Flowering as the Most Highly Sensitive Period of Grapevine 
(*Vitis vinifera* L. cv Mourvèdre) to the Botryosphaeria Dieback 
Agents *Neofusicoccum parvum* and *Diplodia seriata* Infection

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Abstract: Botryosphaeria dieback is a fungal grapevine trunk disease that currently 
represents a threat for viticulture worldwide because of the important economical losses 
due to reduced yield of affected plants and their premature death. *Neofusicoccum parvum* 
and *Diplodia seriata* are among the causal agents. Vine green stems were artificially 
infected with *N. parvum* or *D. seriata* at the onset of three different phenological stages 
(G stage (separated clusters), flowering and veraison). Highest mean lesion lengths were 
recorded at flowering. Major proteome changes associated to artificial infections during 
the three different phenological stages were also reported using two dimensional gel 
electrophoresis (2D)-based analysis. Twenty (G stage), 15 (flowering) and 13 (veraison) 
differentially expressed protein spots were subjected to nanoLC-MS/MS and a total of 247,
54 and 25 proteins were respectively identified. At flowering, a weaker response to the infection was likely activated as compared to the other stages, and some defense-related proteins were even down regulated (e.g., superoxide dismutase, major latex-like protein, and pathogenesis related protein 10). Globally, the flowering period seemed to represent the period of highest sensitivity of grapevine to Botryosphaeria dieback agent infection, possibly being related to the high metabolic activity in the inflorescences.

**Keywords:** Botryosphaeria dieback; *Neofusicoccum parvum*; *Diplodia seriata*; plant proteomics; two dimensional gel electrophoresis; defense-related proteins

### 1. Introduction

Species in the Botryosphaeriaceae [1,2] have a cosmopolitan distribution and occur as endophytes or pathogens for a wide range of annual and perennial hosts [3]. To date, at least 30 Botryosphaeriaceae species are known to infect grapevine [4–9]. Symptoms of Botryosphaeria dieback consist in woody tissues of grey sectorial necrosis with a brown stripe under the bark as well as typical foliar discolorations in white and red cultivars or wilting leaves in some cases [10,11]. *Diplodia seriata* De Not. 1842 and *Neofusicoccum parvum* [1] are among the Botryosphaeria dieback agents more commonly isolated from grapevine-growing regions worldwide ([11] and references therein). Pathogenicity studies indicate that *N. parvum* is among the most virulent Botryosphaeriaceae species to grapevine while *D. seriata* is ranked among those moderately virulent ([7,12,13] and references therein).

Little information is available about the life cycle of Botryosphaeriaceae. The principal sources of inoculum, pycnidia, are located on infected wood, old pruning wounds, and on pruning canes [14]. These fungi are also present at the surface of different organs like canes [15]. The airborne inoculum is present especially during rainfall [16,17] with peak release during the vegetative growth period, especially in France as described by Larignon and Dubos [18] and Kuntzmann et al. [19]. Their manner of penetration remains unclear but the most obvious seems to be through pruning wounds in the vineyard [20], notably for *D. seriata* (Ds) and *N. parvum* (Np). In French vineyards, it was shown that *D. seriata* contaminated pruning wounds more often after the bleeding when the mean temperature was above 10 °C and in the presence of rainfall [21]. In these conditions, the susceptibility of pruning wounds was at least 8 weeks. In Italy, wound susceptibility was shown to remain high for up to 4 months after pruning, even in late spring when vines were bleeding [22]. Moreover, fresh wound susceptibility was shown to be greater in spring than in winter. Other wounds caused in vegetative growth period (i.e., removal of lateral shoots and desuckering) or by climatic events such as strong wind and hail, could be additional means of fungal penetration inside the vine.

Botryosphaeria dieback is one of the major fungal grapevine trunk diseases which represent a threat for vineyards worldwide due to the decreased production of affected plants and their premature death. Therefore, studying the impact of this disease on grapevine physiology represents a key step towards a better knowledge of symptom development leading to possible control strategies. Phytopathogen infection leads to changes in secondary metabolism based on the induction of a defense programme, as
well as to changes in primary metabolism which affect the growth and development of the plant [23]. At the main crop growth stages such as the flowering, the beginning of fruit growth and then their maturity, the physiological status of the plant may be taken into account to elucidate the fine-tuned infection mechanisms. Responsiveness of grapevine infection by some pathogens such as Botrytis cinerea at both flowering and berry ripening may originate from a disruption of defense responses and sugar metabolism [24,25].

In this paper, the differential sensitivity of 15-year-old standing vines cv. Mourvèdre to the Botryosphaeria dieback agent infection depending on the phenological stage was assessed by artificial infections with N. parvum or D. seriata during three vegetative periods (G stage (separated clusters), flowering and veraison). Moreover, in order to gain a better knowledge on the impact of these pathogens on grapevine physiology, major proteome changes in green stems artificially infected were investigated by using a two dimensional gel electrophoresis (2D)-based approach.

2. Results and Discussion

2.1. Pathogenicity Tests

The causal association between lesion development and fungal strains was confirmed by the re-isolation from the edge of the lesions. No fungi were isolated from the lesion of control stems. Highest mean lesion length (39.6 ± 9.1 mm) was associated to the Np infection performed at the onset of the flowering (Figure 1). Mean lesion length associated to the Ds infection in the same period was 14.3 ± 3.9 mm while in the case of the control was 2.5 ± 1.1 mm. Lowest mean lesion lengths were registered for the G stage; 0.6 ± 0.2, 1.0 ± 0.3 and 0.7 ± 0.2 mm were measured for the control, Ds and Np treatments, respectively. Intermediate values of lesion length were registered for the control and Np treatments performed at the onset of the veraison, 1.0 ± 0.3 and 18.1 ± 4.0 mm, respectively. Surprisingly, with a size of 16.6 ± 3.6 mm registered at veraison, the necrosis length provoked by Ds was very close to that observed on stem infected with the same fungus at the flowering stage (Figure 1B,C). These results confirm that the degree of the virulence of Np and Ds is different ([7,12,13] and references therein) and probably not based on the same factors. Necrosis development may be influenced by the virulence of the fungal strain in relation to the plant phenological stage. Thus, if the development of lesions is regarded as the expression of the pathogenic potential of these fungi, this result may represent a further indication of the high sensitivity of grapevine to stresses especially during the flowering stage [24]. In this way, several authors hypothesized that high susceptibility of grapevine flowers to biotic stress (necrotrophic fungus Botrytis cinerea), may be partially related to their poor ability to carry out the induction of efficient defenses (stilbenic compounds, pathogenesis-related protein) [25,26]. The low responsiveness of flowers to activate the defense responses could be explained by the induction of various mechanisms during flowering period, like fertilization, end of pollen maturation and transition in the whole plant physiology, since the carbohydrate source originating from root and trunk reserves is progressively replaced by photosynthesis in the leaves [24,27,28]. Therefore, the concentrations of sucrose, glucose and fructose, the major sugars of both xylem and phloem saps, are important during the flowering [24]. These carbon sources may represent an attractive nutrient for the fungi. A range of
polysaccharide-degrading enzymes and glycosidases were constitutively produced in vitro on a simple carbon source like D-glucose by phytopathogenic fungi [29]. Therefore, high production of hydrolytic enzymes induced by high concentration of sugars in the sap, may also explain the highest lesion length observed in stems in response to artificial infection with Ds or Np. Lastly, lesions associated to infections performed at the onset of the G stage did not evolve during the flowering, indicating that the activation of physical and chemical defenses by the plant during the G stage efficiently limited the fungal colonization even during the flowering.

**Figure 1.** Mean lesion lengths ± SE on green stems after artificial infection with *N. parvum* or *D. seriata* at the onset of the G stage (A); flowering (B) and veraison (C). Control stems were wounded and inoculated with sterile malt agar. Differences among the means were evaluated by the Dunn’s Multiple Comparison Test after that the null hypothesis (equal means) was rejected in the Kruskal–Wallis test, assuming a significance of $p \leq 0.05$. Statistically significant differences between two conditions are indicated by an asterisk.
2.2. Two-Dimensional Gel Electrophoresis (2D) Analysis

2.2.1. Identification of Differentially Expressed Protein Spots by nanoLC-MS/MS Analysis

The highest average number of total spots detected in gels concerned the G stage (139) while 99 and 115 spots were recorded for the flowering and the veraison stages, respectively (Figure 2A). The lowest total number of spots detected for the flowering stage could result from a physiological decrease of metabolic activities in the green stem as a consequence of their increasing in the inflorescences. The flowering stage corresponds to an important transition between two distinct and successive phases on sugar physiology during the grapevine cycle. Phase one corresponds to the mobilization of starch from woody organs which supplies the annual organs with carbohydrates during their early growth. Phase two coincides with net leaf photosynthesis which supports both the continuation of annual organs growth and the replenishment of reserves [24].

**Figure 2.** Number of total spots detected on 2D gels from G stage, flowering and veraison including means (mean of three biological replicates) from all the groups in each stage (A); Number of total spots detected on 2D gels of each group from G stage (B); flowering (C) and veraison (D). Differences among the means were evaluated by the Dunn’s Multiple Comparison Test after that the null hypothesis (equal means) was rejected in the Kruskal-Wallis test, assuming a significance of $p \leq 0.05$. Statistically significant differences between two conditions are indicated by an asterisk.
**Figure 3.** Map of the identified protein spots quantitatively differentially expressed in the green stems of control or infected plants by *Neofusicoccum parvum* (Np) or *Diplodia seriata* (Ds) at G stage (A); flowering (B) and veraison (C). Isoelectric focusing (IEF) was performed on precast dry polyacrylamide 7 cm length gels ReadyStrip IPG (pH 4–7). The relative molecular mass (kDa) was calibrated with standard protein markers (Prestained SDS-PAGE Standards, Bio-Rad, Hercules, CA, USA) after co-second dimensional electrophoresis. Only spots detected in at least two biological replicates were chosen for identification (indicated with a square). Spots that were not detected in any gel of a given group are indicated with a circle.

When an in-stage comparison was performed, no statistically significant differences related to the treatment were recorded for any stage (Figure 2B–D). The difference of the number of spots detected was likely related to the stage but not to treatments. Therefore, the number of spots chosen for nanoLC-MS/MS analysis was not related to the average of total spots detected. In fact, we selected 20, 15 and 13 spots for G, flowering and veraison stages, respectively (Figure 3). This could indicate that proteome changes linked to the treatments (C2, DsG, NpG) depend on the phenological stage as they were more pronounced during the G stage. In detail, very few proteins are found to be common to the various stages of development. Only four of the differentially expressed proteins identified in stems were observed in two or three stages. Among them, three were identified in the stem at both G stage and flowering (isoflavone reductase, pathogenesis-related protein 10 (PRP-10) and chaperonin...
CPN60-2) (Tables 1 and 2), one identified at G stage and veraison (peroxidase 12-like) and one was identified during the three stages studied (l-ascorbate peroxidase 2, ascoPOX2) (Tables 1–3). These results were in accordance with an interesting work on the genome-wide transcriptomic atlas of grapevine, wherein the authors explained that the majority of the genes expressed in berries, tendrils or stems were more common among different organs than at different developmental stages in the same organ [30].

Protein spot abundance in stems infected with Np or Ds was similar and differences apparently related to the specific inoculum were therefore more pronounced in samples from the G stage (Figure 4). Changes occurred not only in stems infected with Np or Ds but also in C2 (stem wounded but non infected), especially at the veraison stage.

**Figure 4.** Pairwise comparison for all the possible pairs of group samples in one spot performed for all the identified spots in each stage. G stage (A); flowering (B), veraison (C). The number of spots where their relative expression was considered to be similar (ratio \( \leq |2| \)) or dissimilar (ratio \( \geq |2| \)) is reported.
The pairwise comparisons C2/Ds, C2/Np and Ds/Np (for stages G, F and V) showed that specific changes were observed in each treatment but were also common to two or three indicating that changes related to the wounding as well as to the specific inocula occurred at the same time.

2.2.2. Differentially Expressed Proteins among the Treatments

The nanoLC-MS/MS analysis allowed the identification of 247, 54 and 25 differentially expressed total proteins from the G, flowering and veraison stages, respectively (Table S1). A selection of these proteins is listed in Tables 1–3.

G Stage (Separated Clusters)

No down accumulation was observed in the samples from the G stage (C2G, DsG and NpG) as compared to C1G. Some qualitative differences were detected only in the case of the G stage, the spot s5526 was not detected in C1G. Among the proteins therein identified were a eukaryotic initiation factor, a protein disulfide isomerase, and a peroxidase 12-like. As shown in Table 1, a group of proteins involved in primary amino-acid metabolism was increased by wounding and/or fungal infection only during G stage (s6509, s7615, s5720, s2327). These metabolisms did not seem to be activated by the same treatments at flowering or veraison stages (Tables 2 and 3). The most notable protein of this group was the $\gamma$-amino-butyric acid (GABA) biosynthetic protein glutamate decarboxylase-like (s6509, Table 1), already observed to be accumulated in trunk of grapevine “Chardonnay” affected by Botryosphaeria dieback [31]. While many studies corroborate the link between primary metabolism and plant defense reactions [32,33], the role of primary amino acid metabolism in modulation of defense responses by the host remains scarce. In this sense, glutamate metabolism is known to play an important role in plant amino acid metabolism, orchestrating metabolic functions such as providing both C skeletons and $\alpha$-amino groups for the biosynthesis of amino acids with key roles in plant defenses: GABA, arginine or proline [34]. Classified in the methionine synthesis pathway, an adenosylhomocysteine isoform 1 (s6509–s7516) was identified in stems and over-accumulated in samples infected by fungi (DsG and NpG). Also associated with the methionine synthesis pathway, S-adenosylmethionine (SAM) synthase isoform 2 was abundant in C2G, DsG and NpG (s5526–s6509) (Table 1), as recently reported in green stem and trunk of apoplectic or esca proper-affected grapevine “Chardonnay” [35,36]. SAM synthase leads to the biosynthesis of SAM, which can be metabolised via various pathways, leading to ethylene, phenylpropanoids and polyamines synthesis, important molecules of plant defense responses [37,38]. In this sense, a putative role of the SAM synthase in the intrinsic resistance capability against Ds and Np may be suggested as well as in Erysiphe necator- and Plasmopara viticola-resistant grapevine cultivar (“Regent”) [39].

Numerous studies highlight that the rate of photosynthesis is reduced locally in response to pathogens [32]. However, in G stage, three proteins involved in photosynthesis were up-regulated in response to fungal infection (Table 1), oxygen-evolving enhancer protein 1 (s1128–s8717), ribulose biphosphate carboxylase (Rubisco) large chain (s7516) and Rubisco large subunit-binding protein subunit alpha (s2629). In grapevine leaves, similar induction of responses related to energy photosynthesis was observed at gene levels, in response to the vascular ascomycete Eutypa lata and it
appeared to be linked to lack of symptom development on leafy shoots of infected grapevine [40]. At this point, it is reasonable to assume that the rate of photosynthesis could increase to supply the carbon skeletons, energy, and reducing equivalents required to support effective plant defense [33]. Under stress conditions, the respiration process could be enhanced as suggested by increased abundance of glycolysis-related proteins; 2,3-biphosphoglycerate-independent phosphoglycerate mutase (s6624, s5628, s6626) and phosphoglucomutase cytoplasmic (s6624) by wounding, Np and Ds; enolase (s6509) by Np and Ds; and Uridine 5'-diphosphate-glucose dehydrogenase (UDP-Glc DH, s7516, by Ds and Np) from the pentose-phosphate pathway (Table 1). Enolase, an integral enzyme in glycolysis that catalyzes the interconversion of 2-phosphoglycerate to phosphoenolpyruvate, has been found to be responsive in grapevine to phytoplasma infection [41]. Up-regulation of catalyzing enzymes might also increase the production of energy, which is needed in response to Ds and Np infection. Additional studies on UDP-Glc DH indicate that the oxidation of UDP-Glc may have a significant role in increasing the pool of UDP-sugars to supply the demand for increased matrix polysaccharide and cellulose synthesis of structural polysaccharides in plants [42,43]. Moreover, accumulation of GDP-mannose 3,5 epimerase (s6426) and beta-xylosidase/alpha-L-arabinofuransidase (s6626) in stems infected by Np confirmed the activation of cell-wall biosynthesis/modification [44–46].

The significant increase in the abundance of proteins related to active oxygen species (AOS), such as ascoPOX2 (s7220), catalase (s7516), and 2-cys-peroxiredoxin (s1128), indicates the induction of an oxidative stress in response to both wounding and fungal infection in stems at G stage (Table 1). In agreement with our observation, most studies have also reported the induction of an antioxidant system (through catalytic activity, proteins expression or transcripts accumulation) on grapevine organs affected by trunk diseases (leaves [40,47–49]; green stem [35]; wood [36]). By scavenging the AOS formations, the plant’s antioxidant system protects against toxic oxygen intermediates [50]. Proteins involved in defense responses (s6024) were over-regulated in both NpG and DsG (PRP-10 and a universal stress protein-1). This is supported by the increase of PR protein 10 gene expression noted in grapevine cell cultures in response to elicitors produced by Phaeomoniella chlamydospora, a fungal agent implicated in esca proper [51]. Universal stress proteins are mediated by defense-related hormone ethylene [52] and gene expression is known to be up-regulated in response to fungal infection [53].

Instead, proteins involved in protein synthesis (60 acidic ribosomal protein P0: s1231, and elongation factor 1-beta 1: s1231, s2327) and in signal transduction (14-3-3 protein: s1231, and 14-3-3 protein 7: s2327) were accumulated especially in DsG (Table 1). The 14-3-3 protein was similarly accumulated in brown striped wood of Botryosphaeria dieback grapevine [31]; this protein is known to function as a regulator of a wide range of target proteins in all eukaryotes, and to accumulate in response to abiotic or biotic stresses in plants [54]. The NpG treatment showed the highest number of proteins over-regulated, 22 out of the 63 proteins (Table 1); this suggests that grapevine metabolism was deeply altered by Np infection. In this sense, the up-regulation of various heat shock proteins (mitochondrial heat shock 70 kDa protein—s5628, heat shock cognate 70 kDa protein isoform 2-2625 and stromal 70 kDa heat shock-related protein—s2625) and a chaperonin CPN60-2 (s4623) were only observed in stems infected by Np. HSPs/chaperonins were involved in protein folding, assembly translocation and degradation, therefore playing a pivotal role in protecting plants against various stresses [55,56]. Herein, the post-translation processes may also be involved in stress resistance.
Moreover, another HSP cognate 70 kDa (with 96% of identity with HSP cognate 70 kDa, s2625) was recently shown to have its expression and phosphorylation levels upregulated in grapevine leaves infected by phytoplasma. During interaction between phytoplasma and apple tree leaves, HSP cognate 70 kDa was found to be necessary in establishing basal expression levels of several abscisic acid (ABA)-responsive genes, suggesting that this chaperone might also be involved in plant stress hormone ABA signalling events [57].

Flowering

Unlike what was observed in the case of green stems infected (Ds or Np) or only wounded (C2) during the G stage, changes that occurred during the flowering consisted of both down- and over-regulation (Table 2). Nearly 50% of the differentially-accumulated proteins were down-regulated during infection (especially with Ds), possibly reflecting the exploitation of cellular resources and/or the suppression of defense responses [58]. A stem-specific protein (SSP) involved in defense and cell rescue, as well as some proteins involved in protein synthesis, glycolysis, photosynthesis, vitamins or nitrogen metabolisms were down regulated in stems infected with Ds or Np. In regard to their low abundances, we can hypothesize a role of these proteins to make a global response insufficient to avoid development of necrosis. For the SSP (s5217), a similar decrease of protein abundance was already observed in the asymptomatic wood of apoplexy-affected grapevine [36]. Only one protein involved in responses to oxidative stresses, a thioredoxin H-type (s3015), was over-regulated in response to wounding (C2F) or fungal infection (DsF and NpF), while the abundance of a superoxide dismutase (SOD, s5018) decreased in the stem infected by Ds. Reduced expression of SOD was also reported in other organs of the vine affected by trunk diseases, green stems [35], leaves [48,49]) and trunk [36]. Similarly to the G stage, PRP-10 (s5016) and HSPs (s3014, s2123) abundance increased in the stem in response to fungal infection, particularly with Ds (Table 2). In accordance with the role of these proteins in plant tolerance against various stresses, the lack of their strong induction in stems infected by Np could explain the longest lesions caused by Np (Figure 1). Moreover, the HSPs accumulated in response to fungal infection are different (18.2 kDa class I HSP isoform 1, 18.2 kDa class I HSP, mitochondrial 23.6 kDa HSP; Table 2) from those accumulated during G stage (70 kDa proteins, Table 1). These results suggest that plant responses to fungal infection may vary significantly with the phenological plant stage. Nevertheless, during the G stage and flowering period, the abundance of isoflavone reductase, a protein involved in secondary metabolism, was increased by two fold in stems infected with Np (Table 1—s5628 and Table 2—s2409). By comparison with the other perturbed metabolisms, the typical secondary metabolism of the vine, represented by the phenylpropanoid pathway, seems to be similarly regulated in response to fungal infection irrespective of the developmental stage (G or F).

Two proteins of the primary metabolism, cytosolic triosephosphate isomerase (TIP, s6222) and pyruvate dehydrogenase (s2409), were among those over-accumulated mainly in NpF. The TPI allows the reversible isomerization between D-glyceraldehyde-3-phosphate (G3P) and dihydroxyacetone phosphate. Note that the G3P is used in the glycolytic pathway to ultimately give pyruvate. At the end of glycolysis, the generated pyruvate will be subsequently decarboxylated and will react with coenzyme A via the action of the pyruvate dehydrogenase, to give acetyl coenzyme A, the entry point
of the Krebs cycle. Accumulation of the proteins associated with glycolysis and the Krebs cycle may support cellular energy requirements for plant defense reactions in response to Np infection [32,33]. Despite their accumulation of proteins, significant necrosis is measured on the stems inoculated with Np which suggests no effect of these proteins in limiting necrosis development.

Veraison

Similarly to the other two stages, HSP was over-accumulated during veraison. In addition, a chloroplastic small HSP (s1102–s1103–s1105) was abundant in stems C2V, DsV and NpV (Table 3). In response to fungal infection, a SAM synthase 5 (s5105), a 22.0 kDa HSP (s5105) and a ascoPOX2 (s6309) protein were over-accumulated in DsV and NpV. Considering the expression of these three proteins, it is obvious that their change in the stem has no effective influence in limiting necrosis development. Still, a glutathione S-transferase F9 (GSTF9, s2204) was among those over-regulated in both C2V and DsV but not in NpV, while a peroxidase 12-like (s4703) was over-accumulated solely in DsV. Nevertheless, the lesion lengths observed on green stems infected by Ds or Np are very similar and suggest that the accumulation of both proteins does not stop necrosis development. For the GST, various studies proposed that GST system is not related to the plant tolerance against pathogenic fungi [36,47]. In this sense, Spagnolo et al. [31] observed no relationship between the up-/down-regulation of GSTF9 and the greater or lesser susceptibility of the grapevine cultivars against Botryosphaeria dieback. Two proteins involved in protein synthesis (s1301, putative transcription factor and nascent polypeptide-associated complex subunit alpha-like) were over- and down-accumulated in C2V and NpV respectively (Table 3). This contrasting response suggests an alteration of the proteins synthesis under infection condition.
Table 1. Identified proteins differentially expressed during the G stage.

| Spot | Ratio to C1G | Matched Protein | Accession Number | Cov. % | M<sub>w</sub> | Functional Category |
|------|--------------|-----------------|------------------|--------|------------|-------------------|
| 5526 | * 9.0 3.2 3.4 | S-adenosylmethionine synthase 2 | gi|23635284 | 40 24.70 | Defense and cell rescue |
| 5526 | * 12.3 4.5 | eukaryotic initiation factor 4A-2 (Vitis vinifera) | gi|22544222 | 46 53.30 | Protein synthesis |
| 5526 | * 6.0 3.4 | protein disulfide isomerase-like 2-3 (Vitis vinifera) | gi|22544717 | 46 32.80 | Protein processing |
| 5526 | * 9.0 3.2 | peroxidase 12-like (Vitis vinifera) | gi|35949319 | 39 33.10 | Defense and cell rescue |
| 1013 | 9.0 4.6 9.5 | actin-depolymerizing factor 2-like isoform 1 (Vitis vinifera) | gi|22543504 | 16 13.70 | Cytoskeleton |
| 1128 | 3.0 7.6 8.3 | 2-Cys peroxiredoxin (Vitis vinifera) | gi|14778975 | 30 52.70 | Defense and cell rescue |
| 1128 | 3.0 7.6 8.3 | oxygen-evolving enhancer protein 1, chloroplastic (Vitis vinifera) | gi|14779185 | 33 20.50 | Photosynthesis |
| 1231 | 4.1 12.3 4.5 | 60S acidic ribosomal protein P0 (Vitis vinifera) | gi|14784326 | 34 12.50 | Protein synthesis |
| 1231 | 4.1 12.3 4.5 | elongation factor 1-beta 1 (Vitis vinifera) | gi|29608391 | 27 38.30 | Protein synthesis |
| 1231 | 4.1 12.3 4.5 | 14-3-3 protein (Vitis vinifera) | gi|35949288 | 29 66.20 | Signal transduction |
| 6624 | 2.4 2.6 4.5 | phosphoglucomutase, cytoplasmic (Vitis vinifera) | gi|22542431 | 63 26.60 | Glycolysis/Gluconeogenesis |
| 6624 | 2.4 2.6 4.5 | succinate dehydrogenase (ubiquinone) flavoprotein subunit 1, mitochondrial isoform 1 (Vitis vinifera) | gi|22543077 | 73 38.80 | Citrate cycle |
| 6624 | 2.4 2.6 4.5 | 2,3-bisphosphoglycerate-independent phosphoglycerate mutase isoform 1 (Vitis vinifera) | gi|22543906 | 61 24.50 | Glycolysis/Gluconeogenesis |
| 6736 | 3.2 3.3 3.4 | eukaryotic initiation factor 4A-2 (Vitis vinifera) | gi|22544222 | 46 22.80 | Protein synthesis |
| 6736 | 3.2 3.3 3.4 | phospholipase D alpha 1 (Vitis vinifera) | gi|22542498 | 92 41.90 | Glycan metabolism |
| 6736 | 3.2 3.3 3.4 | ATP-dependent Clp protease ATP-binding subunit clpA homolog CD4A, chloroplastic-like (Vitis vinifera) | gi|22545647 | 102 46.50 | Cell growth and death |
| 8717 | 3.0 2.0 2.8 | oxygen-evolving enhancer protein 1, chloroplastic (Vitis vinifera) | gi|14779185 | 33 36.60 | Photosynthesis |
| 8717 | 3.0 2.0 2.8 | elongation factor 2-like isoform 1 (Vitis vinifera) | gi|22546216 | 93 44.70 | Protein synthesis |
| 6509 | 1.1 2.1 2.5 | S-adenosylmethionine synthase 2 | gi|22363528 | 43 44.00 | Defense and cell rescue |
| 6509 | 1.1 2.1 2.5 | UDP-glucose 6-dehydrogenase-like isoform 1 (Vitis vinifera) | gi|22542350 | 52 23.70 | Pentose phosphate cycle |
| 6509 | 1.1 2.1 2.5 | adenosylhomocysteinase isoform 1 (Vitis vinifera) | gi|22543350 | 53 30.50 | Amino acid metabolism |
| 6509 | 1.1 2.1 2.5 | enolase 1 (Vitis vinifera) | gi|22544100 | 47 56.80 | Glycolysis/Gluconeogenesis |
| Spot | Ratio to C1G | Matched Protein | Accession Number | Cov. % | M<sub>w</sub> | Functional Category |
|------|--------------|-----------------|------------------|--------|------------|---------------------|
| 6509 | 1.1          | glutamate decarboxylase-like (*Vitis vinifera*) | gi|225462892 | 57 | 25.30 | Amino acid metabolism |
| 6024 | 1.3          | pathogenesis-related protein 10 (Vitis hybrid cultivar) | gi|163914213 | 17 | 49.40 | Defense and cell rescue |
| 6024 | 1.3          | universal stress protein A-like protein isoform 1 (*Vitis vinifera*) | gi|225431940 | 18 | 57.60 | Defense and cell rescue |
| 6024 | 1.3          | eukaryotic translation initiation factor 5A (*Vitis vinifera*) | gi|225468027 | 17 | 28.10 | Protein synthesis |
| 7516 | 1.2          | Ribulose bisphosphate carboxylase large chain (*Vitis vinifera*) | gi|134034997 | 6 | 40.00 | Photosynthesis |
| 7516 | 1.2          | catalase (*Vitis vinifera*) | gi|19070130 | 56 | 12.20 | Glyoxylate and dicarboxylate metabolism |
| 7516 | 1.2          | ATPase subunit 1 (*Vitis vinifera*) | gi|224365668 | 55 | 19.30 | Metabolism and energy |
| 7516 | 1.2          | UDP-glucose 6-dehydrogenase-like isoform 1 (*Vitis vinifera*) | gi|225423507 | 52 | 24.20 | Pentose phosphate cycle |
| 7516 | 1.2          | adenosylhomocysteinase isoform 1 (*Vitis vinifera*) | gi|225433506 | 53 | 38.10 | Amino acid metabolism |
| 7516 | 1.2          | enolase 1 (*Vitis vinifera*) | gi|225441000 | 47 | 40.50 | Glycolysis/Gluconeogenesis |
| 5720 | 2.6          | phospholipase D alpha 1 (*Vitis vinifera*) | gi|225442981 | 92 | 25.50 | Glycer metabolism |
| 5720 | 2.6          | aminopeptidase N-like (*Vitis vinifera*) | gi|359474189 | 101 | 50.60 | Amino acid metabolism |
| 3227 | 2.0          | oxygen-evolving enhancer protein 1, chloroplastic (*Vitis vinifera*) | gi|147791852 | 33 | 57.70 | Photosynthesis |
| 3227 | 2.0          | isoflavone reductase homolog P3 (*Vitis vinifera*) | gi|225458243 | 33 | 34.40 | Secondary metabolism |
| 3227 | 2.0          | putative fructokinase-5-like (*Vitis vinifera*) | gi|225459906 | 34 | 23.20 | Carbohydrate metabolism |
| 2327 | 1.3          | ran-binding protein 1 homolog c (*Vitis vinifera*) | gi|225439378 | 24 | 37.60 | Defense and cell rescue |
| 2327 | 1.3          | 14-3-3 protein 7 (*Vitis vinifera*) | gi|225459292 | 28 | 66.30 | Signal transduction |
| 2327 | 1.3          | elongation factor 1-beta 1 (*Vitis vinifera*) | gi|296083911 | 27 | 55.30 | Photosynthesis |
| 2327 | 1.3          | aspartic proteinase isoform 2 (*Vitis vinifera*) | gi|302144105 | 46 | 23.50 | Amino acid metabolism |
| 7220 | 1.1          | triosephosphate isomerase, cytosolic (*Vitis vinifera*) | gi|147784332 | 27 | 19.30 | Glycolysis/Gluconeogenesis |
| 7220 | 1.1          | L-ascorbate peroxidase 2, cytosolic (*Vitis vinifera*) | gi|225435177 | 27 | 64.80 | Carbohydrate metabolism |
| 7220 | 1.1          | 3-oxoacyl-(acyl-carrier-protein) reductase 1, chloroplastic (*Vitis vinifera*) | gi|225456248 | 34 | 24.80 | Fatty acid metabolism |
| 6426 | 1.3          | eukaryotic initiation factor 4A-3-like (*Vitis vinifera*) | gi|147785805 | 44 | 42.20 | Protein synthesis |
| 6426 | 1.3          | GDP-mannose 3,5-epimerase 1 isoform 1 (*Vitis vinifera*) | gi|147794688 | 42 | 38.30 | Carbohydrate metabolism |
| 6426 | 1.3          | l-adenosylmethionine synthase 2 | gi|223635284 | 43 | 83.00 | Defense and cell rescue |
| Spot | Ratio to C1G | Matched Protein | Accession Number | Cov. % | MW | Functional Category |
|------|-------------|-----------------|-----------------|-------|----|---------------------|
|      | C2G | DsG | NpG |                   |     |    |                     |
| 6426 | 1.3 | 1.3 | 2.4 | alcohol dehydrogenase 1 (Vitis vinifera) | gi225431505 | 41 | 18.40 | Glycolysis/Gluconeogenesis |
| 6426 | 1.3 | 1.3 | 2.4 | isocitrate dehydrogenase (NADP) (Vitis vinifera) | gi225466253 | 42 | 50.00 | Citrate cycle |
| 5628 | 1.0 | 1.1 | 2.3 | heat shock 70 kDa protein, mitochondrial-like (Vitis vinifera) | gi225429228 | 72 | 41.70 | Protein destination |
| 5628 | 1.0 | 1.1 | 2.3 | 2,3-bisphosphoglycerate-independent phosphoglycerate mutase isoform 1 (Vitis vinifera) | gi225439064 | 61 | 22.20 | Glycolysis/Gluconeogenesis |
| 5628 | 1.0 | 1.1 | 2.3 | isoflavone reductase homolog P3 (Vitis vinifera) | gi225458243 | 33 | 14.00 | Secondary metabolism |
| 2625 | 1.3 | 1.5 | 2.1 | luminal-binding protein 5-like (Vitis vinifera) | gi359490716 | 73 | 42.40 | Protein destination |
| 2625 | 1.3 | 1.5 | 2.1 | heat shock cognate 70 kDa protein isoform 2 (Vitis vinifera) | gi359486799 | 75 | 38.60 | Protein destination |
| 2625 | 1.3 | 1.5 | 2.1 | stromal 70 kDa heat shock-related protein, chloroplastic-like (Vitis vinifera) | gi225456004 | 75 | 53.50 | Protein destination |
| 2629 | 1.2 | 0.9 | 2.0 | protein disulfide-isomerase (Vitis vinifera) | gi225459587 | 55 | 51.50 | Protein processing |
| 2629 | 1.2 | 0.9 | 2.0 | ruBisCO large subunit-binding protein subunit alpha, chloroplastic-like (Vitis vinifera) | gi359479362 | 61 | 67.60 | Photosynthesis |
| 6626 | 1.1 | 1.3 | 2.0 | succinate dehydrogenase (ubiquinone) flavoprotein subunit 1, mitochondrial isoform 1 (Vitis vinifera) | gi225430776 | 73 | 12.10 | Citrate cycle |
| 6626 | 1.1 | 1.3 | 2.0 | 2,3-bisphosphoglycerate-independent phosphoglycerate mutase isoform 1 (Vitis vinifera) | gi225439064 | 61 | 64.90 | Glycolysis/Gluconeogenesis |
| 6626 | 1.1 | 1.3 | 2.0 | beta-xyllosidase/alpha-L-arabinofuranosidase 2-like (Vitis vinifera) | gi297745522 | 80 | 16.60 | Carbohydrate metabolism |
| 4623 | 0.7 | 0.6 | 2.0 | d-3-phosphoglycerate dehydrogenase, chloroplastic-like (Vitis vinifera) | gi225428898 | 62 | 10.60 | Glycolysis/Gluconeogenesis |
| 4623 | 0.7 | 0.6 | 2.0 | chaperonin CPN60-2, mitochondrial isoform 1 (Vitis vinifera) | gi225433375 | 61 | 68.70 | Protein destination |
| 4623 | 0.7 | 0.6 | 2.0 | stromal 70 kDa heat shock-related protein, chloroplastic-like (Vitis vinifera) | gi225456004 | 75 | 20.70 | Protein destination |

* spot code as reported in Figure 3; * ratio of spot expression values (relative OD × area%) in C2, Ds and Np stems to the related control (C1G, C1F or C1V). Values indicating over or down expression (ratio ≥ |2|) are highlighted in yellow, respectively. Values were replaced by an asterisk when the spot was not detected in the control; protein identified via the MASCOT and OMSSA search engines against in house made database from NCBI nr database; accession No. of the matched protein as reported in the NCBI database; percentage of the protein sequence covered by the matching peptides; molecular mass (kDa).
Table 2. Identified proteins differentially expressed during flowering.

| Spot  | Ratio to C1F | Matched Protein | Accession Number | Cov. % | MW  | Functional Category |
|-------|--------------|-----------------|------------------|--------|-----|---------------------|
|       | C2F | DSF | NpF |                   |     |     |                     |
| 3015  | 4.56 | 4.69 | 2.95 | thioredoxin H-type isoform 1 (Vitis vinifera) | gi|225458147 | 62 | 12.80 | Protein folding |
| 3014  | 2.80 | 26.5 | 1.78 | 18.2 kDa class I heat shock protein isoform 1 (Vitis vinifera) | gi|225449302 | 46 | 17.02 | Protein destination |
| 2123  | 0.83 | 3.52 | 1.25 | 23.6 kDa heat shock protein, mitochondrial isoform 1 (Vitis vinifera) | gi|225466111 | 42 | 23.74 | Protein destination |
| 1009  | 0.58 | 0.96 | 3.47 | 18.2 kDa class I heat shock protein isoform 1 (Vitis vinifera) | gi|225449302 | 13 | 17.02 | Protein destination |
| 6222  | 1.04 | 0.61 | 3.05 | L-ascorbate peroxidase 2, cytosolic (Vitis vinifera) | gi|225435177 | 65 | 27.56 | Carbohydrate metabolism |
| 6222  | 1.04 | 0.61 | 3.05 | triosephosphate isomerase, cytosolic (Vitis vinifera) | gi|22549541 | 53 | 21.13 | Glycolysis/Gluconeogenesis |
| 2409  | 0.70 | 1.37 | 2.10 | L-galactose dehydrogenase (Vitis vinifera) | gi|146432259 | 47 | 34.64 | Carbohydrate metabolism |
| 2409  | 0.70 | 1.37 | 2.10 | pyruvate dehydrogenase E1 component subunit beta, mitochondrial-like isoform 1 (Vitis vinifera) | gi|225425166 | 42 | 39.49 | Glycolysis/Gluconeogenesis |
| 2409  | 0.70 | 1.37 | 2.10 | PREDICTED: fructokinase-2 (Vitis vinifera) | gi|22543918 | 53 | 35.20 | Carbohydrate metabolism |
| 5016  | 1.08 | 3.31 | 0.46 | pathogenesis-related protein 10 (Vitis hybrid cultivar) | gi|163914213 | 14 | 17.11 | Defense and cell rescue |
| 5016  | 1.08 | 3.31 | 0.46 | putative transcription factor (Vitis vinifera) | gi|22543375 | 45 | 61.37 | Protein destination |
| 4627  | 0.43 | 1.08 | 1.38 | MLP-like protein 34 (Vitis vinifera) | gi|22542477 | 47 | 17.08 | Defense and cell rescue |
| 4627  | 0.43 | 1.08 | 1.38 | ruBisCO large subunit-binding protein subunit beta, chloroplastic (Vitis vinifera) | gi|225435794 | 42 | 64.61 | Photosynthesis |
| 6018  | 1.90 | 0.24 | 1.12 | ubiquitin-conjugating enzyme E2 35 isoform 1 (Vitis vinifera) | gi|225461646 | 51 | 17.22 | Protein degradation |
| 6018  | 1.90 | 0.24 | 1.12 | MLP-like protein 34 (Vitis vinifera) | gi|22542477 | 72 | 17.08 | Defense and cell rescue |
| 6018  | 1.90 | 0.24 | 1.12 | chaperonin CPN60-2, mitochondrial isoform 1 (Vitis vinifera) | gi|22543375 | 27 | 61.37 | Protein degradation |
| 6018  | 1.90 | 0.24 | 1.12 | ubiquitin-conjugating enzyme E2 36 isoform 1 (Vitis vinifera) | gi|225446595 | 38 | 17.22 | Protein degradation |
| 5018  | 1.65 | 0.17 | 0.65 | MLP-like protein 34 (Vitis vinifera) | gi|22542477 | 76 | 17.08 | Defense and cell rescue |
| 5018  | 1.65 | 0.17 | 0.65 | ubiquitin-conjugating enzyme E2 36 isoform 1 (Vitis vinifera) | gi|225446595 | 38 | 17.22 | Protein degradation |
| 5018  | 1.65 | 0.17 | 0.65 | glycine-rich RNA-binding protein GRP1A-like (Vitis vinifera) | gi|359475330 | 17 | 16.33 | Cell growth and death |
| 5018  | 1.65 | 0.17 | 0.65 | superoxide dismutase (Cu-Zn) isoform 2 (Vitis vinifera) | gi|225451120 | 29 | 15.28 | Defense and cell rescue |
| 5216  | 0.73 | 0.50 | 0.42 | putative transcription factor (Vitis vinifera) | gi|14582465 | 31 | 16.70 | Protein synthesis |
## Table 2. Cont.

| Spot  | Ratio to C1F b | Matched Protein c | Accession Number d | Cov. % e | M_W f | Functional Category |
|-------|----------------|-------------------|-------------------|---------|-------|--------------------|
| 5216  | 0.73 0.50 0.42 | ferritin-3, chloroplastic (*Vitis vinifera*) | gi|147784301 | 18 | 25.37 | Metabolism of cofactors and vitamins |
| 5216  | 0.73 0.50 0.42 | triosephosphate isomerase, chloroplastic-like isoform 1 (*Vitis vinifera*) | gi|225427917 | 59 | 34.67 | Glycolysis/Gluconeogenesis |
| 4421  | 0.52 0.47 0.33 | glutelin type-A 1 (*Vitis vinifera*) | gi|225435090 | 44 | 38.33 | - |
| 4421  | 0.52 0.47 0.33 | glutamine synthetase nodule isozyme isoform 1 (*Vitis vinifera*) | gi|225451235 | 19 | 34.37 | Nitrogen metabolism |
| 5217  | 0.61 0.32 0.19 | stem-specific protein TSJT1 (*Vitis vinifera*) | gi|225432548 | 46 | 25.25 | Defense and cell rescue |
| 0207  | 0.20 0.10 0.24 | putative ripening-related protein (*Vitis vinifera*) | gi|7406667 | 23 | 15.39 | - |
| 0309  | 0.35 0.24 0.16 | uncharacterized protein LOC100232885 (*Vitis vinifera*) | gi|225447003 | 78 | 18.41 | - |

a spot code as reported in Figure 3; b ratio of spot expression values (relative OD × area%) in C2, Ds and Np stems to the related control (C1G, C1F or C1V). Values indicating over or down expression (ratio ≥ 2) are highlighted in yellow or grey, respectively. Values were replaced by an asterisk when the spot was not detected in the control; c protein identified via the MASCOT and OMSSA search engines against in house made database from NCBI database; d accession No. of the matched protein as reported in the NCBI database; e percentage of the protein sequence covered by the matching peptides; f molecular mass (kDa).

## Table 3. Identified proteins differentially expressed during the veraison.

| Spot  | Ratio to C1V b | Matched Protein c | Accession Number d | Cov. % e | M_W f | Functional Category |
|-------|----------------|-------------------|-------------------|---------|-------|--------------------|
| 1102  | 5.50 4.35 3.26 | small heat shock protein, chloroplastic (*Vitis vinifera*) | gi|225455238 | 55 | 25.03 | Protein destination |
| 1103  | 12.93 8.10 2.86 | small heat shock protein, chloroplastic (*Vitis vinifera*) | gi|225455238 | 48 | 25.03 | Protein destination |
| 1105  | 33.75 5.75 18.75 | small heat shock protein, chloroplastic (*Vitis vinifera*) | gi|225455238 | 54 | 25.03 | Protein destination |
| 5105  | 1.98 3.02 2.16 | S-adenosylmethionine synthase 5 | gi|223635289 | 10 | 42.79 | Defense and cell rescue |
| 5105  | 1.98 3.02 2.16 | 22.0 kDa heat shock protein (*Vitis vinifera*) | gi|225459900 | 46 | 21.12 | Protein destination |
| 6309  | 1.95 2.16 2.40 | l-ascorbate peroxidase 2, cytosolic (*Vitis vinifera*) | gi|225435177 | 61 | 27.56 | Other carbohydrate metabolism |
| 2204  | 4.13 2.20 1.58 | uncharacterized protein LOC100254632 (*Vitis vinifera*) | gi|225441008 | 34 | 16.79 | - |
| 2204  | 4.13 2.20 1.58 | glutathione S-transferase F9 (*Vitis vinifera*) | gi|225446791 | 29 | 24.91 | Defense and cell rescue |
| 2204  | 4.13 2.20 1.58 | uridylyl kinase isoform 1 (*Vitis vinifera*) | gi|225454048 | 20 | 23.53 | Nucleotide metabolism |
Table 3. Cont.

| Spot | Ratio to C1V<sup>b</sup> | Matched Protein<sup>c</sup> | Accession Number<sup>d</sup> | Cov. %<sup>e</sup> | MW<sup>f</sup> | Functional Category |
|------|-----------------|-------------------------|------------------|----------|----------|-------------------|
| 4703 | 1.31 2.91 0.90 | peroxidase 12-like (<i>Vitis vinifera</i>) | gi|359493149 | 64 | 39.18 | Defense and cell rescue |
| 1301 | 2.15 1.29 0.29 | putative transcription factor (<i>Vitis vinifera</i>) | gi|14582465 | 14 | 16.70 | Protein synthesis |
| 1301 | 2.15 1.29 0.29 | nascent polypeptide-associated complex subunit alpha-like (<i>Vitis vinifera</i>) | gi|225470846 | 42 | 22.03 | Protein synthesis |
| 2709 | 0.00 1.34 0.20 | tubulin alpha-2 chain (<i>Vitis vinifera</i>) | gi|225458970 | 57 | 49.59 | Cytoskeleton |
| 4519 | 0.20 0.70 0.50 | naringenin,2-oxoglutarate 3-dioxygenase (<i>Vitis vinifera</i>) | gi|225431140 | 65 | 40.81 | Secondary metabolism |
| 4519 | 0.20 0.70 0.50 | caffeic acid 3-O-methyltransferase 1-like isofrom 1 (<i>Vitis vinifera</i>) | gi|359490763 | 53 | 39.50 | Secondary metabolism |
| 1304 | 0.44 0.07 0.48 | proteasome subunit alpha type-5 isofrom 1 (<i>Vitis vinifera</i>) | gi|225441985 | 43 | 25.98 | Protein degradation |
| 1402 | 0.12 0.07 0.27 | uncharacterized protein LOC100232885 (<i>Vitis vinifera</i>) | gi|225447003 | 83 | 18.41 | |
| 5602 | 0.03 0.24 0.16 | S-adenosylmethionine synthase 5 | gi|223635289 | 75 | 42.79 | Defense and cell rescue |

<sup>a</sup> spot code as reported in Figure 3; <sup>b</sup> ratio of spot expression values (relative OD × area%) in C2, Ds and Np stems to the related control (C1G, C1F or C1V). Values indicating over or down expression (ratio ≥ |2|) are highlighted in yellow or grey, respectively. Values were replaced by an asterisk when the spot was not detected in the control; <sup>c</sup> protein identified via the MASCOT and OMSSA search engines against in house made database from NCBI database; <sup>d</sup> accession No. of the matched protein as reported in the NCBI database; <sup>e</sup> percentage of the protein sequence covered by the matching peptides; <sup>f</sup> molecular mass (kDa).
Figure 5. The climate trend [minimum, maximum, mean daily temperatures and rainfall (mm)] recorded in the experimental site (Tables 1–3).
3. Experimental Section

3.1. Plant Material, Fungal Strains and Pathogenicity Tests

A 15-year-old vineyard cultivar Mourvèdre/3309 located at Rodilhan (Costières de Nîmes, France) and owned by the Lycée agricole Marie-Durand of Rodilhan was the experimental site. Green stems at the onset of the phenological stage G, flowering and veraison were selected for the study and a total of four treatments were planned: C1, stem non inoculated and non-wounded; C2, stem wounded and inoculated with sterile malt agar plug; Ds, stem inoculated with the D. seriata strain (strain Bo98-1 isolated from symptomatic vines in Pyrénées-Orientales vineyards, France) and Np, stem inoculated with the N. parvum strain (strain Np SV isolated from symptomatic vines in Bouches-du-Rhône, France) (Table 4). Fungal strains were cultivated on 1.5% malt extract agar at 24 °C in the dark. Stems were inoculated at the onset of G stage (3 May 2012), flowering (6 June 2012) and veraison (26 July 2012) (Figure 5) with sterile malt agar plug (control stems) or fungal strains after that a longitudinal wound (8 mm length, 1 mm deep) was performed with a sterile scalpel at level of the third internode. A 5 mm Ø malt agar or mycelial plug from the edge of a 5 days old actively growing fungal culture was then put into the wound and protected with parafilm. A total of 8 repetitions per treatment and one repetition per plant were performed. Observation of lesion development and reisolation tests (5 out of 8 repetitions) were performed on 16 October 2012 for C2, Ds and Np treatments from all phenological stages as described by Larignon and Dubos [59]. Samples for protein extraction (3 out of 8 repetitions), consisting in the inoculated internode (C2, Ds and Np) as well as of the correspondent non wounded internode of C1 stems were collected 20 days after the inoculation. Samples were frozen in the field with liquid nitrogen and subsequently stored at −80 °C. Before protein extraction, the amount of biological sample needed was ground to a fine powder in liquid nitrogen with a Mixer Mill MM 400 (Retsch, Haan, Germany).

Table 4. Description of sample codes.

| Condition          | Phenological Stage |
|--------------------|--------------------|
|                    | G stage | Flowering | Veraison |
| Control 1          | C1G     | C1F       | C1V      |
| Control 2          | C2G     | C2F       | C2V      |
| D. seriata strain Bo98-1 | DsG     | DsF       | DsV      |
| N. parvum strain Np SV | NpG     | NpF       | NpV      |

Example: C1G—(non-wounded stem-G stage); C2G—(stem wounded but inoculated with sterile agar-G stage); DsG—(D. seriata strain Bo98-1-G stage); NpG—(N. parvum strain Np SV -G stage).

3.2. Two-Dimensional Gel Electrophoresis (2D) Analysis

3.2.1. Protein Extraction

Total protein fraction of green stems and related two-Dimensional Gel Electrophoresis (2D) analysis were performed according to Magnin-Robert et al. [36].
3.2.2. Image Analysis

Digitized images at 36.6 μm resolution were obtained using the GS-800 scanner and Quantity One 4.6.2 software (Bio-Rad, Hercules, CA, USA). Computerized 2D gel analysis, including spot detection and quantification, was performed using the PDQuest Basic 8.0.1 software (Bio-Rad, Hercules, CA, USA). The relative molecular mass was calibrated with internal protein markers (Precision Plus Protein Standards, Bio-Rad) after co-migration during the 2nd dimension. Quantification of detected protein spots was performed calculating the relative optical density × area (relative OD × area) in the gels. Normalization was set up according to the total spot density. Three different image analyses (one for each phenological stage) were performed. Since three biological repetitions per treatment (C1, C2, Ds and Np) were considered for the 2D approach, a total of 12 gel images per phenological stage were included in each analysis. Protein spots detected in at least 2 biological repetitions of a given treatment were considered for analysis and compared in all the treatments. Among the differentially expressed protein spots, 20 from the G stage, 15 from the flowering and 13 from the veraison (Figure 3) were subjected to in gel trypsin digestion followed by nanoLC–MS/MS analysis.

The mean relative OD × area ± SD (n = 3) values of each group were finally used to estimate relative expression level (relative OD × area %) of each protein spot among the groups. Differences among the means were evaluated by the Dunn’s Multiple Comparison Test after that the null hypothesis (equal means) was rejected in the Kruskal-Wallis test, assuming a significance of p ≤ 0.05. The relative expression ratio to the control (C1) in the other treatments was also estimated. Values ≥ |2| were discussed.

3.2.3. Protein Identification by Mass Spectrometry

NanoLC-MS/MS analyses were performed on a nanoACQUITY Ultra-Performance-LC system (UPLC) coupled to a Q-TOF mass spectrometer (maXis, Bruker Daltonics, Bremen, Germany) mass spectrometer equipped with a nano electrospray source. The UPLC system was equipped with a Symmetry C18 precolumn (20 × 0.18 mm, 5 μm particle size, Waters, Milford, MA, USA) and an ACQUITY UPLC® BEH130 C18 separation column (75 μm × 250 mm, 1.7 μm particle size, Waters). The solvent system consisted of 0.1% v/v formic acid in water (solvent A) and 0.1% v/v formic acid in acetonitrile (solvent B). Peptides were trapped during 3 min at 5 μL/min with 99% A and 1% B. Elution was performed at 60 °C at a flow rate of 450 nL/min, using a linear gradient of 6%–35% B over 9 min. The mass spectrometer was operating in positive mode, with the following settings: source temperature was set to 150 °C while drying gas flow was at 5 L/min. The nano-electrospray voltage was optimized to −4500 V. External mass calibration of the TOF was achieved before each set of analyses using Tuning Mix (Agilent Technologies, Palo Alto, CA, USA) in the mass range of 322–2722 m/z. Mass correction was achieved by recalibration of acquired spectra to the applied lock masses [methylstearate ([M + H]⁺ 299.2945 m/z) and hexakis (2,2,3,3,-tetrafluoropropoxy) phosphazene ([M + H]⁺ 922.0098 m/z)].

For tandem MS experiments, the system was operated with automatic switching between MS and MS/MS modes in the range of 100–2500 m/z. The two most abundant peptides (absolute intensity threshold of 2000), preferably doubly, triply and quadruply charged ions, were selected from each MS
spectrum for further isolation and CID (Collision Induced Dissociation) fragmentation using nitrogen as collision gas. Ions were excluded after acquisition of two MS/MS spectra and the exclusion was released after 6 seconds. The complete system was fully controlled by Hystar 3.2 (Bruker Daltonics, Bremen, Germany).

Raw data collected during nanoLC-MS/MS analyses were processed, converted into mgf files with DataAnalysis (Bruker Daltonics, Bremen, Germany).

The MS/MS data were analyzed using the MASCOT 2.2.0. algorithm (Matrix Science, London, UK) and OMSSA (Open Mass Spectrometry Search Algorithm, National Institut of Health, Bethesda, MD, USA [60]) for search against an in-house generated protein database composed of protein sequences of *Vitis* and Fungi (taxonomy 3603 and 4751) and known contaminant proteins such as human keratins and trypsin, extracted from the NCBInr database (version 3; September 2013) and combined with reverse sequences for all entries using an in-house database generation toolbox available at https://msda.unistra.fr (total 3.409.536 entries).

Searches were performed without any molecular weight, or isoelectric point restrictions, trypsin was selected as enzyme, carbamidomethylation of cysteine (+57 Da) and oxidation of methionine (+16 Da) were set as variable modifications and mass tolerances on precursor and fragment ions of 10 ppm and 0.02 Da were used, respectively. Mascot and OMSSA results were loaded into the Scaffold software (version 2.2.0, Proteome Software Inc., Portland, OR, USA) and filtered in order to evaluate the false discovery rate [61]. Protein identification was confirmed when at least two peptides with high quality MS/MS spectra (less than 5 points below Mascot’s threshold score of identity at 95% confidence level, or an OMSSA E-value below $-\log(e5)$) were identified. A more stringent filter was applied for single peptide identifications, the score of the unique peptide had to be higher than 10 points above Mascot’s threshold score of identity at 95% confidence level and an OMSSA E-value below $-\log(e10)$ was required. These thresholds led to protein identification with a false discovery rate of less than 1%.

3.2.4. Functional Classification of Identified Proteins

A functional classification of the identified proteins was performed by using GenomeNet Database Resources (http://www.genome.jp/kegg) or according to their role described in the literature.

4. Conclusions

Involvement of the fungal pathogens (*N. parvum* and *D. seriata*) in the necrosis development was confirmed by their reisolation from the edge of the lesions. Results of the pathogenicity tests are in agreement with the proteome changes observed which also report a weakness status of the vine at the flowering stage. Indeed, a general trend of a down-accumulation of proteins (e.g., superoxide dismutase, SSP) was observed in green stems infected with *N. parvum* or *D. seriata*, especially at flowering, except for the over-accumulation of some HSP as well as a PRP 10 in DsF and some proteins involved in the primary metabolism in NpF. Inversely, strongest responses to the infection were particularly activated in G stage through an over-accumulation of primary metabolism proteins, defense and stress-related proteins (e.g., oxygen evolving enhancer, *S*-adenosylmethionine synthetase, 2-cys peroxiredoxin, pathogenesis related protein 10) in correlation with a lower development of the
necrosis. According to the inoculated fungal strains, the disruption of the plant metabolism and the necrosis development are not closely related, indicating that different factors are involved in the virulence of the two fungal strains tested. Globally, the flowering stage seems to be the period of highest sensitivity to Botryosphaeria dieback agents possibly as consequence of the high metabolic activity oriented towards developing inflorescences. Little research has focused on the relationship between primary metabolism and defense responses, and therefore it would be interesting to unravel how primary metabolism occurring during the flowering stage influences defense responses.

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Author Contributions

A.S. conducted grapevine sampling, optimized protein extraction from woody tissues, performed the 2D gels and the image analysis and also wrote most of the manuscript. P.L. performed the inoculation of the plants with the fungi, measured the necrosis sizes, performed the detection of the fungi and participated to the sampling. M.M.R. participated in the writing of the manuscript. A.H. performed the identification of the proteins by mass spectrometry and the bioinformatics analyses. Because of expertise in proteomics on grapevine tissues, C.Ci. suggested the concept of the experimental design and participated in the redaction of the manuscript. C.S.R. helped with bioinformatic analyses and contributed to the interpretation of the results. A.V.D. participated in the correction of the manuscript. C.Cl. took part in the critical revision of the manuscript. FF supervised the study by coordinating the experiments, participated to the interpretation of the results and critically revised the manuscript.

Conflicts of Interest

The authors declare no conflict of interest.

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