

1 **Changes in plant metabolism and accumulation of fungal metabolites in response to**
2 **Esca proper and apoplexy expression in the whole grapevine**
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ABSTRACT

Trunk diseases have become among the most important grapevine diseases worldwide. They are caused by fungal pathogens that attack the permanent woody structure of the vines and cause various symptoms in woody and annual organs. This study examined modifications of plant responses in green stem, cordon and trunk of grapevines expressing Esca proper (E) or apoplexy (A) event, which are the most frequent grapevine trunk disease symptoms observed in Europe. Transcript expression of a set of plant defense- and stress-related genes was monitored by quantitative RT-PCR while plant phytoalexins and fungal metabolites were quantified by HPLC-MS in order to characterize the interaction between the grapevine and trunk disease agents. Expression of genes encoding enzymes of the phenylpropanoid pathway and *trans*-resveratrol content were altered in the three organs of diseased plants, especially in the young tissues of A plants. PR proteins and the antioxidant system were severely modulated in A plants, which indicates a drastic stress effect. In the meantime, fungal polyketides 6-MSA, (*R*)-mellein and (*3R,4R*)-4-hydroxymellein, were accumulated in A plants that suggests their potential effect on plant metabolism during the appearance of foliar symptoms.

KEYWORDS: *green stem, trunk, cordon, phenolic compounds, defense, qRT-PCR, trunk diseases, Chardonnay, black streaked wood, fungal metabolites*

42 INTRODUCTION

43

44 Trunk diseases have become among the most important grapevine diseases in the past 10-15
45 years. Considering a replacement of 1% of plants per year – a considerable underestimate in
46 view of the individual regional data found in the literature – the worldwide annual financial
47 cost of the replacement of death plants due to grapevine trunk diseases is more than €1.5
48 billion (Hofstetter et al. 2012). For example, in France it has been estimated that 13% of
49 vineyards are unproductive due to trunk diseases (Bruez et al. 2013), with an annual cost of
50 €14 million.

51 Esca disease, Eutypa- and Botryosphaeria- dieback are the major grapevine trunk diseases.
52 These diseases are caused by diverse fungal pathogens which attack the woody perennial
53 organs of the vine and ultimately lead to its death (Larignon and Dubos 1997; Mugnai et al.
54 1999; Bertsch et al. 2013). In regard to aetiology, the symptoms that occur in wood and
55 annual organs have been extensively described, revealing that Eutypa dieback symptoms
56 differ from those of Esca disease and Botryosphaeria dieback (Bertsch et al. 2013). Moreover,
57 various fungal species associated with grapevine trunk diseases coexist in the same area of a
58 wood section, *Fomitiporia mediterranea* M. Fischer, *Phaeomoniella chlamydospora* (W.
59 Gams, Crous, M.J. Wingfield & L. Mugnai) P.W. Crous & W. Gams, *Phaeoacremonium*
60 *minimum* (Tul. & C. Tul.) D. Gramaje, L. Mostert & Crous, comb. nov., *Botryosphaeriaceae*
61 species and *Eutypa lata* (Pers; Fr) Tul & C. Tul (Spagnolo et al. 2012). The most frequent
62 symptoms of trunk diseases observed in the Champagne area were assigned to Esca proper
63 (E) and apoplexy (A) (Grosman and Doublet 2012), which have an important impact on
64 French vineyards. Esca proper is the term proposed to indicate the coexistence of two
65 different syndromes, within the Esca disease complex named ‘Esca’ and ‘Grapevine leaf
66 stripe disease’ (GLSD), on a same plant (Surico 2009; Bertsch et al. 2013). Causal agents of

Esca are different wood rot fungi represented mainly by *F. mediterranea* in Europe and the Mediterranean basin (Surico et al. 2008). GLSD is a tracheomycotic widespread syndrome which major causal agents are considered to be *P. chlamydospora* and *P. minimum* (Surico et al. 2008; Surico 2009). Symptoms of GLSD occur in the wood of trunk and cordons, as well as in leaves and berries. Apoplexy, consisting in the partial or complete sudden wilting of the crown, is regarded as the acute form of GLSD and/or E (Surico et al. 2008; Letousey et al. 2010; Bertsch et al. 2013). Thus, affected plants die within a few years (Larignon et al. 2009). Recently, physiological changes in *Vitis* plants affected by E (GLSD) or A were reported especially on leaves, and at a lower extent also in green stem and trunk (Petit et al. 2006; Agrelli et al. 2009; Letousey et al. 2010; Magnin-Robert et al. 2011; Spagnolo et al. 2012; Magnin-Robert et al. 2014; Fontaine et al. 2015). Both leaves and green stems of affected *Vitis vinifera* L. plants showed physiological and metabolic changes related to the external symptoms, although no pathogens associated with E or other trunk diseases have been isolated from these organs (Lima et al. 2010; Magnin-Robert et al. 2011; Spagnolo et al. 2012).

It has been hypothesized that external symptoms of Esca disease complex are caused by toxins produced by fungi in the woody tissues and then translocated to the leaves via the transpiration stream (Mugnai et al. 1999). Polyketides represent an important category of fungal secondary metabolites commonly produced by causal agents of grapevine trunk diseases (*P. minimum*, *P. chlamydospora*, *Neofusicoccum parvum*). These compounds have been identified and have phytotoxic activity on grapevine protoplasts, calli and leaves (Evidente et al. 2000; Tabacchi et al. 2000; Abou-Mansour et al. 2004; Djoukeng et al. 2009; Andolfi et al. 2012, 2014; Abou-Mansour et al. 2015). Their impacts on plant physiology remain a crucial step to appreciate the mechanisms leading to disease emergence. A differential response related to symptoms of E and A was reported on leaves and green stems,

while no relevant differences were recorded in the trunk. In details, the primary metabolism and the defense response were more or less modified according to the form of symptom expression in leaves and in green stems (Petit et al. 2006; Magnin-Robert et al. 2011; Spagnolo et al. 2014). In the trunk, proteins involved in cell growth and defense response are down expressed in asymptomatic wood. Oppositely proteins related to defense were over expressed in the black streaked wood, characterized by the large presence of trunk diseases agents (Magnin-Robert et al. 2014). All these findings are useful to better understand the host-pathogen interactions but they were obtained on separate plants which is not appropriate to provide a reliable overview of symptom development in case of E and A.

For all these reasons, the goal of this work was to investigate plants affected by E and A, through analyzing physiological perturbations on both herbaceous and woody samples in a same plant. We focused on phenylpropanoid pathway by analysing the total phenolic compounds, the stilbene content and the expression of 9 related genes. The expression of stress defense response genes and 2 water-stress related genes as well as the abscisic acid quantification were also performed. Moreover, known fungal metabolites such as 6-methylsalicylic acid, terreutin, scytalone, isosclerone, (*R*)-mellein and (3*R*,4*R*)-4-hydroxymellein were quantified to characterize the fungus-plant interaction.

MATERIAL AND METHODS

Plant material

Fifteen standing vines (cv. Chardonnay/41B) were uprooted in both summer 2010 and 2011 from a vineyard located in the province of Epernay (Champagne-Ardenne region, France) owned by the company Moët & Chandon and planted in 1984. Five plants represented external leaf symptoms (GLSD) or A respectively, and 5 asymptomatic plants were collected each year. Asymptomatic plants were chosen among those that had shown neither GLSD nor

117 A symptoms since 2001, and were thus regarded as visually unaffected plants (control plants,
 118 C). Typical wood symptoms of GLSD and Esca were noted in all plants examined, including
 119 control plants. Therefore, plants showing foliar and wood symptoms of GLSD were
 120 considered as E plants. Four groups of samples were defined for green stems: C (stems from
 121 control plants), A and E (symptomatic stems from apoplectic (A) and Esca proper (E)-
 122 affected plants) and aS (asymptomatic stems from A and E plants) (Fig. 1). In woody tissues,
 123 2 types of samples were studied: asymptomatic and black streaked wood. Black streaking
 124 consists of single or more xylem vessels gathered into individual blackish brown bundles
 125 (Surico et al. 2008). Both woody tissues were sampled in young annual rings (estimated less
 126 than 5-year-old wood, Fig. 1). Finally, six groups of samples were defined for woody organs
 127 (cordon and trunk): asymptomatic wood of control (C1), Esca proper-affected (E1) and
 128 apoplectic (A1) plants; and black streaked wood of control (C2), Esca proper-affected (E2)
 129 and apoplectic (A2) plants (Fig. 1). Plant tissues were frozen in the field with liquid nitrogen
 130 to halt enzymatic activities and stored at -80°C prior to use. Before each analysis, the amount
 131 of biological sample needed was ground to a fine powder in liquid nitrogen with a Mixer Mill
 132 MM 400 (Retsch, Haan, Germany). During 2010 season, various organs of the sampled vines
 133 (trunk, cordon and green stems) were inspected internally for the presence of discolorations
 134 associated with trunk diseases, and subsequently subjected to fungal isolation as described by
 135 Spagnolo et al. (2012). The major causal agents of Esca proper (*P. chlamydospora*, *P.*
 136 *minimum* and *F. mediterranea*) were isolated from discoloured woody tissues of the three
 137 groups of plants (E, A and control plants). *F. mediterranea* was directly linked to white rot.
 138 Other fungi associated with grapevine trunk diseases, such as Botryophariaceae species and
 139 *E. lata* were also isolated. In the opposite, no fungi were detected from either non discolored
 140 wood of trunk and cordons, or discolored and non-discolored woody tissues of one-year-old
 141 stems, as well as from green stems of control or diseased plants (Spagnolo et al. 2012).

142

143 RNA extraction

144 Total RNA was isolated from 2 × 50 mg of powdered green stem tissues and 3 × 50 mg of
145 woody tissues (cordon and trunk) using the Plant RNA Purification Reagent (Invitrogen,
146 Cergy Pontoise, France). The RNA pellet was re suspended in 20 µL of RNase-free water,
147 then treated with RQ1 DNase enzyme (Promega) and quantified by measuring the absorbance
148 at 260 nm following manufacturer's instructions.

149

150 Real-time RT-PCR analysis of gene expression

151 In total, 150 ng of total RNA were reverse-transcribed using the Verso SYBR 2-step QRT
152 ROX enzyme (ABgene, Surrey, UK) according to the manufacturer's protocol. PCR
153 conditions were those described by Bézier et al. (2002). Expressions of 22 targeted genes
154 selected from previous published studies, whose expression and/or accumulation of
155 corresponding proteins are altered either in field-grown grapevine affected by trunk diseases
156 or *in vitro* conditions in response to fungal extracts (Magnin-Robert et al. 2011; Spagnolo et
157 al. 2012; Magnin-Robert et al. 2014; Ramírez-Suero et al. 2014; Spagnolo et al. 2014), were
158 tracked by quantitative Reverse-Transcripts Polymerase Chain Reaction (qRT-PCR) using the
159 primers indicated in Table 1. Reactions were carried out in a real-time PCR detector Chromo
160 4 apparatus (Bio-Rad) using the following thermal profile: 15 s at 95°C (denaturation) and 1
161 min at 60°C (annealing/extension) for 40 cycles. Efficiency of the primer sets was estimated
162 by performing real-time PCR on several dilutions. PCR reactions were performed in
163 duplicate. Results correspond to means ± standard deviation of 10 plants sampled, 5 in 2010
164 and 5 in 2011. The data were analysed using CFX Manager software, and the relative levels
165 of gene expression were determined following the method of Hellemans et al. (2007) with
166 *EF1-α* and *39SRP* serving as the two internal reference genes. The results represent the

relative expression in grapevine tissues of diseased plant (A or E) versus those corresponding to control (C).

Control samples consisted of plant tissues (green stem, C; cordon, C1 and C2; and trunk, C1 and C2) collected from control plants. The analyzed genes were considered significantly up- or down-regulated when change of their expression was $>2\times$ or $<0.5\times$ respectively.

Quantification of total polyphenols and phytoalexins

Sample extraction: Methanolic extracts were prepared from 50 mg of powdered herbaceous and woody tissues mixed with 1 ml of methanol (MeOH) and 25 μL of the internal standard *trans*-4-hydroxystilbene (0.5 mg mL^{-1}) according to Spagnolo et al. (2014).

Quantification of total phenolic compounds: Total phenolics were determined by using the Folin-Ciocalteu method (Singleton and Rossi 1965) downscaled to 96-well-plate (E. Abou-Mansour, personal communication). An aliquot (30 μL) of appropriate dilution (green stem 1:10 (v:v); woody tissues, 1:20 (v:v)) of methanolic extract was mixed with 150 μL of Folin-Ciocalteu reagent (diluted by 10) and after 5 min of incubation at room temperature, 120 μL of sodium carbonate solution (10% w:v) were added. After incubation at room temperature for 2 h the absorbance of the mixture was read against the prepared blank at 750 nm. For each sample, three independent extractions were quantified by three technical replicates. Total phenolics were expressed as mg of gallic acid equivalents (GAE) per g of plant tissues, values presented in the table correspond to means \pm standard deviation of one representative experiment out of two ($n_{2011} = 5$ plants per condition).

Quantification of stilbenes: Standards such as *trans*-piceid, *trans*-resveratrol and *trans*-pterostilbene were purchased from Extrasynthèse (Genay - France). The *trans*- ϵ -viniferin, *trans*-vitisin A and *trans*-vitisin B (Supplemental Figure 2) were extracted from lignified canes of Syrah as described by Spagnolo et al. (2014). Sixty μL of the samples were analysed

on a MN Nucleosil C18 analytical column (250 mm x 4 mm i.d., 100-5) (Machery-Nagel, Duren, Germany) using a flow rate of 0.7 mL min⁻¹ at 27°C. The mobile phase consisted of water/formic acid (0.5%) (solvent A) and acetonitrile (solvent B). The linear gradient started with 5% of B for 5 min and increased to 55% within 25 min reaching 80% at 28 min and 100% at 32 min. Spectral data for all peaks were accumulated in the range between 220 and 600 nm. The quantification of stilbenes was performed at λ 320 nm using internal standard calibration methods. For each sample, three independent extractions were analysed. The data are reported as $\mu\text{g}\cdot\text{g}^{-1}$ of fresh weight, values for stilbenes correspond to means \pm standard deviation of one representative experiment out of two ($n_{2011} = 5$ plants per condition).

Quantification of fungal phytotoxins

Screening of fungal toxins: Fungal phytotoxins scytalone, 4-hydroxyscytalone and isosclerone were obtained from a culture of *P. minimum* as described in Abou-Mansour et al. (2004), (*R*)-mellein, (*3R,4R*)-4-hydroxymellein, 6-methylsalicylic acid (6-MSA) and (-)-terremutin from a culture of *N. parvum* according to the method described in Abou-Mansour et al. (2015).

Identification and quantification of fungal metabolites: 500 μL from the methanolic extract prepared for the stilbene analysis were concentrated to 100 μL and extracted with *n*-hexane 3 x 100 μL to recover (*R*)-mellein, (*3R,4R*)-4-hydroxymellein and 6-MSA. The *n*-hexane extracts were pooled and dried under nitrogen and dissolved in 50 μL of MeOH, finally 10 μL were analysed by HPLC-MS. The remaining methanolic extracts were dried under a stream of nitrogen, dissolved in 80 μL of MeOH and 50 μL injected in the HPLC-MS. Analyses were performed on a HPLC-DAD (Agilent 1100) coupled to a quadrupole mass spectrometer Agilent MSD/SL G1956B. The column used for the analysis of (*R*)-mellein, (*3R,4R*)-4-hydroxymellein and 6-MSA was a MN Nucleodur Phenyl-Hexyl 2.7 μm (150 mm x 4.6 mm i.d.) (Machery-Nagel GmbH) with a mobile phase of water 0.1% formic acid (solvent A) and

217 acetonitrile 0.1% formic acid (solvent B). The gradient started with 5% B for 2 min, B
 218 reached 40% at 12 min, 60% at 23 min and 100% at 26 min, until 30 min. The column is then
 219 re-equilibrated to 5% B during 5 min. The temperature of the column was 30°C and the flow
 220 rate 0.4 mL min⁻¹. Scytalone, 4-hydroxyscytalone, isosclerone and (-)-terremutin were
 221 analysed on a Kromasil C18 column 5 µm (250 mm x 4.6 mm i.d.) with a mobile phase of
 222 water 0.1% formic acid (solvent A) and acetonitrile 0.1% formic acid (solvent B). The
 223 temperature of the column was 35°C, the flow rate 1 mL.min⁻¹ and the split between the DAD
 224 and the MSD of 20% with the same gradient as reported in Abou-Mansour et al. (2015) for
 225 (-)-terremutin analysis. The MS detector consisted of a simple quadrupole mass detector
 226 operated in ESI negative ionisation mode for scytalone, 4-hydroxyscytalone, isosclerone,
 227 (3*R*,4*R*)-4-hydroxymellein, (-)-terremutin and 6-MSA and in positive mode for (*R*)-mellein.
 228 The source was operated with drying gas N₂ at 12 L min⁻¹. The MS parameters were
 229 optimised by injection of the individual solution of the phytotoxins: fragmentor voltage
 230 varying from 80 V to 125 V. The capillary voltage was 3.5 kV in ESI+ and -3.5 kV in ESI-,
 231 the nebulizer pressure 35 psig and the drying gas temperature 300°C. Analyses were carried
 232 in SIM mode. The precursor ion and the most abundant fragment were chosen for
 233 quantification of the phytotoxins, the fragmentation pattern of 6-MSA *m/z* 151/107, of
 234 (-)-terremutin 155/113, of scytalone 193/113, 4-hydroxyscytalone 209/113, isosclerone
 235 177/113, of (*R*)-mellein *m/z* 179/161, (3*R*,4*R*)-4-hydroxymellein 193/149. The limit of
 236 detection was of 0.02 ng for 6-MSA, (*R*)-mellein and (3*R*,4*R*)-4-hydroxymellein and
 237 (-)-terremutin. At 0.1 ng, scytalone and 4-hydroxyscytalone were clearly detected and
 238 isosclerone was at the limit of detection. Compounds were identified according to their
 239 fragmentation pattern coupled to their retention time. Quantification was performed using
 240 external standard calibration method. For each sample, three independent extractions were
 241 analysed. The data are reported as ng g⁻¹ of fresh weight. Values for fungal metabolites

correspond to means \pm standard deviation of one representative experiment out of two (n_{2011} = 5 plants per condition).

Quantification of abscisic acid (ABA)

The method described by Schmelz et al. (2004) was used with some modifications. *Sample preparation*: 300 mg of grapevine powdered tissues were transferred to a screw cap tubes and homogenized twice with 1mL of extraction buffer (1-propanol/H₂O/HCl: 2/1/0.005) at 70°C. Samples were transferred to a glass tube and 100 ng of internal standard abscisic acid-d₆ (Santa Cruz Biotechnology, www.scbt.com) was added. Two mL of methylene chloride was added to each sample and mixed for 15 s with a vortex and centrifuged at 2000 g for 20 min. The lower organic phase was transferred into a 4 ml glass vial and dried by the addition of anhydrous Na₂SO₄. Before derivatisation, the volume of MeCl₂: 1-propanol solvent was reduced until approximately 400 μ L. *Derivatisation*: carboxylic acids including ABA were methylated to their corresponding methyl esters by addition of 50 μ L of methanol and 20 μ L of 2M bis-trimethylsilyldiazomethane (Sigma-Aldrich) at room temperature (RT) for 30 min. Excess of bis-trimethylsilyldiazomethane was quenched by adding 20 μ L of 2M acetic acid during 30 min at RT. *Vapor phase extraction (VOC)*: extraction of the vapor phase was performed using a VOC column (www.ars-fla.com) conditioned with 3 x 1 mL of MeCl₂. The VOC column and a nitrogen needle were fixed on the screw cap of the tube and solvent was evaporated under a nitrogen stream at 70°C, then the tubes were heated for 2.5 min at 200°C. The VOC column was eluted with 1 mL of MeCl₂. Finally the eluate was evaporated and samples dissolved in 60 μ L of hexane before injecting 5 μ L on a capillary column HP1 (25 m x 0.25 mm) GC column (Agilent Technologies) fitted to a Hewlett Packard 5980 GC coupled to a 5970 mass specific detector. The methyl esters of ABA and ABA-d₆ were detected and quantified by selective ion monitoring at m/z 190 and 194 respectively. The amount of ABA

(measured as methyl ABA) was calculated by reference to the amount of internal standard. The results are expressed in $\mu\text{g g}^{-1}$ fresh weight of plant tissue. For each sample, three independent extractions were analysed. ABA values correspond to means \pm standard deviation of one representative experiment out of two ($n_{2011} = 5$ plants per condition).

Statistical analysis

Results from quantification of metabolites (total polyphenols, phytoalexins, ABA and fungal metabolites) correspond to means \pm standard deviation of one representative experiment out of two (n_{2011} plants = 5). For relative expression of targeted genes, each value represents the mean of 10 plants per condition, 5 plants sampled in 2010 and 5 plants in 2011 ($n_{2010}+n_{2011}$). Error bars represent the standard deviation of the mean. A Kruskal Wallis test followed by the Dunn's multiple comparisons test (Prism 5., GraphPad Software, Inc., California, USA) were performed to compare relative genes expression and metabolites content between various conditions. Differences at $P < 0.05$ were considered to be significant.

RESULTS

Expression of genes involved in the phenylpropanoid metabolism

We designed primers for enzymes involved in the phenylpropanoid metabolism, including the proteins leading to the synthesis of phytoalexins (stilbenoids) and phytoanticipins (flavonoids). The expression analysis of 9 genes revealed a perturbation in the transcripts accumulation in the 3 organs (green stems, cordons and trunks) of E and A plants. The amounts of their transcripts in green stems of E plants were similar to those observed in the control plants (Fig. 2). In A plants, the analysis in green stems highlighted an up-regulation of a stilbene synthase (*STS*) and a down-regulation of 4 genes namely a caffeoyl-CoA O-methyltransferase (*CCoAOMT*), 2 isoflavone reductase (*IFRhom* and *IFRL4*) and a peroxidase

(*POX4*) (Fig. 2). In woody tissues, the transcript profile was most perturbed in the black streaked wood of diseased plants (Fig. 2). In cordons, a down-regulation of *IFRhom* and *IFRL4* expression, 2 genes involved in flavonoid pathway, was observed in the black streaked wood of plants affected by Esca proper (E2) and apoplexy (A2), respectively. In the opposite transcript accumulation of *STS* was stimulated in the black streaked wood of diseased plants (A2 and E2; Fig. 2), while that of phenylalanine ammonia lyase gene (*PAL*) was solely increased in the black streaked wood of A plants (A2). Only *CCoAOMT* expression was repressed in the asymptomatic cordon wood of E plants (Fig. 2). Even if the transcript accumulation of the *STS* gene was slightly stimulated in the trunk of A (A1, A2) and E (E1) plants, no significant modification of the transcript profile was noticed for *PAL* (Fig. 2). *IFRL4* and flavonoid-3'-hydroxylase (*F'3H*) genes were up-regulated in asymptomatic wood of A and E plants, respectively (Fig. 2). In the meantime, leucoanthocyanidin dioxygenase (*leucoAND*) was up-regulated in the trunk of A plants (A1, A2, Fig. 2).

Total polyphenolic and stilbene content

The amount of phenolic compounds was analyzed in the three organs (green stems, cordons and trunk) of control and diseased plants, by monitoring total phenolics and stilbenoids that are key molecules in vine defense responses. The content of *trans*-piceid, *trans*-resveratrol, *trans*- ϵ -viniferin, *trans*-vitisin A and *trans*-vitisin B was quantified in various samples (Table 2). The accumulation of the *trans*-piceid was quantified as it is a nontoxic glycosylated derivate of *trans*-resveratrol, which could be a form of *trans*-resveratrol storage in the plant (Belhadj et al. 2006).

In green stems, no difference was recorded in total phenol content between control and diseased plants (Table 2). Focusing on stilbenes, a slight content of *trans*-piceid was detected in green stems with apparent foliar symptoms (A and E). Moreover, *trans*-resveratrol and

317 *trans*-vitisin B were significantly accumulated in green stems of A plants (Table 2). In wood,
 318 the content of phenolics fluctuated depending on the samples. In the cordons, contents of
 319 *trans*-piceid, *trans*- ϵ -viniferin and *trans*-vitisin A were higher in the black streaked wood
 320 when compared to the asymptomatic wood, whatever the control or diseased plants (Table 2).
 321 Moreover, *trans*-vitisin B was quantified in A plants and in the black streaked wood (E2) of E
 322 plants (Table 2). In the trunk, total polyphenolic compounds were more important in the black
 323 streaked wood than in asymptomatic wood, in both control and diseased plants. Total
 324 polyphenolic compounds may indicate an accumulation of phenols related to the age of the
 325 wood, the level of phenols being higher in the trunk than in the cordon (Table 2). An increase
 326 of stilbenes related to the age of the wood was also observed in the black streaked wood (C2,
 327 A2, E2). Nevertheless, we noted a lower level of stilbenes in the asymptomatic wood of trunk
 328 when compared to cordons (C1, A1) (Table 2). These results may suggest that the
 329 accumulation of phenolic compounds in the asymptomatic wood of the trunk cannot be
 330 explained by the accumulation of stilbenes but by other phenols such as flavonoids.
 331 Moreover, the accumulation of total polyphenolic compounds seemed to be affected in the
 332 asymptomatic wood of the trunk from diseased plants, with statistical significance for E1
 333 (Table 2).

334 *PR protein transcript accumulation in green stems and woody tissues*

335 The expression of six genes encoding PR-proteins was investigated: a basic chitinase class I
 336 (*Chit1b*), a chitinase class IV (*Chi4c*), a chitinase class V (*CHV5*), a β -1,3-glucanase (*GLUC*),
 337 a glucan endo-1,3- β -glucosidase (*endoglu*) and a serine proteinase inhibitor (*PR6*). In green
 338 stems, the expression of the genes encoding PR proteins was higher in A than in E and aS
 339 (Fig. 3), although no perturbation of *endoglu* expression was noted in diseased plants. For the
 340 3 genes encoding chitinase, the accumulation of *CHV5* transcripts was the highest in response

to the appearance of foliar symptoms. Moreover, the transcript accumulation of *GLUC* was at least 50-fold higher in A than in E and aS (Fig. 3).

In wood, the transcript analysis indicated a perturbation of gene expressions with higher amplitude in cordons, “young” tissues, than in trunks. For *CHV5*, *Gluc* and *PR6* expression, the highest induction was recorded in cordon of A plants (A1 and A2, Fig. 3). Apart from a slight induction of *Chi4C* gene expression in A1, no significant alterations for *Chi4C* (A2, E1 and E2), *Chit1b* and *endoglu* expression were observed in cordons (Fig. 3). In trunks of diseased plants, the expression of *CHV5*, *Gluc* and *PR6* were increased. For both *Gluc* and *CHV5*, the relative expression was 35-fold higher in asymptomatic wood of A1 and E1 plants (Fig. 3). For *endoglu*, its transcript accumulation was only observed in asymptomatic trunks of A1 and E1, whereas the expression of both *Chi4C* and *Chit1b* genes was not affected in the trunk of diseased plants.

Expression pattern of stress-related genes in green stems and woody tissues

To determine whether stress responses were triggered in A or E grapevines, the expression of 3 detoxification and stress tolerance genes were followed in green stems and woody tissues of diseased plants (Fig. 4): an epoxide hydrolase (*epoxH2*), a glutathion-S-transferase (*GSTI*) and a superoxide dismutase (*SOD*). Two genes encoding Heat Shock Protein (HSP) were also investigated, 70kDa HSP (*HSP70*) and a small chloroplastic HSP (*HSP*).

In green stems, no significant changes of expression of these targeted genes occurred in aS of diseased plants. *GSTI* gene was up-regulated in symptomatic stems by 20- and 4.5-fold higher in A and E plants, respectively. Moreover *epoxH2* was only and slightly up-regulated in symptomatic green stems of A plants (Fig. 4). A repression of *HSP* gene expression was observed in symptomatic stems and was higher in A than in E plants (Fig. 4).

365 Except for the *GSTI* gene, no significant changes occurred in cordons for the stress-related
 366 genes (Fig. 4). *GSTI* expression was 40- and 18-fold stimulated in A1 and A2 plants
 367 respectively, whereas it was repressed by 4-fold in E1 and E2 plants. In the trunk, *GSTI* was
 368 up-regulated in diseased plants by 5-fold in A1, E1 and E2 and by 10-fold in A2 (Fig. 4). The
 369 expression of *epoxH2* was only induced in asymptomatic wood of diseased plants (A1, E1). In
 370 the meantime, a general repression of *HSP* was observed in the trunk of diseased plants (Fig.
 371 4). Nevertheless, the relative expression of *HSP* was significantly induced in E1 (Fig. 4). *SOD*
 372 was down-regulated in asymptomatic wood of A1, while no significant change occurred in
 373 A2, E1 and E2 (Fig. 4).

374

375 *Analysis of fungal metabolite contents*

376 The role of fungal metabolites in causing symptoms on diseased-plants is still unclear.
 377 Whereas the internal inspection of trunk described the presence of major causal agents of
 378 trunk diseases (*P. chlamydospora*, *P. minimum* and *F. mediterranea*, *E. lata* and
 379 Botryophariaceae species) in our experimental vineyard (Spagnolo et al. 2012), we screened
 380 the different samples looking for phytotoxins from these various fungi. We checked fungal
 381 metabolites, such as scytalone, 4-hydroxyscytalone for the presence of *P. minimum* and *P.*
 382 *chlamydospora* (Evidente et al. 2000; Abou-Mansour et al. 2004), isosclerone for the
 383 presence of *P. minimum*, *P. chlamydospora* and *N. parvum* (Evidente et al. 2000; 2010;
 384 Abou-Mansour et al. 2004; Evidente et al , (R)-mellein, (3R,4R)-4-hydroxymellein for *D.*
 385 *seriata* along with 6-MSA and (-)-terremutin for *N. parvum* (Djoukeng et al. 2009; Abou-
 386 Mansour et al. 2015). Scytalone, 4-hydroxyscytalone and isosclerone were not detected in
 387 grapevine tissues of control and diseased plants. (R)-mellein, (3R,4R)-4-hydroxymellein and
 388 6-MSA statistically increased especially (3R,4R)-4-hydroxymellein content in green stems,
 389 cordons (A1) and trunks (A1 and A2) of A vines in comparison with control and E plants

(Fig. 5) and (-)-terremutin was only detected but not quantified in green stems of A (data not shown).

Hormone accumulation and expression of genes involved in water-stress related responses

As the apoplexy is characterized by a sudden wilting of berries and leaves, we further investigated the impact /participation of the hydric stress in symptom expression. In this optic, we designed specific primers for an aquaporine, tonoplast intrinsic proteins (*TIP1*) involved in water transport and a nine-*cis*-epoxycarotenoid dioxygenase 2 (*NCED2*) involved in the biosynthesis of the water stress hormone, abscisic acid (ABA). Moreover, the content of ABA was analyzed in green stems, cordons and trunks. Expression of *TIP1* was up-regulated in green stems (A) and cordons (A1 and A2) of A plants, but not in E plants (Fig. 6). No change of *NCED2* expression profile was observed in diseased plants, except for a down-regulation of *NCED2* in trunks of E2 plants (Fig. 6). The basal content of ABA was 1 ng mg⁻¹ FW in green stems. The content was 2-fold (0.50 ng mg⁻¹ FW) and 4-fold lower (0.25 ng mg⁻¹ FW) in cordons and trunks, respectively (Fig. 7), but no difference was noticed in green stems and cordons between control and diseased plants. For trunks, only a slight accumulation of ABA was detected in black streaked wood of E plants (Fig. 7).

DISCUSSION

No information was previously reported about the whole dynamic process of defense response in the entire plant during external symptom emergence of trunk diseases. The characterization of the stress responses observed in the whole plant in relation with the presence of foliar symptoms is essential to appreciate the impact of these vascular diseases on plant physiology. Moreover, little information is available on the ability of the various plant organs of different ages to activate and develop efficient defenses. In this context, our study clearly demonstrates

415 an alteration of polyphenols contents as well as a modification of stress responses in green
 416 stems and in woody of A and E grapevines, simultaneously at the onset of foliar symptom
 417 development.

418

419 *Alteration of the phenylpropanoid pathway, especially in A plants*

420 To compare the defense responses of grapevine affected by either E or A, we decided to target
 421 the best characterized active defense mechanism, namely the phenylpropanoid pathway.
 422 Preliminary studies described the phenolics accumulation in annual and in woody tissues of
 423 plants affected by trunk disease agents (leaves and berries, Calzarano et al. 2008; Lima et al.
 424 2010; trunks, Amalfitano et al. 2000; Graniti et al. 2000; Del Rio et al. 2004; Martin et al.
 425 2009) and also by other pathogenic agents, such as the xylem-infecting bacterium *Xylella*
 426 *fastidiosa* (Pierce's disease) (Wallis and Chen 2012) or fungal agents involved in grapevine
 427 foliar diseases (*Botrytis cinerea*, *Plasmopara viticola* or *Erysiphe necator*) (Langcake and
 428 Pryce 1976; Dercks and Creasy 1989; Bavaresco et al. 1997; Romero-Perez et al. 2001).
 429 Indeed, a reaction to pathogen attacks is the formation of papillae (poly-phenol rich reactions)
 430 in secondary xylem to compartmentalize pathogens in woody tissues. In this work, alteration
 431 of gene expression involved in phenylpropanoid metabolism included significant
 432 accumulations of *trans*-resveratrol and *trans*-vitisin B, in various tissues of diseased vines,
 433 especially in green stems of A plants. These results indicate a higher effect on the
 434 phenylpropanoid pathway in young grapevine tissues caused by the stress provoked by
 435 apoplexy event in comparison to the onset of E symptoms.

436 Regarding woody tissues, we observed strong variations of phenolic contents between
 437 asymptomatic and black streaked wood, for both control and diseased plants. These
 438 differences can be explained by the presence/absence of trunk diseases agent inocula.
 439 Actually, *P. chlamydospora*, *P. minimum*, *F. mediterranea* and Botryosphaeriaceae species

440 were isolated from black streaked wood, while no fungi were isolated from asymptomatic
 441 wood (Spagnolo et al. 2012). Thus, the black streaked wood corresponds to the zone of
 442 interaction between the plant and the fungal agents but no correlation has been observed
 443 between the emergence of these lesions in the trunk and the capacity of the plant to slow
 444 down the colonization by the fungi. Lambert et al. (2012) described *in vitro* the tolerance of
 445 *P. chlamydospora*, *F. mediterranea* and *P. minimum* to various phenolics. For
 446 Botryosphaeriaceae species, *D. seriata* was described to be more susceptible than *N. parvum*.
 447 All together, these results indicate that the accumulation of phenolics in woody tissues may
 448 participate in plant defense reactions to limit wood colonization by trunk diseases agents,
 449 however their fungicidal activity depends on the pathogenic agents (Lambert et al. 2012). In
 450 our study, the content of phenolic compounds was lower in asymptomatic wood of diseased
 451 plants, except for stilbenes. This suggests that another group(s) of phenolic compounds can be
 452 altered in diseased plants (both A and E) infected by trunk disease causal agents. Expression
 453 of three targeted genes involved in flavonoid and anthocyanin biosynthesis, *leucoAND*
 454 (X75966, also referenced VIT_02s0025g04720), *CHI* (XM_002282072, also referenced
 455 VIT_13s0067g03820) and *F'3H* (XM_002284115, also referenced VIT_17s0000g07210),
 456 were altered in leaves of grapevine plantlets infected with *N. parvum* under greenhouse
 457 conditions (Czemmel et al. 2015). Alteration of *leucoAND* (X75966, also referenced
 458 Vv_10000352) was monitored in grapevine leaves infected by *Eutypa lata*; its up-regulation
 459 was associated to the lack of leaf symptoms (Camps et al. 2010). At metabolite level, the
 460 anthocyanin content was strongly affected in cell cultures exposed to eutypine, a toxin from
 461 *E. lata* (Afifi et al. 2003). Moreover, an interesting proteomic analysis revealed that proteins
 462 involved in isoflavonoid and anthocyanin biosynthesis decreased in asymptomatic wood of
 463 diseased vines (A and E) (Magnin-Robert et al. 2014). Future research is now required to

464 evaluate the shift between stilbenes and flavonoids in various organs connected with the vine
 465 susceptibility to trunk diseases.

466

467 *Modulation of specific defense responses as a consequence of disease expression*

468 In addition to its antimicrobial activity, *trans*-resveratrol can also act as a signaling molecule
 469 by the activation of defense-related responses on *Vitis* cell cultures: alkalinization, mild
 470 elevation of reactive oxygen species (ROS) and PR protein transcript accumulation (Chang et
 471 al. 2011). In this study, the induction level of 3 out of 6 PR-proteins, *CHV5*, *Gluc* and *PR6*,
 472 was higher in the 3 organs tested. This suggests that tissues perceive some elicitor signal
 473 associated with the presence of symptoms and also that these genes are inducible.

474 During xylem infection, typical metabolites changes lead to the accumulation of PR proteins
 475 in xylem sap (Rep et al. 2002; Basha et al. 2010). Similar to our study, the expression of the 3
 476 genes was induced in the black streaked of cordons and trunks where fungi live. Nevertheless,
 477 their induction was also detected in asymptomatic wood, with a higher level for both *Gluc* and
 478 *CHV5* in trunks. These observations may reveal a preventive strategy by the plant to limit
 479 future fungal colonization in asymptomatic wood of the trunk. Despite an up-regulation of
 480 PR-protein gene expression in diseased plants, the grapevine expressed external symptoms (A
 481 or E). Hence, PR-protein gene induction in our plant pathosystem was not sufficient to avoid
 482 symptoms expression. The protection of cellular functions can also be provided by proteins
 483 presenting a chaperone role, like the small heat shock proteins. A general down-regulation of
 484 *HSP* and *HSP70* expression was observed in the organs of A plants. In this sense, a low
 485 accumulation of these proteins was already observed in the brown stripe, a typical wood
 486 discoloration of *Botryosphaeria dieback* (Spagnolo et al. 2014). Their differential level in
 487 Pierce's disease (PD)-resistant and in (PD)-susceptible grapevine genotypes supports the idea
 488 that HSPs might be implicated in resistance (Yang et al. 2011). These results suggest that

489 HSPs are likely related to some cellular dysfunctions associated with the presence of
490 symptoms. During stress conditions, cellular structure is compromised by the formation of
491 excessive ROS due to disruption of cellular homeostasis. The scavenging or detoxification of
492 ROS excess is achieved by efficient enzymatic antioxidants system, like SOD and GST
493 (Bowler et al. 1992; Marrs 1996). Here, *SOD* expression was down-regulated in trunk of A
494 vines. Reductions of SOD protein content and of transcript accumulation were also reported
495 in various organs of A and E plants (Letousey et al. 2010; Magnin-Robert et al. 2011;
496 Spagnolo et al. 2012; Magnin-Robert et al. 2014), which indicates a lack of oxidative stress
497 control. The decrease of SODs expression could therefore be considered as a potential marker
498 of the onset of disease symptom emergence. GSTs are a large superfamily of enzymes, with
499 five classes defined *Theta*, *Zeta*, *Lambda*, *Phi* and *Tau*, the 2 later being specialized in the
500 conjugation of xenobiotics such as toxins (Frova, 2003). Focusing on the *tau* class, *GSTI*
501 expression was up-regulated in the 3 tested organs (this study) and also in the visually healthy
502 leaves of E plants (Magnin-Robert et al. 2011). Moreover, Valtaud et al. (2009) showed an
503 induction of leaf glutathione metabolism simultaneously with the onset of E foliar symptoms
504 appearance. Together, these results suggest that the perturbation of the antioxidant system
505 could be provoked by the presence of fungal metabolites. Phytotoxic metabolites secreted by
506 Esca disease pathogens are considered as toxins circulating in plant tissues and translocated to
507 the leaves *via* the transpiration stream (Mugnai et al. 1999).

508

509 *Activation of detoxification process and accumulation of specific fungal metabolites*

510 The major wood-infesting fungi *P. chlamydospora*, *P. minimum*, *F. mediterranea*, *E. lata* and
511 *Botryosphaeriaceae* sp. are known to produce diverse toxins (Tey-Tuhl et al. 1991; Andolfi et
512 al. 2011; Bertsch et al. 2013). We focused on the epoxHs, which catalyzes the detoxification
513 of xenobiotics by the conversion of epoxides to the corresponding diols (Morisseau and

Hammock 2005). Interestingly, several toxins produced by trunk disease agents are characterized by the presence of epoxides in their chemical structure (Abou-Mansour et al. 2015; Andolfi et al. 2011, 2012). Our study revealed an up-regulation of the *epoxH2* expression in both green stems and trunks of A plants. In the meantime, identification of various fungal metabolites reported as phytotoxins (Abou-Mansour et al. 2015) were undertaken in green stems, cordons and trunk. Our study revealed that the levels of (*R*)-mellein, its derivative (*3R,4R*)-4-hydroxymellein, along with 6-MSA were present at similar high content in tissues of A plants whereas the other toxins were not detected. (*R*)-mellein and its derivatives are widespread in fungi (Chooi et al. 2015) and were reported for an antigerminative activity, suggesting that they may interfere with the cellular pathway involved in germination or hormone signaling in plants (Chooi et al. 2015). In addition, (*R*)-mellein slows down the cell cycle, extending the mitotic phase (Essad and Bousquet 1981). Interestingly 6-MSA, a precursor of (-)-terremutin, is known to activate disease resistance in tobacco inducing accumulation of defense proteins and virus resistance, probably by mimicking SA hormone (Yalpani et al. 2001). High level of 6-MSA in A plants may thus explain the strong activation of defense response, such as PR protein accumulation. 6-MSA may act primarily by modulating the plant defense responses and then the polyketide toxins may affect the antioxidant system, suggesting a coordinated and dramatic effect which compromises the establishment of an appropriate and effective defense response able to avoid disease expression.

A strange relationship between water stress and disease expression

Grapevine xylem is an extreme case of efficiency/ sacrificial strategy for water transport and is therefore particularly vulnerable to drought stress-induced xylem cavitation. Fungal vascular pathogens are able to use wood polymers as energy sources and may alter the xylem

structure, which leads to a loss of xylem function. Infected grapevines may have a greater vulnerability with regard to water stress. Contrary to both defense and anti-oxidant response alteration, the establishment of the typical responses to water stress in diseased plants and especially in A ones seems to be less obvious. In response to water stress, Galmès et al. (2007) observed a rapid down-regulation in *TIP1* transcript accumulation. On the contrary, the less turgid leaves of A vines displayed an up-regulation of *TIP1*. Similar responses were observed in green stems and cordons. Another typical reaction of grapevine to drought stress is the increase of ABA biosynthesis on stems that regulate stomata opening (Christmann et al. 2007). Moreover, a great deal of evidence highlights the importance of ABA as a root-sourced signal transported *via* the xylem and involved in stomatal regulation of drought plants (for review, Dodd et al. 1996). Except for the black streaked wood of trunks from E plants, no significant change in the ABA content was observed in diseased plants. In agreement with Christen et al. (2007), our results suggest that the appearance of foliar symptoms cannot be simply considered as a water-deficit-inducing alteration but that other physiological mechanisms are involved.

To conclude, our results confirm a slight discrimination of plant responses between vines affected by A event and those by E symptoms. A drastic impact on phenylpropanoid pathway (*STS*, *IFRL4*, *IFRhom*), transcript accumulation of PR proteins (*CHV5*, *Gluc PR6*) and anti-oxidant system (*GST1*) was observed in herbaceous and woody tissues of A plants. This work also described the down-regulation of genes such as *HSP* and *SOD*, which suggested that these proteins may likely be related to cellular dysfunctions leading to the onset of foliar symptoms. As previously cited by Djoukeng et al (2009), the (*R*)-mellein and more particularly the hydroxylated derivatives, (*3R,4R*)-4-hydroxymellein, were efficient diagnostic markers of plants affected by the apoplectic form. Whether the detected fungal

564 metabolites play a role in this plant-fungus interaction is a question that warrants future
565 investigation.

566

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575

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820 Table 1: Primers of genes analyzed by real-time reverse-transcription polymerase chain
821 reaction.

	Genes	Primer sequences	Genbank or NCBI accession numbers
housekeeping genes	<i>EF1-α</i> (elongation factor 1-α)	5'-GAACTGGGTGCTTGATAGGC-3' 5'-AACC AAAATATCCGGAGTAAAAGA-3'	GU585871
	<i>39SRP</i> (39S ribosomal protein L41-A)	5'- GACTGACTTCAAGCTTAAACC-3' 5'-GATATAACAGGGAATACAGCAC-3'	XM_002285709.1
phenylpropanoid metabolism	<i>PAL</i> (phenylalanine ammonia lyase)	5'-TCCTCCCGGAAAACAGCTG-3' 5'-TCCTCCAAATGCCTCAAATCA-3'	X75967
	<i>CCoAOMT</i> (caffeoyl-CoA O-methyltransferase)	5'-GTGGACGCAGACAAGGACAAT-3' 5'-CCCGTTCCAGAGGGTGTG-3'	XM_002285070
	<i>STS</i> (stilbene synthase)	5'-AGGAAGCAGCATTGAAGGCTC-3' 5'-TGCACCAGGCATTCTACACC-3'	FJ851185
	<i>CHI</i> (chalcone isomerase)	5'-GCAGAAGCCAAAGCCATTGA-3' 5'-GCCGATGATGGACTCCAGTAC-3'	XM_002282072
	<i>IFRhom</i> (isoflavone reductase homolog-like)	5'-GCAACATACATCAAAACAATTGA-3' 5'-CCGTGAGCTTTTCCCATGTC-3'	XM_002266111
	<i>IFR-L4</i> (isoflavone reductase like protein 4)	5'-GGATCGTGTTAATGCGGTTGA-3' 5'-GCCTGGCTGGACCAATGTAG-3'	BN000709
	<i>F'3H</i> (flavonoid-3'-hydroxylase)	5'-CGCTTGTTTCATGCGTTCAAC-3' 5'-CTATTTTAAATCATGGGCAAACAACCT-3'	XM_002284115
	<i>leucoAND</i> (leucoanthocyanidin dioxygenase)	5'-ATGAGGGCAAGTGGGTGACA-3' 5'-TTGACCAGTCCCTGTGAAGA-3'	X75966
	<i>POX4</i> (peroxidase 4)	5'-AACATCCCCCTCCCACTT-3' 5'-TGCATCTCGCTTGGCTATT-3'	XM_002269882
	<i>Chit1b</i> (class I basic chitinase)	5'-ATGCTGCAGCAAGTTTGTT-3' 5'-CATCCTCCTGTGATGACATT-3'	Z54234
defense protein	<i>Chi4c</i> (class IV chitinase)	5'-TCGAATGCGATGGTGGA-3' 5'-TCCCTGTGCGAAACACCAAG-3'	AY137377
	<i>CHV5</i> (class V chitinase)	5'-CTACAACATATGGCGTGCTG-3' 5'-CCAAAACCATAATGCGGTCT-3'	AF532966
	<i>GLUC</i> (β-1,3-glucanase)	5'-TCAATGGCTGCAATGGTGC-3' 5'-CGGTCGATGTTGCGAGATTTA-3'	DQ267748
	<i>endoglu</i> (glucan endo-1,3-beta-glucosidase)	5'-AGATGGGCAGCTTGGTTACAA-3' 5'-TGAAGGCCAACCACTCTCTGA-3'	XM_002277410
	<i>PR6</i> (serine proteinase inhibitor)	5'- AGGGAACAATCGTTACCCAAG-3' 5'- CCGATGGTAGGGACACTGAT-3'	AY156047
	<i>epoxH2</i> (epoxide hydrolase 2)	5'-TCTGGATTCCGAAGTGCATTG-3' 5'-ACCCATGATTAGCAGCATTGG-3'	XM_002270484
detoxication and stress tolerance	<i>GSTI</i> (glutathion-S-transferase, tau form)	5'-TGCATGGAGGAGGAGTTCGT-3' 5'-CAAGGCTATATCCCCATTTCTTC-3'	AY156048
	<i>SOD</i> (superoxide dismutase)	5'- GTGGACCTAATGCAGTGATTGGA-3' 5'- TGCCAGTGGAAGGCTAAGTTCA-3'	AF056622
	<i>HSP70</i> (heat shock protein 70kDa)	5'- CAACATGAGGAACACTGTCAAAGAC-3' 5'-TGCAAGCTGGTTACTGTCCAA-3'	XM_002283496
	<i>HSP</i> (alpha crystalline heat shock protein)	5'-TCGGTGGAGGATGACTTGCT-3' 5'-CGTGTGCTGTACGAGCTGAAG-3'	XM_002272382
	<i>TIP1</i> (tonoplast intrinsic aquaporin)	5'-ATCACC AACCTCATT CATATGC-3' 5'-GTTGTTGTCTCAACCCATTTC-3'	AF271661
water stress	<i>NCED2</i> (9-cis-epoxycarotenoid dioxygenase 2)	5'-CTCTTGCCATGTCGGAAGA-3' 5'-CGGAGCTGCTGTGCGAAGTC-3'	XM_003632982.1

Table 2: Total phenolics and stilbene compounds concentrations in green stem: control stem (C), asymptomatic (aS) and symptomatic stems (A and E), and in both cordon and trunk: asymptomatic (C1, A1 and E1) and black streaked (C2, A2 and E2) wood of apoplectic (A) and Esca proper-affected (E) 26-year-old standing vines cv. Chardonnay.

Samples		Total phenolic compounds (mg GAE g ⁻¹ FW)	Stilbene compounds (mg g ⁻¹ FW)				
			<i>trans</i> -piceids	<i>trans</i> -resveratrol	<i>trans</i> -ε-viniferin	<i>trans</i> -vitisin A	<i>trans</i> -vitisin B
Green stem	C	8.11 ± 0.79 a	nd a	0.003 ± 0.002 a	0.093 ± 0.085 a	0.025 ± 0.019 a	nd a
	aS	9.72 ± 4.45 a	nd a	0.03 ± 0.02 a	0.29 ± 0.28 a	0.014 ± 0.013 a	0.008 ± 0.007 a
	A	9.33 ± 2.41 a	0.004 ± 0.002 a	0.55 ± 0.10 b	0.15 ± 0.06 a	0.064 ± 0.017 a	0.191 ± 0.048 b
	E	9.79 ± 1.23 a	0.006 ± 0.004 a	0.06 ± 0.04 a	0.44 ± 0.33 a	0.051 ± 0.011 a	0.013 ± 0.011 a
Cordon	C1	4.90 ± 0.61 a	0.13 ± 0.03 a	0.04 ± 0.02 a	0.17 ± 0.08 a	0.02 ± 0.01 a	nd a
	C2	8.61 ± 1.69 b	0.27 ± 0.02 b	0.52 ± 0.20 a	2.31 ± 0.76 b	0.35 ± 0.12 a	nd a
	A1	6.33 ± 2.28 ac	0.17 ± 0.06 ac	0.85 ± 0.43 a	0.55 ± 0.23 a	0.07 ± 0.04 a	0.07 ± 0.04 a
	A2	8.04 ± 1.37 bc	0.28 ± 0.09 bc	0.97 ± 0.23 a	1.47 ± 0.35 ab	0.16 ± 0.01 a	0.09 ± 0.03 a
	E1	4.58 ± 0.32 a	0.15 ± 0.02 a	0.07 ± 0.01 a	0.17 ± 0.1 a	0.02 ± 0.01 a	nd a
	E2	9.00 ± 1.89 b	0.41 ± 0.15 b	0.68 ± 0.35 a	1.6 ± 0.4 ab	0.21 ± 0.05 a	0.007 ± 0.005 a
Trunk	C1	24.81 ± 4.16 a	0.04 ± 0.02 a	0.004 ± 0.002 a	0.06 ± 0.03 a	0.008 ± 0.008 a	nd a
	C2	29.09 ± 7.30 ab	0.42 ± 0.04 b	0.65 ± 0.11 b	2.80 ± 0.14 b	0.69 ± 0.05 b	nd a
	A1	20.85 ± 2.72 ac	0.16 ± 0.04 a	0.12 ± 0.04 a	0.19 ± 0.08 a	0.02 ± 0.01 a	0.045 ± 0.030 a
	A2	34.25 ± 12.79 bc	0.38 ± 0.05 b	0.98 ± 0.29 b	4.00 ± 1.79 b	1.00 ± 0.50 b	nd a
	E1	16.16 ± 4.88 c	0.12 ± 0.01 a	0.09 ± 0.02 a	0.15 ± 0.03 a	0.04 ± 0.02 a	0.005 ± 0.003 a
	E2	32.81 ± 3.66 b	0.40 ± 0.04 b	1.69 ± 0.66 b	3.54 ± 1.19	1.06 ± 0.54 b	nd a

Lowercase letters a, b and c indicate significant difference ($\alpha=0.05$) for the concentrations found for each organs (Dunn's multiple comparison test, $P \leq 0.05$),

nd indicates no-detected. Total phenolics were expressed as milligram of gallic acid equivalents (GAE) per gram of plant tissues.

829 Figures caption

830

831 Fig 1: Description of sampling plants and sample codes used.

832

833 Fig. 2: Expression level of 9 selected genes involved in phenylpropanoid metabolism
834 determined by quantitative reverse-transcription polymerase chain reaction were observed in
835 green stems **A**, symptomatic (A and E) and asymptomatic (aS) green stems, cordon **B**, and
836 trunk **C**, asymptomatic (A1 and E1) and black streaked (A2 and E2) wood of apoplectic (A)
837 and Esca proper-affected (E) 26-year-old standing vines cv. Chardonnay. Gene expression
838 was considered as significantly up- or down-regulated to the 1x appropriate controls (dotted
839 lines), when changes in relative expression were $>2x$ or $<0.5x$, respectively. Results
840 correspond to means \pm S.D. (10 plants, $n_{2010}=5$ and $n_{2011}=5$). Columns headed by the same
841 letter are not significantly different (Dunn's multiple comparison test, $P<0.05$).

842

843 Fig. 3: Expression level of 6 selected genes encoded for PR proteins determined by
844 quantitative reverse-transcription polymerase chain reaction were observed in green stems **A**,
845 symptomatic (A and E) and asymptomatic (aS) green stems, cordon **B**, and trunk **C**,
846 asymptomatic (A1 and E1) and black streaked (A2 and E2) wood of apoplectic (A) and Esca
847 proper-affected (E) 26-year-old standing vines cv. Chardonnay. Gene expression was
848 considered as significantly up- or down-regulated to the 1x appropriate controls (dotted lines),
849 when changes in relative expression were $>2x$ or $<0.5x$, respectively. Results correspond to
850 means \pm S.D. (10 plants, $n_{2010}=5$ and $n_{2011}=5$). Columns headed by the same letter are not
851 significantly different (Dunn's multiple comparison test, $P<0.05$).

852

853 Fig. 4: Expression level of five selected genes implicated in detoxification and stress tolerance
 854 determined by quantitative reverse-transcription polymerase chain reaction were observed in
 855 green stems **A**, symptomatic (A and E) and asymptomatic (aS) green stems, cordon **B**, and
 856 trunk **C**, asymptomatic (A1 and E1) and black streaked (A2 and E2) wood of apoplectic (A)
 857 and Esca proper-affected (E) 26-year-old standing vines cv. Chardonnay. Gene expression
 858 was considered as significantly up- or down-regulated to the 1x appropriate controls (dotted
 859 lines), when changes in relative expression were $>2x$ or $<0.5x$, respectively. Results
 860 correspond to means \pm S.D. (10 plants, $n_{2010}=5$ and $n_{2011}=5$). Columns headed by the same
 861 letter are not significantly different (Dunn's multiple comparison test, $P<0.05$).

862

863 Fig. 5: Fungal metabolites (6-MSA, (*R*)-mellein and (3*R*,4*R*)-4-hydroxymellein) contents
 864 expressed in ng g^{-1} FW were determined in green stems: control stems (C) and symptomatic
 865 stems (A and E), both cordon and trunk: asymptomatic (C1, A1 and E1) and black streaked
 866 (C2, A2 and E2) wood of apoplectic (A) and Esca proper-affected (E) 26-year-old standing
 867 vines cv. Chardonnay. Results correspond to means \pm S.D. ($n_{2011}=3$). Data correspond to the
 868 fungal metabolites content evaluated in stems sampled in vineyard during the 2011
 869 season. *One asterisk indicates a statistically significant difference to the respective control
 870 sample (Dunn's multiple comparison test, $P<0.05$).

871

872 Fig. 6: Expression level of two selected genes implicated in water transport and ABA
 873 synthesis determined by quantitative reverse-transcription polymerase chain reaction were
 874 observed in green stems **A**, symptomatic (A and E) and asymptomatic (aS) green stems,
 875 cordon **B**, and trunk **C**, asymptomatic (A1 and E1) and black streaked (A2 and E2) wood of
 876 apoplectic (A) and Esca proper-affected (E) 26-year-old standing vines cv. Chardonnay. Gene
 877 expression was considered as significantly up- or down-regulated to the 1x appropriate

878 controls (dotted lines), when changes in relative expression were $>2x$ or $<0.5x$, respectively.
879 Results correspond to means \pm S.D. (10 plants, $n_{2010}=5$ and $n_{2011}=5$). Columns headed by the
880 same letter are not significantly different (Dunn's multiple comparison test, $P<0.05$).

881

882 Fig. 7: ABA contents expressed in $ng\ mg^{-1}$ FW were determined in green stems **A**, control
883 stems (C), asymptomatic (aS) and symptomatic stems (A and E), cordon **B**, and trunk **C**,
884 asymptomatic (C1, A1 and E1) and black streaked (C2, A2 and E2) wood of apoplectic (A)
885 and Esca proper-affected (E) 26-year-old standing vines cv. Chardonnay. Results correspond
886 to means \pm S.D. ($n=5$). Results correspond to means \pm S.D. from one representative ($n_{2011}=5$)
887 out of two. Columns headed by the same letter are not significantly different (Dunn's multiple
888 comparison test, $P<0.05$).

889

890 Supplemental Fig 1: Structures of studied stilbenic compounds: **1**: *trans*-piceid and *trans*-
891 resveratrol, **2**: *trans*- ϵ -viniferin, **3**: *trans*-vitisin A, **4**: *trans*-vitisin B.

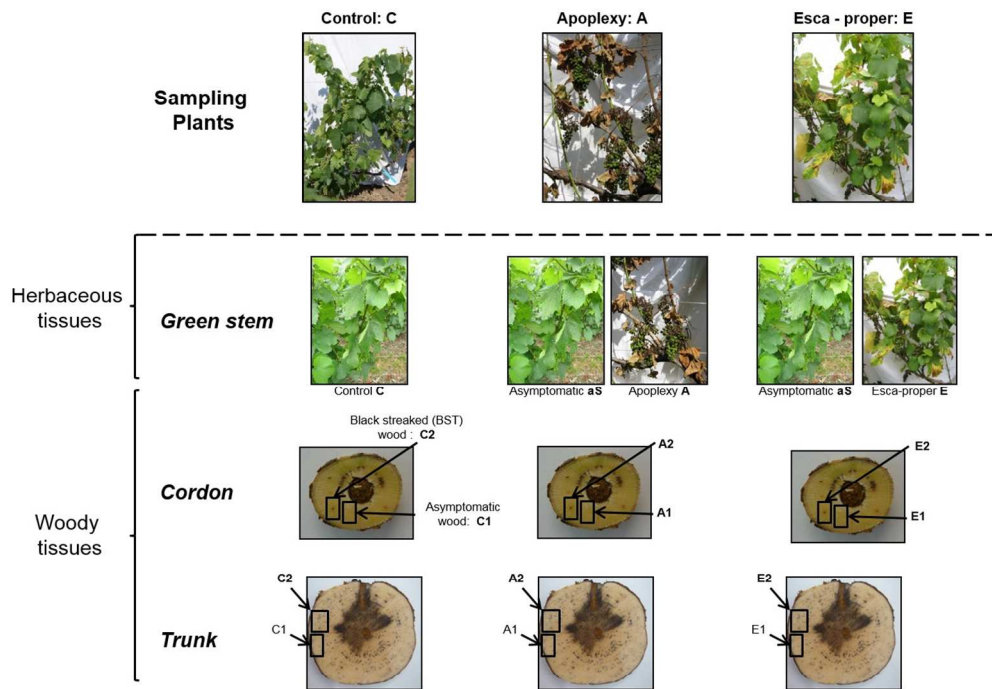


Fig. 1, Magnin-Robert, Phytopathology

Fig 1
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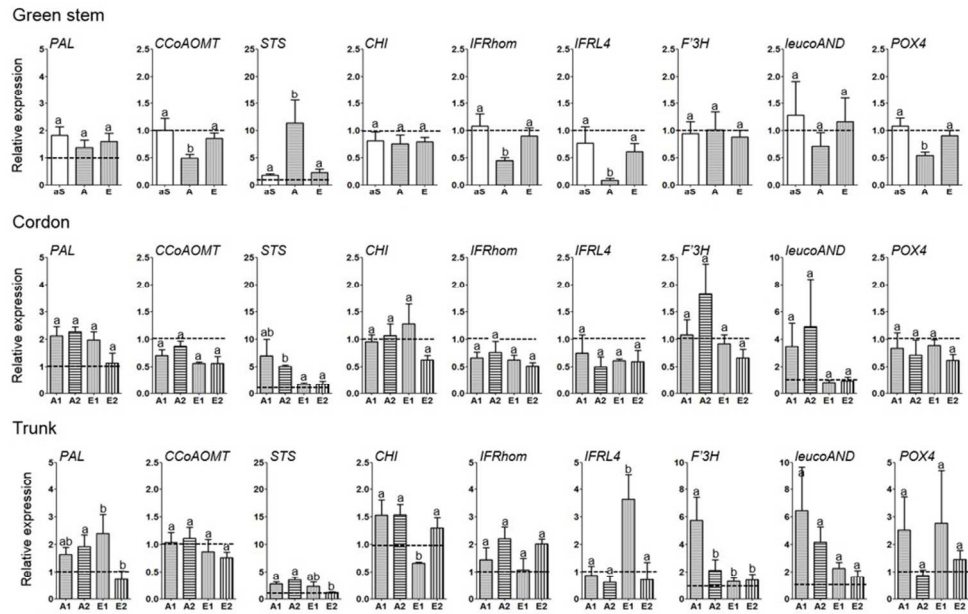


Fig. 2, Magnin-Robert, *Phytopathology*

Fig 2
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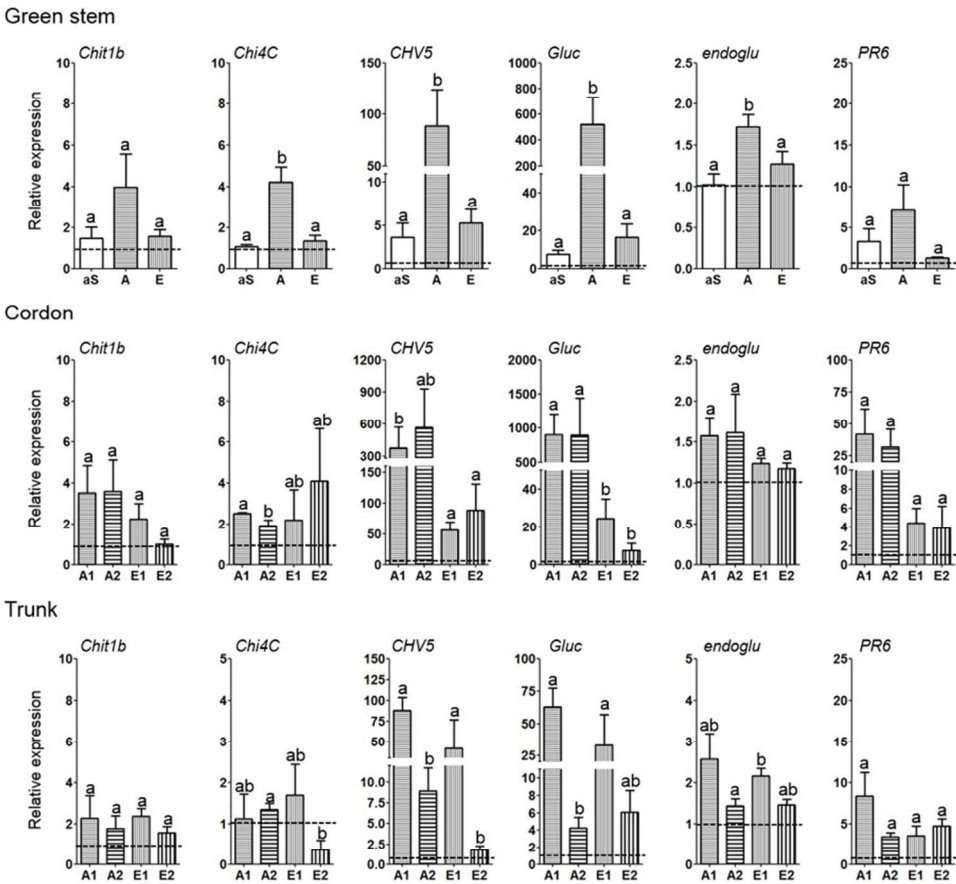
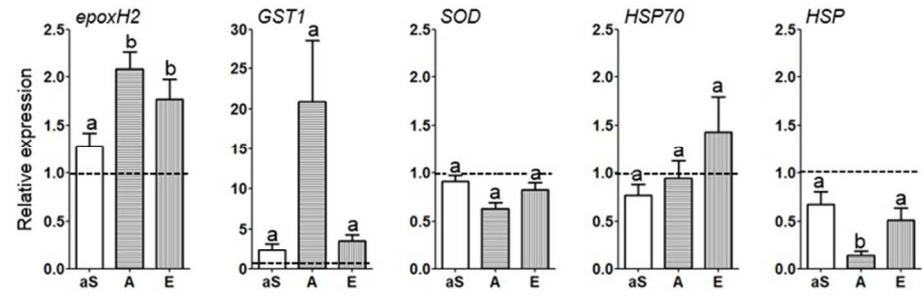


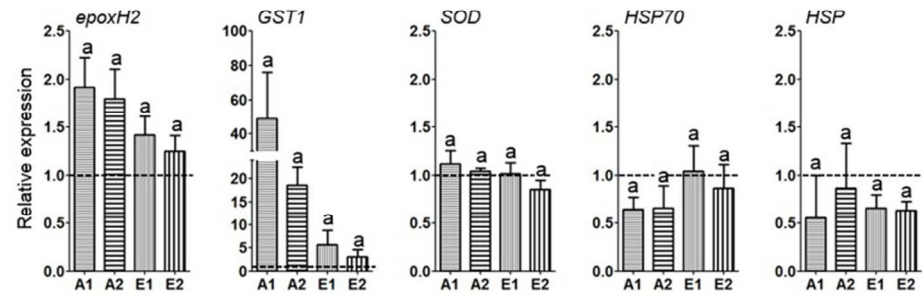
Fig. 3, Magnin-Robert, *Phytopathology*

Fig 3
160x153mm (150 x 150 DPI)

Green stem



Cordon



Trunk

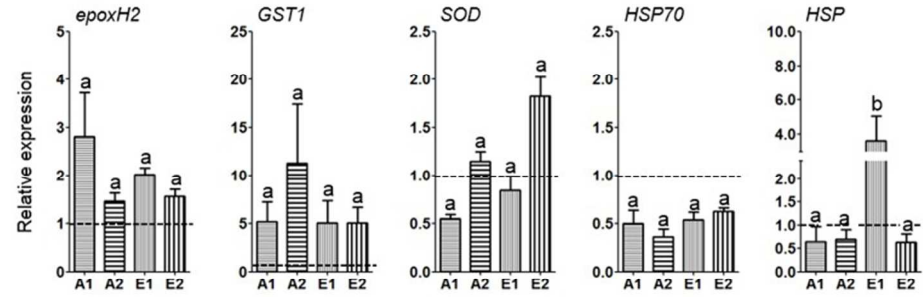


Fig. 4, Magnin-Robert, *Phytopathology*

Fig 4
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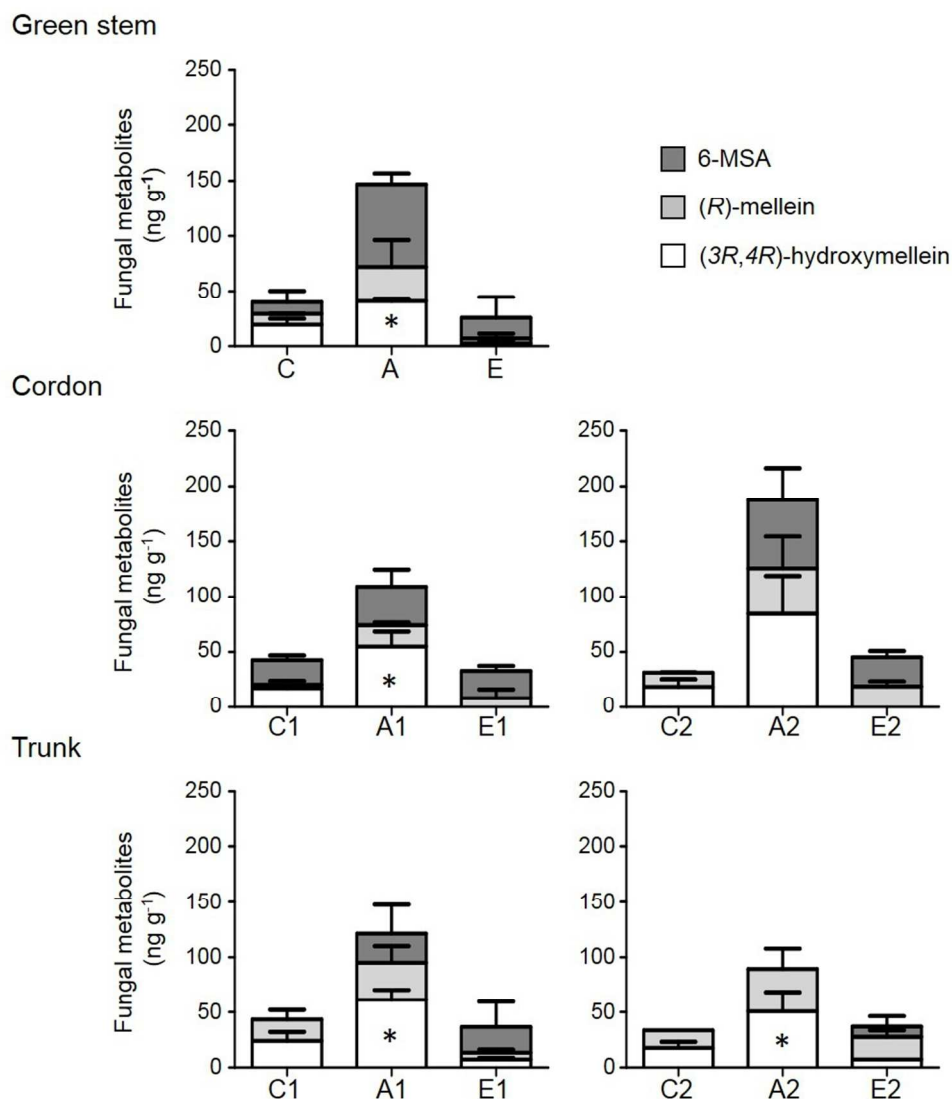
Fig. 5, Magnin-Robert, *Phytopathology*

Fig. 5
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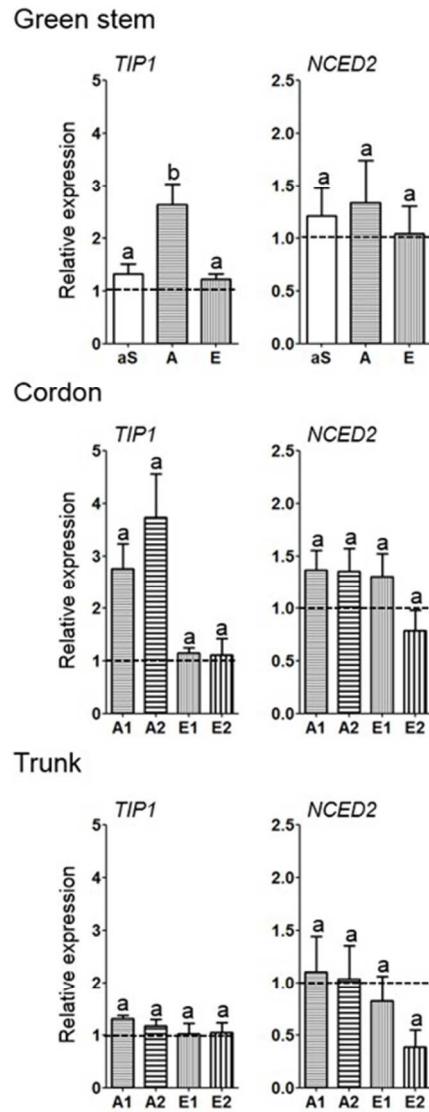


Fig. 6, Magnin-Robert, *Phytopathology*

Fig 6
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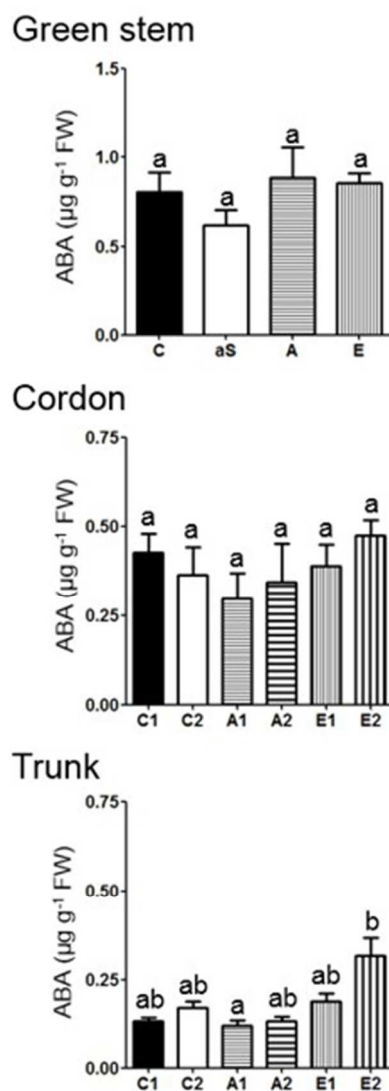
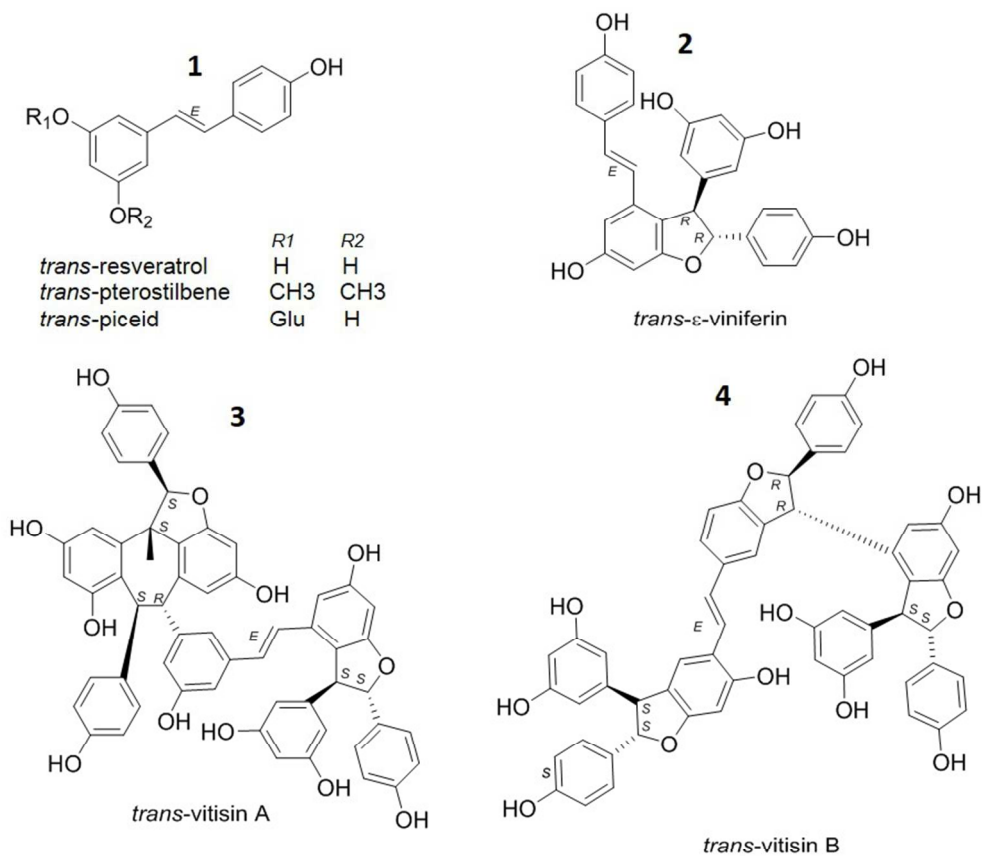


Fig. 7, Magnin-Robert, *Phytopathology*

Fig 7
65x121mm (150 x 150 DPI)



Supplemental Fig. 1, Magnin-Robert, Phytopathology

153x144mm (150 x 150 DPI)