to be involved in the resistance to both biotic and abiotic stresses.\cite{43}

The availability of the genome sequences of a highly homozygous clone PN40024 of Pinot Noir\cite{44} and the cultivated Pinot Noir clone ENTAV 115\cite{45} represents a strong foundation for a vast range of genetic and functional studies of entire gene families. In the present study, a total of 59 VvWRKY genes were identified and manually annotated from the grapevine genome. Detailed analysis, including gene classification, annotation, phylogenetic evaluation, conserved motif determination and expression profiling based on previously published microarray data were performed on all members of the family. Additionally, the expression patterns of Group II-a and III WRKY transcription factors (VvWRKYs) in response to abiotic and biotic stresses were further investigated using quantitative real-time reverse transcription PCR (qRT-PCR). Our results provide a foundation for further comparative genomics and functional studies of this important class of transcriptional regulators in grapevine.

**MATERIALS AND METHODS**

Identification of putative WRKY proteins in grapevine

To identify a complete list of grapevine WRKY genes, annotated grapevine proteins were downloaded from three public databases: the National Centre for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/), the Grapevine Genome Browser (http://www.genoscope.cns.fr/externe/GenomeBrowser/Vitis/) and the Grapevine Genome CRIIBi Biotech website (http://genomes.criibi.uniprd.it/). Putative grapevine WRKY proteins were identified as described by Wu et al.\cite{46} The HMM profile of the WRKY DNA-binding domain (PF03106) was downloaded from the Pfam protein families database (http://pfam.janelia.org) and used to survey all grapevine proteins in the 12× coverage assembly of the V. vinifera PN40024 genome. All non-redundant WRKY protein sequences were considered and used in further analysis. Length of sequences, molecular weights and isoelectric points of deduced polypeptides were calculated by using tools provided at the ExPasy website (http://web.expasy.org/protparam/). Finally, manual annotation was performed to resolve any discrepancy between incorrectly predicted genes and the actual chromosomal locations of involved genes in question. The Arabidopsis WRKY gene family database was downloaded from the Database of Arabidopsis Transcription Factorshttp://datf.cbi.pku.edu.cn/) and used for comparative analysis.

Phylogenetic analysis and classification of the grapevine WRKY family

All identified grapevine WRKY genes (VvWRKYs) were classified into different groups based on the AtWRKY classification scheme and the alignment of VvWRKY and AtWRKY DNA-binding domains using Clustal X 2.1\cite{47} with default settings. The phylogenetic trees were inferred using MEGA 5.0\cite{48} with the neighbour-joining method. Bootstrap values were calculated for 1000 iterations.

Chromosomal location of VvWRKYs

All VvWRKY genes were mapped to grapevine chromosomes based on information available at the Grape Genome CRIIBi website (http://genomes.criibi.uniprd.it/). The map was drafted using MapInspect software (http://www.plantbreeding.wur.nl/uk/software-mapinspect.html). Tandem duplicated VvWRKY genes in the grape genome were identified by checking their physical locations on individual chromosomes and were defined as adjacent paralogous on a grape chromosome, with no more than one intervening gene.\cite{49} MCScanX software (http://chibba.pgm.uga.edu/mcscan2/) was used to detect the gene duplication events, with the E-value set below 1×10$^{-5}$. Physical chromosomal locations were graphically represented by scaling the 19 chromosomes.

Analysis and distribution of conserved motifs and exon–intron structures

The exon–intron organisation of VvWRKY genes was determined by comparing predicted coding sequences with their corresponding genomic sequences using the GSDS software (http://gds.cbi.pku.edu.cn). The Fancy Gene online tool (http://bio.ieo.eu/fancygene/) was used to illustrate the exon–intron structure of VvWRKYs.\cite{50} The Expectation Maximization for Motif Elicitation online program (http://meme.sdsc.edu/meme/intro.html)\cite{51} for DNA and protein sequence analysis was used to identify conserved motifs in the 59 VvWRKY-deduced proteins. The optimized parameters of Expectation Maximization for Motif Elicitation were set as follows: the number of repetitions, any; the maximum number of motifs, 50; and the optimum width of each motif, between 6 and 300 residues. The online program 2ZIP (http://2zip.molgen.mpg.de/index.html) was used for predicting Leu zipper motifs, while HARF, LXXLL and LXXLXX motifs were identified by manual inspection.

Microarray data analysis

The expression profiles of VvWRKY genes predicted from an analysis of the grapevine genome releases was analysed in a V. vinifera cv ‘Corvina’ (clone 48) gene expression atlas of different organs at various developmental stages. Microarray data were downloaded from Gene Expression Omnibus (GEO) database (GSE3775).

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**Grapevine WRKY gene family analysis**

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under the series entry GSE36128 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=lfcrxseycqgsjoandacc=GSE36128). Expression analyses in response to abiotic and biotic stresses were based on microarray data (series matrix accession numbers GSE31594, GSE31677, GSE6404, GSE12842 and GSE31660) downloaded from the NCBI gene expression omnibus (GEO) datasets. Data were analysed and graphically represented using the mean of Multi Experiment Viewer software.23 Apart from data obtained from the V. vinifera cv. Corvinus expression atlas, which were normalized based on the mean expression value of each gene in all tissues/organisms analysed, all other expression data were calculated as log, fold change in treated vs. untreated samples.

Plant material and treatments

In vitro grapevine plants (V. vinifera, genotype PN40024) were kindly provided by Dr Anne-Françoise Adam-Blondon (INRA, Castanet Tolosan cedex France) and maintained in vitro on half-strength MS medium supplemented with 0.3 mg L\(^{-1}\) indole 3-butyric acid (Sigma Chemical Co., St. Louis, MO, USA), at 25°C in a culture room under 16 h light/8 h dark photoperiod cycle. To validate VvWRKYs microarray expression data obtained from the Corvinus expression atlas in different tissues and developmental stages, three biological replicates of roots, stems, leaves and shoot tips, were independently gathered in subcultured young shoots. The following program was used: 94°C for 6 s, which was normalized based on the mean expression value of each gene in all tissues/organisms analysed, all other expression data were calculated as log, fold change in treated vs. untreated samples.

To validate VvWRKYs microarray data in response to various stresses, Group I and Group III VvWRKY genes were selected for analysis using qRT-PCR. For drought stress treatment, five-week-old in vitro PN40024 plants were transplanted into pots and acclimatized in a growth incubator until about 40 cm tall with 14 leaves. To validate VvWRKYs microarray expression data obtained from the Corvinus expression atlas in different tissues and developmental stages, three biological replicates of roots, stems, leaves and shoot tips, were independently gathered in subcultured young shoots. The following program was used: 94°C for 6 s, which was normalized based on the mean expression value of each gene in all tissues/organisms analysed, all other expression data were calculated as log, fold change in treated vs. untreated samples.

RESULTS

Identification and annotation of VvWRKY genes in the grapevine genome

A total of 59 full-length genes encoding putative WRKY proteins were identified in the grapevine genome. Among them, 58 gene sequences were obtained from the 12X V1 prediction of the PN40024 grape genome sequence which is available at the CRIBI Biotech website (http://genomes.cribi.unipd.it/). One additional gene, corresponding to accession XP_002277383.1 in the NCBI GenBank database, was found in the old version of the PN40024 genotype representing the 8X coverage of the grape genome (http://www.genoscope.cns.fr/externe/GenomeBrowser/Vitis/). A homologous sequence (CAN70150) was also identified in the PN ENTA V115 grape genome sequence, indicating that this WRKY member is real and was most likely missed in the PN40024 12X V1 prediction of proteins. Three WRKY genes identified as VIT_11s0052g04450, VIT_15s0021g01310 and VIT_06s0040g00230 in the 12X V1 prediction, were found to have truncated coding sequences and manual annotation was performed to correct their sequences. All 59 identified WRKY genes were named as VvWRKY1 to VvWRKY59 based on their chromosome location according to Vannozzi et al.25 The parameters used to describe the VvWRKY proteins are listed in Table 1 and included the deduced protein length, the molecular weight, the isoelectric point, the aliphatic index and the grand average of hydropathicity. The deduced length of VvWRKY proteins ranged from a minimum of 101 amino acids (VvWRKY12) to a maximum of 612 residues (VvWRKY59), whereas the PI ranged from 5.02 (VvWRKY38) to 9.91 (VvWRKY34). This range of variability implies that different VvWRKY proteins might operate in different microenvironments.

Chromosomal distribution and exon–intron organisation of VvWRKY genes

Using Tblastn, 57 out of the 59 VvWRKY genes could be mapped to 18 out of 19 grapevine chromosomes (chr), with the exclusion of chr 3, indicating a comprehensive distribution of VvWRKYs within the grapevine genome (Figure 1). VvWRKY58 and 59, which mapped to chr1_random and chrUn, respectively, are not showed. The para
dologous chromosome segments resulting from whole-genome duplication or fusion events are illustrated in the same colour as in Jaillon et al.43 Chromosome 4, with eight VvWRKY genes, possesses the largest number followed by chr 7 with six genes.

According to the definition of a gene cluster provided by Holub,56 13 VvWRKY genes could be placed in six clusters. Among the six clusters, two were found on chromosomes 4 and 7. While only one cluster was located on chromosomes 13 and 15. Tandem and segmental duplications have been suggested to be two of the main causes for gene family expansion in plants.57 We identified two WRKY tandem duplication clusters (VvWRKY11/VvWRKY12 and VvWRKY40/VvWRKY41) on grapevine chromosome 4 (Figure 1). We then examined the synteny block within the grape genome and found that 16 VvWRKY genes were located in 11 pairs of duplicated genome regions (Figure 1). In summary, 20 of 59 VvWRKY genes were associated in either tandem and segmental duplication events.

The exon–intron structure of all the identified VvWRKY genes was analysed in order to gain more insight into the evolution of the WRKY family in grapevine. As results indicated that a large number VvWRKY genes contain two to four introns (Figure 2), twenty-seven VvWRKY genes were found to possess two introns, eight members
### Table 1. Grapevine WRKY genes identified in the PN40024

| Proposed name | PN40024 12× V1 ID | Chr | ORF (aa) | MW (kDa) | pl | Al | GRAVY | Group | Conserved motif | Domain pattern | Zinc finger |
|---------------|-------------------|-----|----------|----------|----|----|-------|-------|----------------|--------------|------------|
| VVWRKY01      | VIT_01s0011g00720 | 1   | 305      | 33.3585  | 5.62| 47.67| −0.882| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY02      | VIT_01s0026g01730 | 1   | 594      | 64.3902  | 6.02| 57.24| −0.689| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY03      | VIT_01s0010g03930 | 1   | 189      | 21.2588  | 9.13| 54.13| −0.723| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY04      | VIT_02s0025g00420 | 2   | 329      | 36.3666  | 4.71| 40.62| −0.602| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY05      | VIT_02s0025g01280 | 2   | 342      | 38.5618  | 5.05| 55.03| −0.793| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY06      | VIT_04s0008g01470 | 4   | 166      | 18.1917  | 5.2 | 48.19| −1.122| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY07      | VIT_04s0008g00570 | 4   | 261      | 28.8756  | 9.1 | 65.4 | −0.716| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY08      | VIT_04s0008g00576 | 4   | 317      | 35.2637  | 8.22| 73.19| −0.718| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |
| VVWRKY09      | VIT_04s0008g06600 | 4   | 491      | 54.3129  | 8.88| 57.8 | −0.837| Ile   | WRKYQGK       | C-X-C-X-H-X-H-H | C2H2       |

Continued
had three introns and eleven had four introns. Five VvWRKY genes had five introns, one had six introns and seven VvWRKY genes had only one intron. The intron located in the conserved WRKY domain can be classified as either an R-type intron or V-type intron, similar to what has been reported in rice.46 The R-type intron, a phase-2 intron, is spliced prior to an arginine residue (R). However, the V-type intron, a phase-0 intron, is spliced before a valine residue, six amino acids after the second cysteine residue of the zinc finger C2H2 motif46 (Supplementary Fig. S1). The V-type intron was only observed in the WRKY domains of genes belonging to groups II-a and II-b, whereas the R-type intron is extensively distributed in all the other VvWRKY groups (I-C, II-c, II-d, II-e and III) (Supplementary Fig. S1). No introns were found in the sequence encoding the Group I N-terminal WRKY domains.

Phylogenetic analysis, classification and motifs of the grapevine WRKY gene family

In order to examine the phylogenetic relationship between the predicted WRKY DNA-binding domains in grapevine and Arabidopsis, an unrooted phylogenetic tree was inferred based on the alignment of 131 WRKY domains from the two species (excluding N-terminal domains of Group I) (Figure 3). Based on AtWRKY classification, VvWRKYs were classified into three major groups. Group I is comprised of 12 proteins, each containing two WRKY domains and the C2H2-type zinc-finger motifs (C–X4–C–X22–H–X1–H). The only exception within this group is VvWRKY57, which does not possess any zinc-finger motifs at the C-terminus. Thirty-nine VvWRKY proteins possessing a single WRKY domain were assigned to Group II in which the C2H2-type zinc-finger structure is C–X5–C–X23–H–X1–H. Group II was further divided into five subgroups based on their primary amino-acid sequence and was comprised of Groups II-a, -b, -c, -d and -e with 3, 8, 15, 6 and 7 members, respectively. Finally, six VvWRKYs, each with a single WRKY domain, were assigned to Group III, differing from Group II VvWRKYs based on their C2HC zinc-finger structure (C–X4–C–X23–H–X1–C). VvWRKY15 and 27 exhibit sequence divergence in the unique WRKY domain; therefore, they were not classified into any group. The grouping of VvWRKY proteins produced the following pairwise relationships between grapevine and Arabidopsis: VvWRKY24 and...
Figure 2. Structures of VvWRKY genes. Exon–intron composition of VvWRKY genes. Names of genes are indicated on the left. Exons, represented by black or red boxes, were drawn to scale. Dashed lines connecting two exons represent an intron. Intron phases 0, 1 and 2 are indicated by numbers 0, 1 and 2, respectively. WRKY domains in VvWRKY proteins are marked in red.
AtWRKY33 in Group I; VvWRKY22 and AtWRKY51 in Group II-c; and VvWRKY49 and AtWRKY22 in Group II-e. A strong relationship among some of the grapevine WRKY proteins may imply the occurrence of gene duplication. For example, VvWRKY11 and 12 belong to the same cluster, and VvWRKY44, 53 and 02 were located within a synteny block (Figure 1). A consensus phylogenetic tree obtained by aligning all the deduced grapevine WRKY domains (including both N-terminal and C-terminal domains) is shown in Figure 4, while the details of the alignment of domains are reported in Supplementary Figure S1. Out of a total number of 71 WRKY domains, 54 WRKY proteins contain WRKYGQK domains that were perfectly conserved, while the other domains differ in one amino acid in the conserved WRKY signature. The WRKY domain sequence in VvWRKY6, 11, 12 and 22 all belong to the Group II-c. While the WRKY domain sequence in VvWRKY15, which does not fall into one of the three WRKY groups, is WKKYGQK.

Conserved motifs other than the WRKY domain were detected manually in the VvWRKY proteins (Supplementary Table S1). A ‘Leucine zipper’ motif, present in Group II-a and II-b WRKY proteins of Arabidopsis and rice, was detected in VvWRKY07 belonging to Group II-a. Four members (VvWRKY10, 21, 34 and 54) of Group II-d contain the novel HARF (RTGHARFRR/A/G/P) motif, whose function has not been clearly determined. The LxLxLx repressor and the LxxLL co-activator motifs were found in seven (VvWRKY08, 14, 21, 34, 41, 48 and 54) and ten (VvWRKY02, 07, 08, 15, 25, 34, 39, 44, 53 and 55) VvWRKY proteins, respectively. Interestingly, VvWRKY08 contains both co-activator and active repressor motifs.

Twenty other distinct motifs (Table 2 and Figure 4) were identified by using the Multiple Expectation Maximization for Motif Elicitation online tool (http://meme.sdsc.edu/meme/intro.html). As illustrated in Figure 4, other than motifs 1 and 3 which are the WRKY domains widely distributed in all 59 proteins, VvWRKY members within the same subgroups were generally found to share a common motif composition, as illustrated in Figure 4. For instance, motif 14 is unique to Group III, whereas motifs 13 and 19 are specific to Group II-d. Group II-a and II-b both contain motifs 7 and 8 except for VvWRKY44 which possesses only motifs 8. As observed for the amino-acid motifs detected in VvWRKY deduced proteins, WRKY genes within the same group generally also possess similar exon–intron arrangements.
Figure 4. Phylogenetic tree of deduced VvWRKY domains associated with the motif compositions in the amino-acid sequences. Consensus phylogenetic tree generated after alignment of deduced VvWRKY domains (both N-terminal and C-terminal domains) with ClustalX 2.1 using the NJ method. The phylogenetic trees were constructed using MEGA 5 software. Reliability of the predicted tree was tested using bootstrapping with 1000 replicates. Numbers at the nodes indicate how often the group to the right appeared among bootstrap replicates. Subtrees branch lines are coloured indicating different WRKY subgroups. Group I N-terminal WRKY domains are indicated with the suffix N after the name, whereas Group I C-terminal WRKY domains are indicated with the suffix C. The motif composition related to each VvWRKY protein is displayed on the right-hand side. The motifs, numbered 1–20, are displayed in different coloured boxes. The sequence information for each motif is provided in Table 2. NJ, neighbour-joining.
Table 2. Analysis and distribution of conserved motifs in grapevine VvWRKYSs

| Motif | E-value | Width | Best possible match |
|-------|---------|-------|---------------------|
| 1     | 1.6×10⁻⁹⁴⁴ | 26   | DGYRWRKYQGVKQGQP|FNIPRSYRCT |
| 2     | 3.8×10⁻⁹⁵⁴ | 41   | (SAAGCCP|VA|RK|V|KR|QHVR|V|EQ|IR|ASC|IF|SA|EDPS|IM|VL|IV|TV|YEG|EH|NH|PD|LPARAx|ASxAS |
| 3     | 1.4×10⁻³⁵⁹ | 21   | (YFW|SA|WRKYGGKQPK|KLGS|PK|YY|PR|GS|YY |
| 4     | 1.3×10⁻³⁵⁹ | 21   | E|KR|TK|IV|ER|EK|PR|VF|AV|FQ|RT|KR|SEYDLD |
| 5     | 3.1×10⁻¹⁰⁵ | 26   | GQ|IV|TE|IV|Y|KGTH|ND|HP|K|QPN|RK|RS|AKS|LG |
| 6     | 1.0×10⁻⁸⁶ | 34   | SG|R|CHSK|KR|KL|RL|VK|SR|TV|IV|V|RK|V|PIAIS|NS|K|M|AD|PI|PA|ED |
| 7     | 2.5×10⁻⁸⁶ | 41   | (EK|ILT|V|EG|V|VGS|L|LAI|Q|KKV|AE|V|EL|M|EN|E|V|RT|ES|M|ENS|Q|I|M|MV|TCM|KNE|ND|I|NS|ADT|LQ|M|HQ|L|FV|IV|ET|L|LYM |

The number of VvWRKY genes belonging to each subgroup was compared with the number in other plant species where this gene family has been fully characterized, i.e., Arabidopsis, rice, castor bean, poplar and tomato (Table 3). The number of genes in Group I WRKY family is similar in grapevine, Arabidopsis, rice and tomato, but has undergone a significant expansion in poplar. The number of members belonging to the different Group II subclades are also comparable between all the considered species, although an obvious expansion was detected in poplar Group II-d and tomato Group II-e WRKYSs. Lastly, apart from Castor bean, grapevine showed a lower number of members belonging to Group III if compared to the other species, especially rice, which has the largest number of WRKY proteins in this subfamily.

VvWRKYs expression in different organs, tissues and developmental stages

The expression patterns of VvWRKY genes were analysed in the V. vinifera cv. Corvina global gene expression atlas, which consists of 45 different organs/tissues at various developmental stages obtained by microarray analysis (Supplementary Table S2). All VvWRKYs, except VvWRKY36, had corresponding probes on the NimbleGen array. Figure 5a presents a graphical representation of the expression pattern of each VvWRKY gene in which both genes and samples were ordered based on a hierarchical clustering analysis.

This analysis indicates that only some members within the same group share a similar expression profile in grapevine organs/tissues during development (Figure 5a). For example, VvWRKY14, 19 and 52, belonging to Group II-c, are highly expressed in berries and rachis during or following véraison (V, MR and R). Other members of this group, such as VvWRKY23 and 46, exhibit high levels of expression in overwinter organs tissues, such as buds (Bud-W) and stems (Stem-W), suggesting an involvement of Group II-c members in berry ripening and cold acclimation.

Members of Group III VvWRKYs, including VvWRKY05, 47, 41 and 25, and Group II-a, including VvWRKY07, 08 and 28, are characterized by high levels of expression in a heterogeneous group of tissues including inflorescence, leaf, rachis, tendril and berry in pre-véraison phases and are less expressed (especially VvWRKY05 and 47) in berries and the rachis after véraison phase. The expression pattern

Table 3. Size of the WRKY groups and sub-groups in different plant species

|       | Group I |       |       |       |       | Group III | NG | Total |
|-------|---------|-------|-------|-------|-------|-----------|----|-------|
|       |         |       |       |       |       |           |    |       |
| Grapevine | 12 | 3 | 8 | 15 | 6 | 7 | 6 | 2 | 59 |
| Arabidopsis | 16 | 3 | 8 | 17 | 7 | 8 | 14 | 1 | 74 |
| Rice | 15 | 4 | 8 | 15 | 7 | 11 | 11 | 5 | 96 |
| Castor bean | 9 | 3 | 10 | 12 | 5 | 15 | 10 | 4 | 47 |
| Poplar | 50 | 5 | 9 | 13 | 13 | 4 | 10 | 3 | 104 |
| Tomato | 15 | 5 | 8 | 16 | 6 | 17 | 11 | 3 | 81 |

*a* According to Ulker and Somssich.6

*b* According to Xie et al.36

*c* According to Li et al.59

*d* According to He et al.65

*e* According to Huang et al.66

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Figure 5. Expression profiles of the grapevine VvWRKY genes in different grapevine organs, tissues and developmental stages. (a) Expression of VvWRKY genes in the V. vinifera cv ‘Corvina’ atlas (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=lfcrxeyciggsjvandacc=GSE36128). Data were normalized based on the mean expression value of each gene in all tissues analysed. Genes were hierarchically clustered based on average Pearson’s distance metric and ‘average linkage’ method. Yellow and blue boxes indicate high and low expression levels, respectively, for each gene. (b) qRT-PCR validation of VvWRKY expression in roots, stems, leaves and shoot tips obtained from 4-week-old in vitro explants of the V. vinifera PN40024 genotype. Transcripts were normalized to the expression of the actin gene. The mean ± s.d. of three biological replicates are presented. Asterisks indicate that the corresponding gene was significantly up- or downregulated in a given organ (*p < 0.05, **p < 0.01). Bud-AB, bud after burst; Bud-B, Bud burst; Bud-W, winter bud; Bud-L, latent bud; Bud-S, bud swell; Flower-F, flowering; Flower-FB, flowering begins; FS, fruit set; Inflorescence-Y, young inflorescence with single flowers separated; Inflorescence-WD, well-developed inflorescence; Leaf-FS, mature leaf; Leaf-S, senescing leaf; Leaf-Y, young leaf; MR, mid-ripening; R, ripening; PFS, post fruit set; Stem-G, green stem; Stem-W, woody stem; Tendril-FS, mature tendril; Tendril-WD, well-developed tendril; Tendril-Y, young tendril; V, véraison.
of these genes was also evaluated by qRT-PCR in selected organs of the V. vinifera PN40024 genotype, confirming their high level of expression in leaves (Figure 5b). Low modulation in the levels of expression were detected for VvWRKY40 and 51 (Group III) in all tissues in the Corvina atlas. However, results on expression levels obtained by qRT-PCR revealed high transcript accumulation in leaves (Figure 5b). Another interesting gene cluster in Group I, composed of VvWRKY13, 59 and 57, was all characterized by a high level of expression that was restricted to stamen and pollen tissues. A high level of modulation was observed in some VvWRKYs in senescing leaves (S). These include VvWRKY22 (Group II-c), VvWRKY 02 (Group II-b) and VvWRKY03 (Group II-c). Although the upregulation of VvWRKY22 and 02 was also present in previous leaf developmental stages (Y and F5), high levels of expression of VvWRKY03 were restricted to the senescing stage of leaves. VvWRKY17 (Group II-d) and VvWRKY27 (NG) were induced in the very early phases of seed development, especially at post-fruit-set, while VvWRKY44, 48 and 53 were upregulated in roots.

Expression patterns of VvWRKY genes in response to biotic and abiotic stresses

The expression of VvWRKY genes in response to both biotic and abiotic stresses was investigated using microarray data from several previously published papers. Regarding abiotic stresses, expression datasets were obtained from two studies (GSE31594 and GSE31677) conducted on transcriptomic response of leaves of V. vinifera cv ‘Cabernet Sauvignon’ to short-term salt, water and cold stress, and long-term water and salt stress. Since the Affymetrix array used in these analysis was based on the few VvWRKYs sequences (cDNA and ESTs) known at the time, the expression of only a limited number of VvWRKY genes could be determined (Supplementary Tables S3 and S4). In general, cold stress induced the most rapid upregulation of VvWRKY genes of all the abiotic stress treatments (Figure 6a). In fact, a relatively large group of VvWRKY genes, including VvWRKY08, 14, 16 and 24, was induced since the first hour after the imposition of the stress, whereas the majority of genes induced by salt (SS) and water (WS) stresses, were upregulated after 24 h. Anyway, the quicker transcriptional response observed upon cold treatment could be related to a more rapid perception of the cold compared to the gradual imposition of the water stress. More in detail, VvWRKY24 was found to be a cold-specific responsive gene since it was induced at all time points of the cold treatment, but was not in response to either the water or salt stresses. Conversely, VvWRKY30 and 52 were both strongly induced 24 h after water and salt stress, but not in response to cold stress. VvWRKY29 appeared to be weakly induced since in the first hours following salt and cold treatments, but interestingly was strongly repressed 1 h after the water stress treatment. The data presented in Figure 6b also confirmed the putative involvement of VvWRKY30 and 52 in water and salt stress responses together other genes such as VvWRKY03 and 16. Interestingly, all the VvWRKY genes that were induced within 24 h in response to short-term water and salt stress (Figure 6a) appeared to be more highly induced in the last point of the long-term stress treatments (16 days after treatment, Figure 6b). VvWRKY29 was again downregulation in almost all time points of the water stress treatment, whereas it appeared to be slightly induced in response to salt stress (Figure 6b).

In order to validate the results obtained from the analysis of the microarray data, qRT-PCR was performed on PN40024 leaves infected by E. necator in order to validate and integrate microarray data (Figure 7d). In general, the expression profiles of VvWRKY genes in response of grapevine to GLRaV-3 infection in both véraison and ripening phases revealed a general repression of many members of the VvWRKY family (Figure 7c). However, a weak-upregulation, limited to VvWRKY30, 35 and 29, specifically induced during the ripening phase.

DISCUSSION

Organisation of the grapevine VvWRKY gene family

WRKY genes encode a class of transcription factors that are ubiquitous to all plant species and have been surveyed in several species whose genomes have been sequenced, such as Arabidopsis,\textsuperscript{35} rice,\textsuperscript{36} cucumber,\textsuperscript{37} poplar,\textsuperscript{38} tomato,\textsuperscript{49} sunflower\textsuperscript{77} and coffee.\textsuperscript{68} In the current study, a search for WRKY genes in the last and previous versions of the genome assembly of the grape PN40024 genotype resulted in the identification of 59 members, which were designated VvWRKY01 through VvWRKY59 on the basis of their chromosomal location.

The structure of the phylogenetic tree obtained from an alignment of the deduced WRKY domains of grapevine and Arabidopsis indicated that the 59 VvWRKY genes identified in grapevine can be

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divided into the three major groups (I, II and III) as previously described in other plant species. Members within the same group, or subgroup within group II, shared a similar gene structure (intron/exon organisation), length and amino-acid motif composition, indicating their close evolutionary relationship.

Group I VvWRKYs, which are characterized by the presence of two WRKY domains, contain approximately 20% of the entire VvWRKY family and is comparable in size to Group I in Arabidopsis, rice, tomato and castor bean (Table 3). This group has undergone a significant expansion in poplar, where it contains approximately...
50% of the entire PtWRKY family. Group II subgroups in Arabidopsis, rice and grapevine were comparable in size, whereas a sensible expansion was detected for poplar Group II-d and tomato group II-e. In grapevine, the members of Group II-a and II-b are characterized by the presence of motifs 7 and 8, whereas Group II-d contains motifs 13 and 19 and the HARF sequence. WRKY Group III, containing the motif 14, was found to be smaller in number than in Arabidopsis or rice, with only 10% of the entire VvWRKY family falling in this clade. Similar percentages were detected for castor bean, poplar and tomato. Gene orthology may represent a starting point for genetical studies which aim to define the function of a candidate gene. Some of the predicted WRKY domains from

Figure 7. Expression profiles of VvWRKY genes in response to biotic stresses. (a) V. vinifera cv ‘Cabernet sauvignon’ and V. aestivalis cv ‘Norton’ plants were grown in an environmental chamber and inoculated with Erysiphe necator conidiospores (PM). Inoculated leaves were harvested at 0, 4, 8, 12, 24 and 48 h after inoculation (GEO series GSE6404). (b) Field-grown plants of V. vinifera cv ‘Chardonnay’ and ‘Incrocio Manzoni’ naturally infected with Bois Noir phytoplasma (BN), compared to healthy samples (GEO series GSE12842). (c) V. vinifera cv ‘Cabernet Sauvignon’ was infected with GLRaV-3 during véraison and the ripening stages of berry development (GEO series GSE31660). (d) qRT-PCR expression analysis of VvWRKY genes in Pinot PN40024 genotype infected with powdery mildew. Transcripts were normalized to actin gene expression. The means ± s.d. of three biological replicates are presented. Asterisks indicate that the gene was significantly up- or downregulated in a given organ (*p ≤ 0.05, **p ≤ 0.01). GLRaV-3, grapevine leaf roll-associated virus-3.
grapevine and Arabidopsis showed pairwise relationships, such as the those between VvWRKY22 and AtWRKY51, VvWRKY24 and AtWRKY33, and VvWRKY49 and AtWRKY22. Indeed, the function of these Arabidopsis genes was previously inferred. AtWRKY51 proteins appear to be involved in the SA-dependent and low oleic acid-dependent repression of JA signaling pathway. AtWRKY33 plays a role in Arabidopsis tolerance to high concentrations of NaCl concentration, while AtWRKY22 appears to be involved in the regulation of dark-induced leaf senescence in Arabidopsis.

Gene duplication and divergence events have been suggested to be the main contributors to evolutionary momentum. An analysis of gene duplication in the current study indicated that 20 of 59 (33.89%) VvWRKY genes have arisen through either tandem or segmental duplication. Two pair of the WRKY genes, VvWRKY11/VvWRKY12 and VvWRKY40/VvWRKY41, appeared to have undergone tandem duplication (Figure 4). Due to the genome of grapevine originated from a hexaploidisation, several synteny blocks would be expected to exist in this genome. Among the 11 collinearity relationships, only three of them, VvWRKY2/44/53 were identified as triplicates (Figure 1). Taken together, this implies that low tandem and high segmental duplications existed in VvWRKY genes, which is in consistent with the results in Arabidopsis. Segmental duplicates may be more often retained due to subfunctionalisation, without increasing the likelihood of gene rearrangement.

VvWRKY genes involved in grapevine growth and development

Patterns of expression of all of the predicted coding members of the VvWRKY family in different grapevine tissues and at different developmental stages were examined using an expression atlas of V. vinifera cv Corvina. The analysis revealed that many VvWRKY genes can be grouped together based on their abundant expression in specific grapevine tissues, possibly reflecting their involvement in a common metabolic and/or developmental process. In addition to the well-known response of WRKY genes to biotic and abiotic stresses, a number of studies have provided evidence of a role for WRKY TFs in developmental processes, such as seed and trichome development, dormancy and germination, and senescence.

The expression analysis revealed that VvWRKY23 and 46, belonging to Group II-c, were highly expressed in ‘woody’ tissues, suggesting a putative role in the regulation of lignin biosynthesis. Involvement of WRKY genes in the lignin synthesis has been previously reported for one grapevine WRKY within the same group (VvWRKY01). In tobacco, VvWRKY01 was able to induce lignin accumulation by regulating lignin biosynthetic genes when overexpressed.

VvWRKY02 of Group II-b was found to be specifically expressed in leaf tissues and highly accumulated during senescence. In Arabidopsis, AtWRKY6, belonging to the same group as VvWRKY02, was strongly upregulated during leaf senescence and targets a senescence-specific receptor-like kinase (SIRK/FRK1). Several other Arabidopsis WRKY genes were also upregulated in the Arabidopsis leaf transcriptome during senescence. VvWRKY03, another gene belonging to Group II-c, was specifically induced during senescence and would be an interesting candidate for functional characterisation.

AtWRKY53 and AtWRKY70 play a critical roles in leaf senescence; however, none of the grapevine VvWRKY genes belonging to the same group (III) were determined to be highly expressed in senescencing leaves. Both VvWRKY13, 59 and 57, belonging to Group I, were highly expressed in stamen and pollen tissues and are closely related to Arabidopsis AtWRKY34, which was shown to have a role in coordinating gene expression during the formation of the tapetum and microspores.

Stress induced VvWRKY expression in grapevine

The involvement of WRKY TFs in the regulation of plant response to biotic and abiotic stresses has been described in many species. Recent studies have indicated that hormones involved in plant defence signalling pathways are mediated by the induction of WRKY TFs. The transcriptomic databases generated in previous studies of grapevine subjected to biotic and abiotic stresses, together with our qRT-PCR analysis, allowed us to identify VvWRKY genes putatively involved in stress response. Since Arabidopsis WRKYs involved in stress response mainly belong to Groups II-a and III, the current focus is also on VvWRKYs within the same groups.

All members of Group II-a, the least abundant subgroup in both grapevine and Arabidopsis, are associated with stress response. In particular, VvWRKY07, 08 and 28, had the strongest upregulation in response to drought stress. Water stress induces the accumulation of abscisic acid (ABA) which triggers closing of stomata, thus reducing water loss. In Arabidopsis, AtWRKY18, 40 and 60 of Group II-a exhibited a complex pattern of expression in plant responses to ABA and abiotic stresses. Although an ABA treatment was not included in our survey, VvWRKY08, which is closely related to these Arabidopsis WRKYs, was induced by cold, salt and drought stresses, and by SA and ethylene treatments. VvWRKY08 shares 99% amino-acid sequence identity with VpWRKY3, a Group II-a member of V. pseudoreticulata, which has been shown to be involved in the ABA signalling pathway, in response to salt stress, and to increase resistance to Ralstonia solanacearum when overexpressed in tobacco. The similar expression response of VvWRKY08 and VvWRKY3 in response to drought, SA and ethylene treatments, and E. necator (powdery mildew) infection suggests a putative role for VvWRKY08 in grapevine plant stress tolerance that deserves further investigation.

Regarding biotic stresses, the low response of WRKY genes detected in resistant varieties compared to susceptible genotypes is not surprising since overall changes in the global transcriptome were generally lower in the resistant genotypes. In the case of E. necator infection, it is important to note that despite showing a low fold change in infected vs. mock-inoculated leaves in ‘Norton’ (resistant genotype), the majority of VvWRKY genes showed a higher level of constitutive expression in the mock-inoculated leaves compared to what was detected in ‘Cabernet sauvignon’ (susceptible genotype) mock-inoculated leaves. This indicates that, although a significant induction of VvWRKYs did not occur in response to infection, the baseline levels of these transcript are always higher in the resistant genotype than in the susceptible genotype.

VvWRKY07 and VvWRKY08, two members of Group II-a, are significantly upregulated in response to E. necator infection, suggesting that these two genes may be involved in the pathogen response pathway. The involvement of Group II-a WRKY genes in biotic responses has been previously shown in Arabidopsis where AtWRKY18 and AtWRKY40 play a significant role in controlling the basal defense. All members of Group III VvWRKYs were induced by both abiotic and biotic stresses, except for VvWRKY41. In Arabidopsis, AtWRKY70 and AtWRKY54, also in Group III, play an important role in plant defence against E. amylovora and in osmotic stress signalling. Interestingly, VvWRKY25, which is orthologous to AtWRKY54, is significantly induced by cold and drought treatments, and by E. necator infection. VvWRKY25 shares 88.66% amino-acid sequence identity with VpWRKY1 of V. pseudoreticulata was also reported to be rapidly induced by SA treatment and E. necator infection. This suggests that VvWRKY25 may also play a role in mediating plant defense response in grapevine V. vinifera.

Despite many recent advances in functional studies of WRKYs in grapevine, the biological function of most VvWRKY genes in physiological and developmental processes and plant defence still needs to be elucidated. The bioinformatic analysis and expression patterns of the VvWRKY gene family conducted in the present study...
provide an overall picture of the composition and expression of WRKY genes in grapevine that will facilitate selecting candidate genes for cloning and further functional characterisation.

CONCLUSIONS
Fifty-nine WRKY genes in grapevine were bioinformatically identified and characterized. The VvWRKY family genes were classified into three main groups, with high similar exon–intron structures and motif compositions within the same groups and subgroups. VvWRKY genes play important roles in grapevine growth and development as indicated by their spatial and temporal expression profiles. Group II-a and Group III genes exhibited different patterns of expression in response to different stresses. Most members in Group II-a and Group III were expressed under drought stress, and represent good candidate genes for exploring the role of WRKYs in grapevine stress response pathways.

Supplemental information for this article can be found on the Horticulture Research website (http://www.nature.com/hortres).

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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