Gene expression trade-offs between defence and growth in English elm induced by Ophiostoma novo-ulmi

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INTRODUCTION

Trees are exposed to many instances of environmental interaction and stress throughout their life cycle. These include pests and diseases that can cause dramatic mortality events in forests and loss of foundation species and iconic trees (e.g., Ellison et al., 2005). Diseases affecting tree vascular systems are usually very harmful due to the crucial role of these tissues in transporting water, nutrients, hormones, and other compounds. Dutch elm disease (DED) is a vascular wilt disease caused by the pathogenic fungi Ophiostoma ulmi (Buisman) Melin & Nannf and the more virulent wilt disease caused by the pathogenic fungi Ophiostoma novo-ulmi. Dutch elm disease (DED) is a vascular crucia role of these tissues in transporting water, nutrients, hormones, and other compounds. Dutch elm disease (DED) is a vascular disease caused by the fungus Ophiostoma novo-ulmi, destroyed most of North American and European elm populations in the 20th century. The highly susceptible English elm, also known as Atinian clone, suffered the highest mortality rates during the last pandemic event, probably due to its lack of genetic diversity. To study the DED pathosystem, we inoculated English elm ramets with O. novo-ulmi and evaluated xylem anatomy, molecular response, and disease symptoms. The high DED susceptibility of the clone was linked to xylem structure. The transcript levels changed significantly for 1,696 genes during O. novo-ulmi invasion. Genes covering different steps of the plant immune system were identified, many of which showed homology with Arabidopsis thaliana genes involved in systemic acquired resistance. Induction of several pathogenesis-related proteins and repression of fasciclin-like arabinogalactan proteins and other cell wall biosynthesis pathways evidence unbalanced costs between growth and defence mechanisms far from the inoculation point. This study sheds light on elm molecular defence mechanisms against DED.

KEYWORDS
Dutch elm disease, microarrays, plant immune system, systemic acquired resistance, Ulmus minor, vascular pathogen

Abstract
Wilt diseases caused by vascular pathogens include some of the most damaging stresses affecting trees. Dutch elm disease (DED), caused by the fungus Ophiostoma novo-ulmi, destroyed most of North American and European elm populations in the 20th century. The highly susceptible English elm, also known as Atinian clone, suffered the highest mortality rates during the last pandemic event, probably due to its lack of genetic diversity. To study the DED pathosystem, we inoculated English elm ramets with O. novo-ulmi and evaluated xylem anatomy, molecular response, and disease symptoms. The high DED susceptibility of the clone was linked to xylem structure. The transcript levels changed significantly for 1,696 genes during O. novo-ulmi invasion. Genes covering different steps of the plant immune system were identified, many of which showed homology with Arabidopsis thaliana genes involved in systemic acquired resistance. Induction of several pathogenesis-related proteins and repression of fasciclin-like arabinogalactan proteins and other cell wall biosynthesis pathways evidence unbalanced costs between growth and defence mechanisms far from the inoculation point. This study sheds light on elm molecular defence mechanisms against DED.

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DED may have favoured the spread of the disease during pandemic events (Gil et al., 2004).

Despite the major impact of DED pandemics on Atinian elms, some iconic ornamental trees surviving in cities and villages and other individuals in seminatural elm populations such as Rivas-Vaciamadrid (Madrid, Spain) belong to this clone (Gil et al., 2004; Martín, Solla, Burón, López-Almansa, & Gil, 2006). Atinian elms commonly occur as immature trees in most European countries affected by DED due to the ability of the clone to resprout from roots (Dunn, 2000). The prevalence of this susceptible clone and its role in the spread of DED pandemics make it particularly suitable for advancing knowledge of how *O. novo-ulmi* interacts with *U. minor* during vascular colonization.

Diseases affecting plant vascular systems, known as wilt diseases, cause severe damage and are difficult to study because of their complexity (Yadeta & Thomma, 2013). As Bernier et al. (2015) reported, the pathosystem formed by interaction between *Ulmus* spp. and *O. ulmi* or *O. novo-ulmi* has often been analysed in a very reductionist manner. The limited molecular knowledge of the DED pathosystem contrasts with current knowledge of plant–pathogen interactions in model plants. Research on these organisms has elucidated the molecular bases of these interactions. Plants have developed various molecular pathways integrated into a complex plant immune system. Plant cells sense invading pathogens via extracellular receptors known as pattern recognition receptors. This innate immune system identifies general or specific conserved microbe-specific molecules, referred to as microbe- or pathogen-associated molecular patterns (MAMPs or PAMPs), which induce a first cascade of defence called MAMP- or PAMP-triggered immunity (MTI or PTI) (Jones & Dangl, 2006).

A second line of plant defence is activated after direct or indirect recognition of a given effector by a set of plant resistance (R) gene products, resulting in effector-triggered immunity (ETI; Jones & Dangl, 2006). R proteins are commonly intracellular receptors containing leucine-rich repeats (LRRs), although other R genes without this domain have been described (Gururani et al., 2012). Once plant cell receptors sense a pathogen, both MTI/PTI and ETI are activated to coordinate the plant defence response, essentially as correlative phenomena. The major signalling pathways involved in posterior signal transduction in response to pathogens are mediated by kinases, reactive oxygen species (ROS), and proteins modulating signalling driven by concentrations of calcium ions (Ca\(^{2+}\)). These cascades modulate the expression of transcription factors (TFs) or genes involved in the synthesis of new regulators, mainly salicylic acid (SA) and jasmonic acid (JA; Moore, Loake, & Spoel, 2011; Pieterse, Leon-Reyes, Van der Ent, & Van Wees, 2009). SA and JA are thought to form the backbone of the plant defence system. Other hormones, such as ethylene, abscisic acid, gibberellins, auxins, and cytokinins, are able to modulate the SA- and JA-dependent signalling pathways (Pieterse, Van der Does, Zamioudis, Leon-Reyes, & Van Wees, 2012). Accumulation of endogenous SA is required to activate systemic acquired resistance (SAR), a long-lasting, broad-spectrum induced resistance characterized by coordinated activation of a specific set of pathogenesis-related (PR) proteins, most of which have antimicrobial activity (Van Loon, Rep, & Pieterse, 2006).

Earlier studies on elm responses to DED focused on individual components produced by the tree, such as phytoalexins (Jeng, Alfenas, Hubbes, & Dumas, 1983), lignin and suberin (Aoun, Rioux, Simard, & Bernier, 2009; Martín, Solla, Woodward, & Gil, 2007), and phenolic compounds (Aoun et al., 2009; Krause, Ichida, Schreiber, & Domír, 1996; Rioux & Ouellette, 1991), or on fungal molecules such as cerato-ulmin (Takei & Hiratsuka, 1984) and peptideorhamnomannans (Nordin & Strobel, 1981). More recently, in vitro and in vivo studies have started to unravel *Ulmus–Ophiostoma* interaction from a molecular perspective, reporting variation in expression of several defence-related genes encoding enzymes involved in phytoalexin biosynthesis, various PR proteins, and genes associated with the phenylpropanoid pathway (Aoun et al., 2009; Nasmith, Jeng, & Hubbes, 2008; Sherif, Shukla, Murch, Bernier, & Saxena, 2016). New omics technologies have significantly increased the genetic information available for *Ulmus* spp. Resources include recent sequencing of an expressed sequence tag (EST) library in response to elm leaf beetle (*Xanthogaleruca luteola*) infestation and spraying with methyl jasmonates (Büchel et al., 2012) and a first transcriptome in response to abiotic and biotic stresses (Perdiguero, Venturas, Cervera, Gil, & Collada, 2015b). In the latter case, *U. minor* genotypes with contrasting DED tolerance were inoculated with *O. ulmi* and *O. novo-ulmi* and subjected to other stresses.

In this study, we hypothesize that *O. novo-ulmi* is a hemibiotrophic pathogen, which induces JA- and/or SA-mediated signalling pathways in xylem tissues of infected plants. As a result, these induced pathways lead to the overexpression of specific defence responses in distal parts of the plant. As *O. novo-ulmi* is a vascular pathogen, we also hypothesize that the xylem anatomical features of the English elm and the structural defences induced by *O. novo-ulmi* in its vascular tissues are related to the high susceptibility of this clone. To test these hypotheses, 6-year-old Atinian elm ramets were inoculated with *O. novo-ulmi* to evaluate their defence responses. Xylem anatomical structures (where the pathogen spreads) and the long-distance (systemic) molecular responses activated during plant colonization were analysed. To elucidate the genes involved in the *U. minor* immune system and the molecular changes after inoculation, oligomicroarrays were constructed using the data from the transcriptome available in the Dryad database (Perdiguero, Venturas, Cervera, Gil, & Collada, 2015a) and hybridized with cDNA obtained from the ramets over a time course post inoculation. To the best of our knowledge, this is the first study to address the global differential expression of *U. minor* genes in *in vivo* colonization of *O. novo-ulmi*. Numerous genes and pathways showed significant changes, some previously reported in other plant species. The results allow us to advance knowledge of molecular responses that may be specific to vascular wilt diseases and, in particular, of the recognition of genes unique to the DED pathosystem.

2 | MATERIALS AND METHODS

2.1 | Plant material

Twelve ramets from elm genotype VA-AP38 (Arrabal del Portillo, Valladolid) propagated in 2008 from hardwood cuttings were...
planted in a research plot at Puerta de Hierro Forest Breeding Centre (Madrid, Spain; 40.456238 N, 3.751823 W) in spring 2009. Ramets were planted 1.0 × 0.75 m apart. The plot was watered during spring and summer to ensure plant establishment and growth and avoid drought stress.

Genotype VA-AP38 was identified as Atinian elm using molecular markers, following the process described by Martín et al. (2015). Leaf DNA was extracted following the protocol of Dumolin, Demesure, and Petit (1995). The chloroplast lineage was analysed with two markers (SFM and ccmp2), and 10 nuclear microsatellites (Ulm1-98 and Ulm1-165; Ulm2, Ulm3, and Ulm8; UR 123, UR 141, UR 153, UR 158, and UR 159) were used to determine whether the genotype belonged to the Atinian elm clone. The primer and polymerase chain reaction (PCR) conditions are shown in Table S1.

### 2.2 Inoculation, sampling times, and symptom analysis

At the beginning of May 2013, when the ramets were 6 years old, an aggressive local strain of O. novo-ulmi (Z-BU1) was inoculated into the selected genotype following the procedure described by Solla, Martín, Ouellette, and Gil (2005). Inoculation was performed about 20 days after full leaf expansion by introducing at midday (during maximum transpiration rates) 0.1 ml of a suspension of O. novo-ulmi spores (10⁶ spores ml⁻¹) into the sap stream through a wound made at the base of the trunk with a sharp blade. Each experimental unit, randomly distributed in the plot, consisted of two consecutive ramets. One ramet was inoculated with O. novo-ulmi spore suspension and the other with distilled water as a control. A healthy 3-year-old branch at approximately 2 m height was collected at 1, 3, 7, 14, and 21 days post inoculation (dpi) from inoculated and control plants at midday. The first branch segment (5 cm) was collected for anatomical structure analysis. Branches were tagged and immediately frozen in liquid nitrogen and stored at −80 °C. Disease development was evaluated in previously selected branches by estimating the percentage of wilting leaves. These observations were performed at sampling times (1, 3, 7, 14, and 21 dpi) and at 28, 60, and 120 dpi.

### 2.3 Characterization of anatomical structure

Anatomical parameters were measured in two control ramets. Two additional ramets inoculated with the pathogen were sampled to visualize xylem responses to infection by O. novo-ulmi. Four 3-year-old twig segments (3 cm length, 1.3–1.9 cm diameter) were taken from the cardinal points of the crown of each tree and immediately placed in formalin–acetic alcohol (formaldehyde:acetic acid:70% ethanol, 5:5:9 v/v/v) fixative. Samples were taken from control ramets at Day 0 and from inoculated plants at Day 120 post inoculation.

Twig samples were sectioned in radial, transverse, and tangential cuts (20 μm thick) using a Leica SM2400 Sledge Microtome. Cuts were dehydrated through an alcohol series, placed on glass slides, stained with safranine 0.1% (w/v), and fixed with Canada balsam. Sections were examined under an Olympus BX50 light microscope connected to a personal computer. Measurements were taken from the 2013 growth increment using ImageJ software (http://rsweb.nih.gov/ij/).

Bordered pit measurements were made on radial sections using pits located in earlywood vessels with radial diameters of 60–100 μm. Pit membrane diameter and pit aperture area were recorded. All earlywood vessels in the radial section of the 2013 annual ring were included. At least eight sections per sample, comprising 50–60 measurements per ramet, were evaluated.

Earlywood vessel measurements were made on transverse sections, randomly selecting four radial 500-μm-wide sectors 90° apart in the earlywood. Vessel diameters were measured radially in each sector, disregarding vessels less than 20 μm in diameter. The Hagen–Poiseuille equation (Giordano, Salleo, Salleo, & Wanderlingh, 1978) was used to estimate the relative theoretical hydraulic conductance (THC) of vessels by dividing the sum of the fourth power of internal vessel radii by the sector area. Vessel transactional area was obtained by dividing the area occupied by the vessels in a sector by the total area of the sector then multiplying by 100. At least four sections per sample, comprising 100–200 measurements per ramet, were used.

Tangential sections were used to measure ray tangential area (area occupied by the rays in a sector by the total area of the sector, multiplied by 100). At least eight sections per sample, comprising 50–60 measurements per ramet, were examined.

### 2.4 Microarray design and hybridization

Encoding a range of proteins, 14,898 isotigs were obtained from a transcriptome described by Perdiguero et al. (2015a) and included in the microarray design (Agilent 8 × 60 K, Agilent Technologies, CA, USA). For each isotig, one 60-base-long probe was designed and spotted at least 3 times on the slide. Probes designed for Populus, mouse, and human ESTs available in public databases were included as negative controls.

Total RNA for each sampling time was extracted from branches following the CTAB–LiCl precipitation method (Chang, Puryear, & Cairney, 1993). RNA was purified using the Qiagen RNAeasy kit (Qiagen, CA). RNA amplification and labelling were performed following Adie et al. (2007). Three biological replicates for control and inoculated plants for each sampling time (1, 3, 7, 14, and 21 dpi) were hybridized using two colours (inoculated vs. control). “The manual two-colour microarray-based gene expression analysis” protocol (Agilent Technologies, CA, USA) was followed for hybridizations. Images from Cy3 and Hyper5 channels were equilibrated and captured with a GenePix 4000B (Axon, CA, USA) and spots quantified using the GenePix software (Axon, CA, USA). The microarray design and hybridization data were deposited in ArrayExpress database under accession numbers A-MTAB-611 and E-MTAB-5654, respectively.

### 2.5 Data analysis

Background correction and normalization of expression data were performed using Linear Models for Microarray Data (Smyth, 2004; Smyth & Speed, 2003), part of “Bioconductor, an R language project”
Differentially expressed genes (DEGs) were evaluated by the non-parametric algorithm "Rank Products," available as the "RankProd" package at "Bioconductor, an R language project" (Breitling, Armengaud, Ammann, & Herzyk, 2004; Hong et al., 2006). This method detects genes that are consistently high ranked in a number of replicated experiments independently of their numerical intensities. Results are provided in the form of p, defined as the probability that a given gene is ranked in the observed position by chance. The expected false discovery rate was controlled to be less than 5 %.

Changes in the expression of a gene relative to control plants were estimated using the average signal intensity across the dataset from three biological replicates. Based on the statistical analysis, a gene was considered to be significantly up- or down-regulated for a genotype if it met all three of the following criteria: (a) FDR Rank Prod p < .05; (b) fold change ≥ ±1.6 for up-regulation or ≤ -1.6 for down-regulation at any sampling time, and (c) the trend (up- or down-regulation) was consistent in all datasets for each isolate.

Hierarchical clustering of DEGs was performed using the log2 of average expression value from three biological replicates and K-means method, selecting 12 clusters as threshold of the MeV 4.4 software (Saeed et al., 2006).

2.6 Single enrichment and gene set enrichment analysis

Full GO (Gene Ontology) term annotation and KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways information from the U. minor transcriptome available in Dryad database (Perdiguer et al., 2015a) were used in functional analysis of DEGs.

GO term enrichment was analysed for every trend according to clustering by comparison with the remaining selected significant genes identified during O. novo-ulmi invasion, using Fatigo (Al-Shahrour, Diaz-Uriarte, & Dopazo, 2004) implemented in Blast2GO software. This programme conducts a Fisher's exact test for 2 x 2 contingency tables to check for significant over-representation of GO annotations. GO term enrichment was significant at p < .05.

KEGG pathway enrichment analysis was performed using GSEA software (Subramanian et al., 2005). Every gene set represents a full KEGG pathway and was defined as the aggregation of all genes identified along the pathway. A data matrix containing the log2 of average expression value from three biological replicates was used to analyse pathways positively or negatively correlated with the main trend. Time course analysis using Pearson's correlation was applied with a 1,000 gene set permutation. Change in pathways was significant at p < .05.

2.7 Real-time quantitative PCR

Validation of the transcriptome profiling experiment was performed on selected genes identified from the microarray experiment by real-time reverse transcription polymerase chain reactions (RT-PCRs) using RNA from one of the samples used as a biological replicate in the microarrays. For RT-PCR, RNA was treated with DNase Turbo (Ambion; Applied Biosystems, Life Technologies, CA, USA). First-strand cDNA was synthesized from 1 μg total RNA from each sample using PowerScriptIII reverse transcriptase (Invitrogen) according to the supplier's manual. After verifying that the signal intensity remained unchanged across Ophiostoma invasion, 18S rRNA was used as a control.

The primers for the selected genes are shown in Table S2. PCRs were performed in an optical 96-well plate with a CFX 96 detection system (Bio-Rad), using EvaGreen to monitor dsDNA synthesis. Reactions containing 2× SsoFast EvaGreen Supermix reagent (Bio-Rad, CA, USA), 12.5 ng cDNA, and 500 nM of primers in a final volume of 10 μl were subjected to the following standard thermal profile: 95 °C for 3 min, 40 cycles of 95 °C for 10 s, and 60 °C for 10 s. Three technical replicates were performed for each PCR run. To compare the data from different PCR runs or cDNA samples, the mean of the threshold cycle (CT) values of the three technical replicates was normalized to the mean CT value of R18S. The expression ratios were then obtained using the ΔΔCT method corrected for the PCR efficiency for each gene (Pfaffl, 2001).

3 RESULTS

3.1 Identification of Atinian elm by molecular markers

Chloroplast DNA analysis showed that VA-AP38 belongs to the same Italian lineage as the Atinian clone (Table 1), and the alleles found for the 10 nuclear loci tested revealed that VA-AP38 had the same genotype as Atinian clone reference samples from England and Spain (Gil et al., 2004).

3.2 Symptom analysis

Plants were visually analysed to evaluate symptoms in the 4 months post inoculation. O. novo-ulmi inoculated plants showed the first DED symptoms at 14 dpi, when 10–15 % of their leaves were wilted. Wilting values were above 50 % at 21 dpi, the final sampling time for molecular analysis. At later dates (28, 60, and 120 dpi), all O. novo-ulmi infected plants showed wilting values above 80 %. No DED symptoms were observed in control plants during the experiment (Figure 1).

3.3 Anatomical structure

Atinian clone showed wide earlywood vessels and high values of vessel transsectional area and THC compared with data reported for twig samples of Siberian elm (Ulmus pumila) and U. minor clones tolerant to DED (Table 2). The pit membrane diameter of Atinian elm was similar to values found in U. pumila and tolerant U. minor, but pit aperture area was larger. The tangential section had large medullary rays and therefore a higher proportion of ray parenchyma cells than in U. pumila and tolerant U. minor. In the ramets inoculated with O. novo-ulmi,
abundant tylosis formation was observed in earlywood vessels at 120 dpi (Figure 2).

### 3.4 Identification, clustering, and functional analysis of DEGs

Time course analysis revealed that 1,696 of the isotigs included in the microarray (approximately 12%) showed significant differences in transcription levels during *O. novo-ulmi* infection; these were considered DEGs. DEGs were clustered in 12 main trends by transcription patterns (Figure 3 and Table S3). Most of the DEGs correspond to two profiles: six clusters (C1, C2, C6, C9, C11, and C12; 42.33% of DEGs) predominantly showed inductions in mean trends, whereas the other six (C3, C4, C5, C7, C8, and C10) mainly presented decreases in transcript levels during *O. novo-ulmi* colonization. C1 included the genes most induced during infection. The transcript levels of these genes increased progressively from the first dpi. A similar trend was observed in C12, although the expression level was lower than C1. In contrast, genes included in C2 showed highest induction during the first 24 hr post inoculation. C6 grouped genes with moderate increases in transcript levels at intermediate sampling times, mainly 3 and 7 dpi. The final clusters showing induction of genes, that is, C9 and C11, included genes with strong increases in transcript levels at 21 dpi, with higher expression values in C11. Clusters with repression in expression patterns, C3, C4, and C5, evidenced a progressive decrease in transcript levels, with lower expression values at 21 dpi. C5 was the cluster that included the most repressed genes. Clusters C7 and C8 grouped genes with decreases in transcript levels at 14 dpi, with C7 maintaining the repression at 21 dpi, whereas C8 recovered similar expression to control plants at 21 dpi. C10 included genes with moderate repression at 3 and 7 dpi.

Single enrichment analysis was performed by comparing functionalities assigned to the 1,696 DEGs and those identified within the 14,898 genes included in the array design. In the class "cellular component," the functional categories "cell wall," "extracellular region," and "plasma membrane" were over-represented in DEGs.

### TABLE 1 Genetic characterization of VA-AP38

<table>
<thead>
<tr>
<th></th>
<th>SFm</th>
<th>ccpm2</th>
<th>Ulm1-98</th>
<th>Ulm1-16S</th>
<th>Ulm2</th>
<th>Ulm3</th>
<th>Ulm8</th>
<th>UR 123</th>
<th>UR 141</th>
<th>UR 153</th>
<th>UR 158</th>
<th>UR 159</th>
</tr>
</thead>
<tbody>
<tr>
<td>VA-AP38</td>
<td>297</td>
<td>215</td>
<td>126</td>
<td>151</td>
<td>146</td>
<td>156</td>
<td>102</td>
<td>108</td>
<td>176</td>
<td>176</td>
<td>196</td>
<td>196</td>
</tr>
</tbody>
</table>

Note. Alleles for the two chloroplast DNA markers and 10 nuclear microsatellites used are shown.

### TABLE 2 Mean xylem parameters (and range values) measured in 3-year-old branches of Atinian elm and *Ulmus pumila* and tolerant *Ulmus minor* clones reported in a previous work

<table>
<thead>
<tr>
<th>Anatomical trait</th>
<th>Atinian elm</th>
<th><em>U. pumila</em></th>
<th>Tolerant <em>U. minor</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Earlywood vessel diameter (μm)</td>
<td>55.8 (20.4–125.8)</td>
<td>49.5 (27.0–119.6)</td>
<td>48.5 (27.4–89.4)</td>
</tr>
<tr>
<td>VTA (%)</td>
<td>32.9 (17.0–68.0)</td>
<td>25.2 (21.3–30.6)</td>
<td>16.9 (12.4–28.9)</td>
</tr>
<tr>
<td>THC (μm²)</td>
<td>251.9 (93.8–429.1)</td>
<td>129.2 (21.3–30.6)</td>
<td>79.5 (12.4–28.9)</td>
</tr>
<tr>
<td>Pit membrane diameter (μm)</td>
<td>7.0 (5.3–9.5)</td>
<td>7.2 (4.8–9.4)</td>
<td>7.2 (4.5–11.3)</td>
</tr>
<tr>
<td>Pit aperture area (μm²)</td>
<td>4.6 (2.9–6.3)</td>
<td>2.4 (0.8–4.8)</td>
<td>2.7 (0.8–5.8)</td>
</tr>
<tr>
<td>Ray tangential area (%)</td>
<td>10.1 (6.2–14.7)</td>
<td>7.5 (6.3–9.2)</td>
<td>8.5 (7.4–10.0)</td>
</tr>
</tbody>
</table>

*Values reported in Martín et al. (2009).*
Also, in the class "biological process," all the functional categories related with response ("stress," "abiotic," "biotic," "extracellular," and "endogenous stimulus") and other categories including "secondary metabolic process," "signal transduction," "cell growth," "cell death," or "transport" were significantly enriched within DEGs (Figure S1).

Several GO term categories were associated with trends previously identified in clustering analysis. Global functional analysis of clusters at Level 2 in the class "biological process" (Figure 3) showed that the GO terms "signalling," "response to stimulus," and "biological regulation" are therefore slightly over-represented in clusters showing clear induction during disease development. In contrast, in clusters showing repression throughout infection, GO terms "reproduction," "multicellular organismal process," "growth," and "developmental process" presented higher percentages. Enrichment analysis between clusters, comparing GO terms from biological processes or molecular functions, showed clear enrichment in some functionalities for every cluster (Figure 4). Significant enrichment (p < .05) in GO terms "cell death," "response to biotic stimulus," "response to stress," "response to endogenous stimulus," and "response to external stimulus" was found in all clusters with overexpression patterns. Specific functionalities were observed in certain clusters, for example, "oxygen binding," "sequence-specific DNA binding transcription factor activity," and "signal transduction" in C2, "protein binding" in C6, and "transport," "transporter activity," and "transferase activity" in C9. In repressed gene clusters, significant enrichment of GO terms "anatomical structure morphogenesis," "cell growth," and "cell differentiation" was observed in C3 and C4, whereas in C7, C8, and C10, enrichment occurred in GO term "response to abiotic stimulus" and in other terms at a lower percentage.

### 3.5 U. minor genes involved in response to O. novo-ulmi

Genes notably overexpressed or repressed compared to noninoculated plants were identified from the 1,696 DEGs during O. novo-ulmi invasion, mainly at 14–21 dpi. The 25 top up-regulated genes were all included in Cluster 1 or 11. These genes reached 50- to 350-fold higher average expression values in infected plants than in controls. Various PR proteins were encoded by 16 of the 25 induced genes. The remaining nine genes encoded five enzymes (valencene synthase, UDP-glycosyltransferase, 2-methylene-furan-3-one reductase, phenylalanine ammonia-lyase, and feruloyl ortho-hydroxylase) and one plant basic secretory family protein, and three genes were annotated as hypothetical or uncharacterized proteins. In contrast, the transcript levels of the 25 top down-regulated genes reached values of −30- to −170-fold in infected relative to control plants (Table 3). The most repressed genes included five encoding fasciclin-like arabinogalactan proteins (FLAs), one encoding the enzyme germacrene D synthase, one encoding an aquaporin, and one encoding a tubulin. Notably, no homology with known proteins was found for eight strongly repressed genes.

In addition to these PR proteins, other well-known genes were identified in the DEGs, including receptors potentially involved in the perception of O. novo-ulmi, kinases and other genes involved in signal
transduction, and genes encoding TFs potentially involved in activation of defence genes.

3.5.1 Genes involved in perception
In total, 76 U. minor genes, grouped in clusters by expression pattern, were identified as receptors or annotated with GO term “receptor activity” (GO:0004872). Of these, 45 showed increased transcript levels during infection and 31 showed decreased levels (Table S4). At least 39 genes contained an LRR domain. Three U. minor genes up-regulated during O. novo-ulmi invasion were identified as R protein receptors, nucleotide-binding site-leucine-rich receptor-interleukin-1 receptors (NB-LRR-TIR), and nucleotide-binding site-leucine-rich receptor-coiled coil (NB-LRR-CC; Table S4).

3.5.2 Genes involved in signal transduction
During pathogen infection, at least eight genes showing significant changes in transcript levels were identified as kinases potentially involved in signal transduction, including various mitogen-activated protein kinases. A group of 17 genes were identified as calcium-related or calcium-binding proteins, some of which showed remarkable induction at different sampling times during disease development (Table S5).

3.5.3 Transcription factors
Of the 1,696 DEGs, 108 were identified as TFs, and 89 of them were annotated with GO term “transcription factor activity” (GO:0003700). No significant enrichment was identified for this GO term during global functional analysis. However, specific analysis
within this group showed a significant enrichment in specific families such as WRKY TF, ethylene-responsive TF, and nac-domain TF. In addition, more than 80% of the members from these three families were overexpressed during the response to the pathogen. In other families such as dehydration-responsive element-binding protein or heat stress TF, most of the members included in the array were also identified as DEGs (Figure S2). Seventy-one TFs presented increased transcript levels at one or more sampling times during disease development. An nac-domain TF reached the highest induction for TFs at 14 dpi (Table S6).

3.5.4 Genes involved in defence
A group of induced DEGs was related to regulation mediated by SA. Two up-regulated genes, isotig02962 and isotig10240, annotated as senescence-associated carboxylesterase 101-like (SAG101), are homologous to Arabidopsis thaliana enhanced disease susceptibility 1 (EDS1) and phytoalexin deficient 4 (PAD4), respectively, genes acting as molecular players in SA pathways. Three correspond to genes identified as SA-binding proteins. Isotig11511 was significantly induced in the days immediately following inoculation (7 and 4 dpi) and was annotated as npr1 nim1-interacting.
### TABLE 3  Top 25 up-regulated and down-regulated *Ulmus minor* genes during *Ophiostoma novo-ulmi* invasion

<table>
<thead>
<tr>
<th>Locus TAIR</th>
<th>Description Blast2GO</th>
<th>Cluster number</th>
<th>Fold change Day 1</th>
<th>Day 3</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
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Note. Locus TAIR indicates protein from *Arabidopsis thaliana* showing highest homology with the *Ulmus minor* sequence. For determination of significance, see Section 2.
protein, which is potentially involved in late regulation of SAR (Table S7).

Forty-eight genes from *U. minor* with significant induction were identified as members of different PR protein families. A remarkable increase in transcript levels was observed in members of certain families, mainly in *PR2*-type (glucan endo-beta-glucosidases), *PR4*-type (chitinases), *PR5*-type (thuaumin), *PR9*-type (peroxidases), and *PR14*-type (non-specific lipid transfer proteins). Notable overexpression was observed in some genes included in the unclassified PR proteins group, for example, an early nodulin, a late embryogenesis abundant hydroxyproline-rich glycoprotein, and an α-amylase (Table S8).

Another group of induced genes, catalogued as transporters, showed the highest induction in genes involved in polyols, ammonium, sugars, and zinc transport. Numerous genes encoding ATP binding cassette (ABC) transporters also showed remarkable increase in transcript levels. Genes related to biotic stress response (major allergen or some *gdsl* esterase lipase genes) showed significantly increased transcript levels. Moderate variation in transcript levels, usually down-regulated during *O. novo-ulmi* invasion, was observed in seven genes that presented homology with WAT1 (*Walls are Thin1*) gene from *A. thaliana* (Table S7).

### 3.6 Pathways correlated with disease development

Genes encoding enzymes involved in various pathways included in KEGG were analysed globally to identify specific metabolic pathways correlated with disease development. A pathway is considered significantly enriched when a group of genes encoding enzymes in the pathway are positively or negatively correlated with the main trend observed in expression patterns. A group of metabolic processes with significant enrichment (*p* < .05) were identified during the analysis (Figure 5 and Table S9).

Twelve of the metabolic pathways identified were increasingly up-regulated over time. The pathways included biosynthesis of terpenoids, anthocyanins, flavonoids, isosquoline alkaloids, and indole alkaloids, as well as metabolism of butanone, propanoate, and glyoxylate and dicarboxylate.

Pathways related to metabolism of amino acids such as valine, alanine, and tyrosine were also significantly enriched. Four pathways were down-regulated over time, mainly showing repression in enzymes related to pentose and glucuronate interconversions, starch and sucrose metabolism, sphingolipid metabolism, and fatty acid biosynthesis.

### 3.7 Validation of microarray data by RT-PCR

To validate the microarray results, quantitative RT-PCR was performed to determine the expression levels of eight *Ulmus* genes selected to cover several functional groups and expression patterns from the list of DEGs in response to *O. novo-ulmi*. RT-PCR confirmed the microarray expression patterns (Figure 6). Seven of the eight genes analysed showed a Pearson's correlation value higher than 0.9, including values close to 1 in some genes. Only one gene showed a lower Pearson's correlation value (0.6), which could be attributed to the different dynamic range of the two techniques.

### 4 DISCUSSION

The main objective of this work was to characterize the expression profiles of plant defence genes in *U. minor* over the time course of vessel invasion by *O. novo-ulmi*, the causal agent of DED. We examined the Atinian elm, also known as English elm, an iconic tree widely distributed throughout Europe since Roman times. This clone is known to be highly susceptible to *O. novo-ulmi* attack (Martín, Solla, Coimbra, & Gil, 2008), as confirmed in this experiment. Inoculated plants showed severe wilting and almost total canopy desiccation by the end of the observation period (120 dpi). The evolution of the disease has been correlated to physiological parameters such as water potentials, gas exchange, and hydraulic conductance, all of which decreased drastically for this clone in infected compared to control plants (Li et al., 2016). This study focuses on the molecular response of this clone to DED and the influence of its xylem anatomical traits.

The anatomical structure of Atinian elm is perfectly adapted to maximize hydraulic conductance in early spring. The earlywood conducting system is formed almost exclusively of wide earlywood vessels, endowing it with high THC values. In 3-year-old branches, THC values are particularly high in Atinian elm compared to values in elms tolerant to *O. novo-ulmi* (Table 2) and other susceptible clones (Martín, Solla, Esteban, de Palacios, & Gil, 2009; Venturas et al., 2014). Anatomical structure probably contributes to the high susceptibility of the clone to *O. novo-ulmi*, favouring the spread of fungal bud cells, toxins, and cell wall-degrading enzymes, ultimately leading to vessel occlusion and/or embolism. The large pit apertures in the clone could also facilitate the spread of fungal products in the xylem sap and penetration of the pit membrane by fungal hyphae (Martín et al., 2009). The high proportion of ray parenchyma cells is important, because this tissue has a decisive role in (a) exudation of tyloses, gels, and gums into the vessels through pit connections (Figure 2; Chattaway, 1949) and (b) storage of nonstructural carbohydrates (NSCs; Magel, Jay-Allemand, & Ziegler, 1994; Martín et al., 2008). In relation to tylose formation, a strong reaction of the Atinian elm to *O. novo-ulmi* inoculation was detected in cross sections at 120 dpi, with a large proportion of earlywood vessels occluded by tyloses and gums (Figure 2). This reaction corresponds to Wall 1 of the CODIT model (Shigo & Marx, 1977) and may restrict upward and downward movement of the pathogen. However, this reaction can be so efficient that vertical sap transport to the canopy becomes blocked, contributing to tree death (Ouellette, Rioux, Simard, & Cherif, 2004). Therefore, the higher proportion of ray cells in the Atinian elm than in tolerant elms could give this clone greater ability to block vessels that are infected or near infected tissue. The anatomical structure of Atinian elm also suggests a high storage capacity of NSCs in medullary rays, possibly related to the great capability of this clone to occlude vessels with tyloses and gels or gums. These components include accumulation of suberin and polyphenolic compounds, which are formed from stored NSCs and are toxic to microbes (Shain, 1979). NSCs can, however, be a source of easily assimilable metabolites for fungal growth.
once the fungus is able to penetrate living cells (Martín et al., 2009; Ouellette, 2008). Despite uncertainty as to the exact role of xylem parenchyma cells in DED pathogenesis, these tissues may be responsible for major defence responses triggered in an attempt to compartmentalize the DED pathogen.

Relationships between vascular wilt pathogens and their host plants are complicated to study because they take place inside plants, usually in specific niches (e.g., conducting elements). From a molecular perspective, response to wilt pathogens triggers activation of the plant immune system and induction or repression of specific genes in wilt pathogen-infected tissues (Yadeta & Thomma, 2013, and references therein). Therefore, the molecular response of *U. minor* to *O. novo-ulmi* should induce common and specific patterns during disease development. To date, arguably the most in-depth molecular study analysing the DED pathosystem was performed by Aoun, Jacobi, Boyle, and Bernier (2010) using *in vitro* interactions between *Ulmus americana* cells and *O. novo-ulmi* to identify a set of DEGs. An interesting group of transcripts were identified, some of which have recently been searched for in mature trees to analyse their putative relationship with DED tolerance (Sherif et al., 2016). Using high-throughput sequencing, the genetic information available for *Ulmus* spp. has been considerably increased with the creation of an EST library in response to feeding or methyl jasmonates (Büchel et al., 2012) and a *U. minor* transcriptome in response to abiotic stress (drought) and biotic stresses (including inoculation with *Ophiostoma* spp. and other pathogens; Perdiguero et al., 2015a, 2015b). However, to be able to identify genes of interest in these libraries, it is necessary to conduct differential expression analysis focusing on a specific problem.

Herein, we report 1,696 DEGs during *O. novo-ulmi* invasion, of which 89% showed homology with proteins from a set of plant model species contained in RefSeq database according to the *U. minor* transcriptome used as reference (Perdiguero et al., 2015a). Using clustering analysis, DEGs were grouped into 12 clusters with shared expression patterns. This approach could be a basis for constructing useful coexpression networks highlighting putative protein–protein interactions during *O. novo-ulmi* invasion. Interestingly, coherence was identified in results from functional enrichment analysis. Clusters showing up-regulation in expression patterns were associated with GO terms “response to biotic stimulus” and “response to stress,” and also “cell death,” the final consequence of immune system activation (Jones & Dangl, 2006). Enrichment shown by Cluster 2, including the terms “oxygen binding,” “sequence-specific DNA binding transcription factor,” and “signal transduction,” coincident with high induction 24 h post inoculation, indicates rapid activation of molecular response to the

![FIGURE 5](image-url)
presence of the pathogen even at considerable distance from the pathogen entry point: Branches collected for the study had a healthy appearance and were 2 m from the inoculation point.

The immune system of infected plants produces plant hormones that actively regulate response to infection by controlling signalling pathways. SA is a signal molecule required for SAR. It is a generally observed pattern that SA regulates resistance to biotrophs and hemibiotrophs, whereas JA regulates resistance to necrotrophs (Pieterse et al., 2009). O. novo-ulmi is considered a hemibiotroph, and therefore, an SA predominant response should be expected, although both types of plant resistance responses could be involved. SA and JA accumulations have been reported in U. americana plants inoculated with O. novo-ulmi (Sherif et al., 2016). In the molecular response of Atinian elm, overexpression of genes related to SAR to pathogens from 7 to 21 dpi is noteworthy. These SAR-related genes also showed a progressive increase in expression over time. This response to pathogens (generally bacteria or fungi) has been widely described in other plants and is dependent on SA (Vlot, Dempsey, & Klessig, 2009). After localized pathogen infection, several signals are triggered in distal parts of the plant to enhance defence against invasion in the entire plant. In our study, we found several up-regulated genes with high homology to SAR-related genes described in Arabidopsis. These genes are mainly grouped in Clusters 1 and 11 (Table S3). Isotig02962 and isotig10240, homologous to EDS1 and PAD4, respectively, showed similar changes in expression at the same sampling times. Isotig03130, homologous to SAG101, also showed a significant increase in transcript levels from

FIGURE 6 Validation of microarrays by reverse transcription polymerase chain reaction (RT-PCR). Expression profiles obtained with microarray (bars) and RT-PCR techniques (continuous line) for eight selected differentially expressed genes. Pearson’s correlation (r) between techniques is indicated for each gene in each panel.
7 to 20 dpi. This situation corresponds precisely to the common SAR response described in other plants infected by fungal pathogens. In A. thaliana infected by Fusarium graminearum, proteins EDS1, PAD4, and SAG101 act in a complex that promotes SA accumulation and, therefore, SA-mediated defence, limiting infection by the pathogen (Makandar et al., 2015). In this complex, EDS1 is essential to recognize pathogens through R proteins and, consequently, triggers resistance at pathogen infection foci (Feys, Moisan, Newman, & Parker, 2001). EDS1 and PAD4 are partners in basal resistance to various pathogens. These proteins are regulators of ETI, which is mediated by resistance (R) genes (Feys et al., 2005). In our analysis, several isotigs were annotated as genes involved in this signalling pathway (Table S4). Genes coding for LRRs or TIRs were principally induced in Ophiostoma-inoculated trees from 7 dpi onwards, and this activation coincides with increased expression of SAR-marked genes, such as phenylalanine ammonia-lyase and PR genes (Table 3). These genes have been described as overexpressed during DED development (Aoun et al., 2009; Nasmith et al., 2008) and were the most up-regulated genes in our study. Seven of the 25 top up-regulated genes (Table 3) were identified by Aoun et al. (2010) in U. americana callus.

In the classic SAR response, activation of PR genes contributes to enhance whole-plant resistance (Pieterse et al., 2014). Several genes annotated as PR were found with high and progressively increased expression from 3 or 7 dpi to 21 dpi in inoculated trees. We highlight the up-regulation of genes with homology to Arabidopsis PR proteins, such as PR2 with β-1,3-glucanase activity, several chitinases and PR4, thaumatin-like proteins belonging to the PR5 family, and non-specific lipid transfer proteins classified as PR14 (Table S8). Although PR1 is a good marker of SAR, its function is still unknown; however, the other PR families are all proteins with antifungal activity (Van Loon & Van Strien, 1999). We therefore confirmed that inoculation with O. novo-ulmi triggered activation of several genes encoding antifungal activities in distal tissues in VA-AP38.

This activation occurred with increased expression of various WRKY TFs directly related to SAR regulation. Homologs to WRKY18, WRKY40, and WRKY70 appeared up-regulated in Ophiostoma-inoculated trees; these three TFs are positive regulators of SA-mediated defence and negative regulators of JA-mediated responses (Hu, Dong, & Yu, 2012; Pandey, Roccaro, Schön, Logemann, & Somssich, 2010). We also found a TF homologous to WRKY33 that regulates camalexin biosynthesis by inducing genes of this pathway, including various CYP71 genes (cytochrome p450 with monoxygenase activity; Mao et al., 2011) that were also up-regulated in this study (Clusters 1 and 11).

Enrichment analysis using pathways permitted identification of interesting processes during O. novo-ulmi invasion. The significant up-regulation observed in biosynthesis of terpenoids matched the accumulation of lignin, and the up-regulation of anthocyanin and flavonoid biosynthesis was consistent with accumulation of phenolic compounds (i.e., in vessel lumina for restricting axial movement of the pathogen), both reported in U. minor trees after O. novo-ulmi infection (Martin et al., 2007, 2008; Ouellette et al., 2004).

Several up-regulated genes associated with oxidative stress were identified at 14 and 21 dpi (Clusters 9 and 12). Genes encoding peroxidases, glutaredoxins, glutathione S-transferases, and nac-TFs are expressed at these sampling times, indicating an increase in oxidative stress induced by the pathogen. As there was no evidence of pathogen presence at the sampling time, a hypothetical situation could be explained by an increase of signalling mediated by ROS. These ROS are produced mainly by the NADPH oxidases in cells of direct contact with the pathogen. This process leads to great increases in the superoxide anion (O2−), which is reduced to H2O2 by superoxide dismutases. H2O2 can diffuse into cells and move long distances, acting as a signal with SA and nitric oxide in many defence mechanisms in plants (Mittler, 2002).

Clusters with down-regulated genes (not previously analysed in other elm studies) showed functionalities related to growth or anatomical structure and morphogenesis. These changes reflect differences in the energy use strategy of infected plants. In the absence of virulent pathogens, plants invest energy in growth, but when their physiological functions are challenged by virulent pathogens, they use energy to form defence structures and substances (Huot, Yao, Montgomery, & He, 2014). A group of strongly repressed genes (Cluster 4) encoded FLAs that appear to contribute specifically to the biomechanical properties of stems through their impact on the synthesis and architecture of the secondary cell wall. Analyses with inactivated or mutant FLA transgenic plants (MacMillan, Mansfield, Stachurski, Evans, & Southerton, 2010; Wang et al., 2015) showed down-regulation of some xylem-specific genes involved in cell wall formation that resulted in altered stem biomechanics of transgenic plants. In view of the key role of FLA in stem composition and biomechanics, we hypothesize that strong down-regulation during disease development favoured invasion of vessels by the pathogen.

Among down-regulated pathways, pentose and glucuronate interconversions and starch and sucrose metabolism are represented by numerous groups of genes involved in starch formation and regulation of pectin esterification. Both processes are important in cell wall development and defence responses. This discovery, along with other down-regulated genes related to secondary metabolism, lipid metabolism, and cellular cycle processes, suggests that the SAR response is directly associated with a decrease in development, differentiation, and vegetative growth. These results agree with earlier research demonstrating depletion of U. minor starch reserves during O. novo-ulmi infection, which was more intense in susceptible than in resistant genotypes (Martin, Solla, Coimbra, & Gil, 2005).

The down-regulation of genes related to auxin and gibberellin signalling pathways (Clusters 5, 7, and 8) in Ophiostoma-inoculated trees from 14 to 21 dpi is noteworthy. It is well known that these hormones control growth in plants by promoting induction of a large number of growth-related genes (Davies, 2010). However, some pathogens also produce auxins or manipulate auxin biosynthesis in plants. To avoid pathogen growth, plants produce SA to repress the auxin and gibberellin signalling pathways and stabilize auxin response repressors as part of the defence mechanisms (Wang, Pajerowska-Mukhtar, Culler, & Dong, 2007). The repressed auxin-related genes found in our study could be a direct antagonistic effect of SAR in trees inoculated with O. novo-ulmi.

Identification and classification of defence-related genes are considered a task of primary importance for the forest research community (Kovalchuk et al., 2013). This work is a first global overview
Figure 7 Schematic representation of putative molecular responses in Atinian elms inoculated with *Ophiostoma novo-ulmi*. Inoculation at the base of the *Ulmus minor* trunk induces a local response where pathogen-associated molecular patterns (PAMPs) and resistance (R) proteins are expressed, generating two resistance responses: effector-triggered immunity (ETI) and PAMP-triggered immunity (PTI). This process activates the salicylic acid (SA) pathway in the affected area and a similar rapid response in distal parts of the tree through signalling by SA and reactive oxygen species such as nitric oxide (NO) and hydrogen peroxide (H$_2$O$_2$). This response in distal healthy tissues is known as systemic acquired resistance (SAR). The systemic response activates pathways involved in the increase of SA levels through induction of the first enzyme in the phenylpropanoid pathway (phenylalanine ammonia lyase; PAL) and the combined action of three proteins: senescence-associated carboxylesterase 101-like (SAG101), enhanced disease susceptibility 1 (EDS1), and phytoalexin deficient 4 (PAD4). This complex is regulated by two transcription factors, WRKY18 and WRKY40. Increased SA levels activate the expression of pathogenesis-related genes (PR) that codify for different proteins with antifungal activity, whereas SA-related pathways down-regulate genes involved in jasmonic acid (JA)-mediated signalling and growth and development. Some of these genes are related to the actions of other phytohormones such as auxins and gibberellins, for example, indoleacetic acid-induced protein 9 (IAA9) and gibberellin (GA)-regulated protein, whereas others are directly involved in plant development, for example, the fasciulin-like arabinogalactan proteins (FLAs). In the systemic response, other pathways are induced, producing positive feedback on SA-related pathways and suppressing JA-related pathways, as with camalexin synthesis regulated by transcription factor WRKY33 and a cytochrome p450 of the CYP71 family of the molecular response of *U. minor* saplings to xylem invasion by *O. novo-ulmi*. The DEGs reported in this analysis are the first expression level study in elm saplings infected with *O. novo-ulmi*. Among these DEGs, a significant number of genes were identified across the steps that occur during plant immune system activation. Numerous genes potentially involved in perception, signal transduction, and transcription modulation, as well as defence genes, were identified and analysed. The results highlight several receptors, kinases, and TFs in various families that are certain to provide interesting material for advancing knowledge of the DED pathosystem. Based on the results of this study, Figure 7 shows a putative model of a systemic response induced in *U. minor* inoculated with *O. novo-ulmi* and highlights pathways with greater differential expression between inoculated and control plants.

A considerable number of the DEGs reported are homologous to genes that have been described as key regulators during plant–pathogen interactions in other species, including interesting genes identified in other wilt diseases (Yadeta & Thomma, 2013). For instance, WAT1 encodes a tonoplast localized auxin transporter that has been associated with resistance to vascular pathogens in *A. thaliana*, including the fungus *Verticillium dahliae* (Denancé et al., 2013). A large group of genes with remarkable up- or down-regulation during disease development showed no homology to proteins from other plants. These genes pose new challenges for researchers interested in identifying specific genes involved in wilt diseases of trees or specifically involved in *Ulmus* sp. and *O. novo-ulmi* interaction. Most of the up-regulated and down-regulated genes found in this study indicate that the response in healthy branches located distally from the inoculation point resembles the general SAR responses described for plant defence. Gruner, Griebel, Návarová, Attaran, and Zeier (2013) presented an in-depth study showing all the up- and down-regulated genes in SAR-associated defence in *Arabidopsis* against *Pseudomonas syringae*. Their work strongly supports our data, because many of the genes described in *Arabidopsis* were found here in *U. minor*. Our results support our initial hypotheses, because the pathogen promoted a systemic response in distal parts of the plant, and this response corresponded with the typical SAR mediated by SA. In order to determine the traits behind the high susceptibility of this clone to *O. novo-ulmi*, analyses of tissues in the vicinity of the inoculation point are needed to clearly discern the differences between early local response and systemic responses. Possibly, factors such as the timing and intensity of the local defence responses are crucial to determine the degree of susceptibility to the pathogen. The described anatomical structure of this clone probably contributed to its high susceptibility, favouring pathogen spread in vascular tissues. Our results significantly advance knowledge of vascular diseases of trees and the plant defence mechanism against pathogens. This knowledge is crucial in advancing towards molecular assisted techniques to select and breed trees resistant to vascular pathogens.

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CONFLICT OF INTEREST
The authors have no conflict of interest to declare.

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