

Comprehensive analysis of *Verticillium nonalfalfae* in silico secretome uncovers putative effector proteins expressed during hop invasion

1 Kristina Marton¹, Marko Flajšman¹, Sebastjan Radišek², Katarina Košmelj¹, Jernej Jakše¹,
2 Branka Javornik¹, Sabina Berne^{1*}

3 ¹Department of Agronomy, Biotechnical Faculty, University of Ljubljana, Ljubljana, Slovenia

4 ²Slovenian Institute of Hop Research and Brewing, Žalec, Slovenia

5 * Correspondence:

6 Sabina Berne

7 sabina.berne@bf.uni-lj.si

8 Abstract

9 Background: The vascular plant pathogen *Verticillium nonalfalfae* causes Verticillium wilt in several
10 important crops. *VnaSSP4.2* was recently discovered as a *V. nonalfalfae* virulence effector protein in
11 the xylem sap of infected hop. Here, we expanded our search for candidate secreted effector proteins
12 (CSEPs) in the *V. nonalfalfae* predicted secretome using a bioinformatic pipeline built on *V.*
13 *nonalfalfae* genome data, RNA-Seq and proteomic studies of the interaction with hop.

14 Results: The secretome, rich in carbohydrate active enzymes, proteases, redox proteins and proteins
15 involved in secondary metabolism, cellular processing and signaling, includes 263 CSEPs. Several
16 homologs of known fungal effectors (LysM, NLPs, Hce2, Cerato-platanins, Cyanovirin-N lectins,
17 hydrophobins and CFEM domain containing proteins) and avirulence determinants in the PHI
18 database (Avr-Pita1 and MgSM1) were found. The majority of CSEPs were non-annotated and were

narrowed down to 44 top priority candidates based on their likelihood of being effectors. These were examined by spatio-temporal gene expression profiling of infected hop. Among the highest *in planta* expressed CSEPs, five deletion mutants were tested in pathogenicity assays. A deletion mutant of *VnaUn.279*, a lethal pathotype specific gene with sequence similarity to SAM-dependent methyltransferase (*LaeA*), had lower infectivity and showed highly reduced virulence, but no changes in morphology, fungal growth or conidiation were observed.

Conclusions: Several putative secreted effector proteins that probably contribute to *V. nonalfalfae* colonization of hop were identified in this study. Among them, *LaeA* gene homolog was found to act as a potential novel virulence effector of *V. nonalfalfae*. The combined results will serve for future characterization of *V. nonalfalfae* effectors, which will advance our understanding of Verticillium wilt disease.

Keywords: *Verticillium*, hop, plant-pathogen interaction, effector, secretome, ATMT, virulence, *LaeA*

1 Background

Soil-born vascular plant pathogens, members of the *Verticillium* genus [1], cause Verticillium wilt in several economically important crops, including tomato, potato, cotton, hop, sunflower and woody perennials [2,3]. Studies of *Verticillium* – host interactions and disease processes, particular those caused by *V. dahliae*, have significantly contributed to the understanding of *Verticillium* spp. pathogenicity, although more research is needed for successful implementation of Verticillium wilt disease control [4–6].

Plant-colonizing fungi secrete a number of effectors, consisting among others of hydrolytic enzymes, toxic proteins and small molecules, to alter the host cell structure and function, thereby facilitating

infection and/or triggering defense responses [7]. The assortment of effector molecules is complex and highly dynamic, reflecting the fungal pathogenic lifestyle [8] and leading to pathogen perpetuation and development of disease. Research into plant-pathogen interactions has significantly advanced, with an increasing number of sequenced microbial genomes, which have enabled computational prediction of effectors and subsequent functional and structural characterization of selected candidates. However, the prediction of fungal effectors, mainly small secreted proteins, which typically lack conserved sequence motifs and structural folds, is challenging and largely based on broad criteria, such as the presence of a secretion signal, no similarities with other protein domains, relatively small size, high cysteine content and species-specificity [8–10]. Using these features to mine predicted secretomes for candidate effectors has been valuable, but has not produced a one-size-fits-all solution [11]. Various approaches that combine several bioinformatics tools and also consider features such as diversifying selection, genome location and expression *in planta* [12–15], have had mixed outcomes. The EffectorP application has recently been presented as the first machine learning approach to predicting fungal effectors with over 80% sensitivity and specificity [16].

The genome sequences of five *Verticillium* species (*V. dahliae*, *V. alfalfae*, *V. tricorpus*, *V. longisporum* and *V. nonalfalfae*) and their strains have been published [17–22], providing a wealth of genomic information for various studies. Klosterman et al. [17] queried *V. dahliae* (strain VdLs.17) and *V. alfalfae* (strain VaMs.102) genomes for potential effectors and other secreted proteins based on subcellular localization and the presence of signal peptide. A similar number of secreted proteins was found in both genomes (780 and 759 for *V. dahliae* and *V. alfalfae*, respectively), and comparable to other fungi. These secretomes were further examined for effector candidates, obtaining 127 for *V. dahliae* and 119 for *V. alfalfae* proteins, based on the assumption that fungal effectors are small cysteine-rich proteins (SSCPs) with fewer than 400 amino acids and more than

four cysteine residues. Siedl et al. [19] later re-examined the secretomes of two *V. dahliae* strains (VdLs.17 and JR2) and *V. alfalfae* (VaMs.102), and predicted a higher number of secreted proteins and smaller number of SSCPs, due to improved gene annotation and restricted criteria for SSCPs. Interestingly, in their comparison of highly pathogenic *Verticillium* species with saprophytic and weak pathogen *V. tricornutum*, a similar content of the secretome (cca 8.5%) in their respective proteomes was obtained. Orthologs of known effectors of *F. oxysporum*, *C. fulvum* or oomycete *Phytophthora infestans* were not found in *V. dahliae* and *V. alfalfae* genomes [17], except for *C. fulvum* LysM effector Ecp6 [23] (7 and 6 genes in *V. dahliae* and *V. alfalfae*, respectively) and *C. fulvum* virulence factor Ecp2 [24]. Several LysM effectors, widespread fungal proteins recognized by the LysM domain, have also been characterized as suppressors of PTI (PAMP-triggered immunity) through their chitin binding ability [25–28]. Reexamination of *Verticillium* LysM effectors has corroborated only three LysM effectors as core proteins in the genomes of *V. dahliae* strains with a function other than fungal pathogenicity, and one strain specific (VdLs.17) virulence associated LysM protein [18,27]. An increase in NLP (necrosis and ethylene-inducing protein (NEP-1)-like proteins) gene homologs has also been found among secreted proteins in *Verticillium* genomes (8 and 7 in *V. dahliae* and *V. alfalfae*, respectively) [17,19]. Zhou et al. [29] and Santhanam et al. [30] showed that only two of them displayed cytotoxic activity in tomato, cotton, *Arabidopsis* and *Nicotiana benthamiana*, while reduced virulence has been demonstrated for the deletion mutant of *VdNLP1* and *VdNLP2* in tomato and *Arabidopsis*.

Although numerous secreted proteins with unknown function or sequence similarity have been reported for *Verticillium* spp, only a handful have been characterized. Virulence effector Ave1 is a 134 aa long secreted protein with 4 conserved cysteines and an Expansin-like EG45 domain, discovered by comparative genomics in *V. dahliae* race 1 strains [31]. Ave1 recognition by tomato receptor-like protein Ve1 triggered immune signaling pathways leading to resistance to *V. dahliae*

race 1 strains [32,33]. The other reported *V. dahliae* effector protein, PevD1, induced a hypersensitive response in tobacco [34,35] and triggered innate immunity in cotton plants, as demonstrated by upregulation of defense-related genes, metabolic substance deposition and cell wall modifications [36]. Zhang et al. [37] recently characterized a novel effector protein, VdSCP7, as a host nucleus targeting protein, which induced a plant immune response and altered the plant's susceptibility to fungal and oomycete pathogens.

In addition to *V. dahliae* effectors, a small secreted protein, VnaSSP4.2, with an important role in fungal virulence, has been discovered in xylem sap during *V. nonalfalfae* infection of hops [38]. *V. nonalfalfae* is another pathogenic species of the genus *Verticillium*, although with a narrower host range than *V. dahlia* [1]. However, it causes Verticillium wilt and plant death in several important crops [4]. The occurrence of different virulent strains has been well documented in hop (*Humulus lupulus* L.), in which two pathotypes of *V. nonalfalfae* with different aggressiveness have been isolated, causing mild (fluctuating) and lethal (progressive) disease forms [39–43]. The disease is demonstrated by plant wilting, foliar chlorosis and necrosis, vascular browning and rapid plant withering and dieback in the lethal disease form [43]. Sporadic outbreaks of the *V. nonalfalfae* lethal pathotype in European hop gardens are of major concern, since there is no effective disease control except host resistance and strict phytosanitary measures. Despite the high economic losses caused by the lethal Verticillium wilt, the development of highly aggressive *V. nonalfalfae* pathotypes, as well as the genetics of hop resistance, remains enigmatic.

In the present study, a comprehensive biological database was generated using data from recently sequenced *V. nonalfalfae* genomes [22], transcriptomic and proteomic research of fungal growth on xylem stimulating medium [44] and RNA-seq studies of *V. nonalfalfae* – hop interactions [45]. A customized bioinformatics platform was used to set up a pipeline for prediction and characterization of the *V. nonalfalfae* secretome and select the best candidate secreted effector proteins (CSEPs) for

functional studies. From a total of 263 CSEPs in the final dataset, the gene expression of the 44 highest ranking CSEPs was assessed by spatio-temporal RT-qPCR profiling of infected hop. Furthermore, deletion mutants of five selected CSEPs were analyzed in pathogenicity assays, with one of them exhibiting reduced virulence on hop plants. Our findings should assist further characterization of *V. nonalfalfae* effectors in an attempt to understand the molecular mechanisms of Verticillium wilt disease.

2 Results

2.1 The *V. nonalfalfae* in silico secretome is rich in carbohydrate-active enzymes, proteases and candidate secreted effector proteins (CSEPs)

The *V. nonalfalfae* genome comprises 9,269 predicted protein-encoding genes. Among these putative proteins, 944 are classically secreted proteins with signal peptide and no more than one transmembrane domain, representing 10.2% of the *V. nonalfalfae* predicted proteome (Figure 1 and Table S1). The accuracy of prediction was evaluated by comparing this dataset to a set of 91 unique sequences obtained by proteomic analysis (2D-DIGE) of *V. nonalfalfae* proteins secreted in xylem simulating medium [44], resulting in a 81.3% match (Table S1). Using TMHMM and Phobius [46] for transmembrane (TM) domain prediction, 801 proteins without a TM domain and 161 proteins harboring one TM domain were determined (Table S1). Based on subcellular localization predictions, 709 extracellular proteins ('extr' > 17) were acquired with WoLF PSORT [47], 450 proteins residing in the apoplast were determined with ApoplastP [48], while Localizer [49] identified 52 proteins harboring a chloroplast targeting signal and 12 proteins with a signal sequence for localization in mitochondria (Table S1).

After similarity searches to known proteins in various databases, hypothetical functions were assigned to 727 (77%) putatively secreted *V. nonalfalfae* proteins. A superfamily annotation scheme

[50] was used to classify the *V. nonalfalfae* secretome into seven functional groups (Figure 2). In the 'Metabolism' group, pectin lyase-like proteins and proteins with a cellulose-binding domain were over-represented within the 'Polysaccharide metabolism and transport' category, while (trans)glycosidases and six-hairpin glycosidases were predominant in the 'Carbohydrate metabolism and transport' category. Other abundant proteins in the 'Metabolism' group were reductases and heme-dependent peroxidases in the 'Redox' category, Concavalin A-like lectins/glucanases in the 'Secondary metabolism' category and 'Transferases'. The major constituents of the 'Intracellular processes' group were 'Proteases' (especially acid proteases, Zn-dependent exopeptidases, subtilisin-like proteases and metalloproteases), cupredoxins in the 'Ion metabolism and transport' category and phospholipases in the 'Phospholipid metabolism and transport' category. Proteins involved in 'Cell adhesion' were over-represented within 'Extracellular processes', while most proteins in the 'Information' group classified into the 'DNA replication/repair' category. In the 'General' group, FAD-binding/transporter-associated domain-like proteins and proteins with a FAD/NAD(P)-binding domain were major constituents in the 'Small molecule binding' category. Analysis of the EuKaryotic Orthologous Groups (KOG) (Table S2) revealed a high number of proteins in the 'Cellular processing and signaling' category, associated with posttranslational modification, protein turnover, chaperones and signal transduction mechanisms, as well as cell wall/membrane/envelope biogenesis and intracellular trafficking, secretion, and vesicular transport, while the protein composition of the 'Metabolism' category mirrored that of the Superfamily.

Since 'Carbohydrate metabolic process' and 'Peptidase activity' were overrepresented terms after Blast2GO analysis of the *V. nonalfalfae* secretome (Figure S1), they were investigated more thoroughly. Almost one third of the putative *V. nonalfalfae* secretome was CAZymes, of which 255 were expressed *in planta* and distributed as follows: glycoside hydrolases (129 GH), carbohydrate esterases (47 CE), redox enzymes that act in conjunction with CAZymes (49 AA; proteins with

auxiliary activities), proteins with carbohydrate-binding modules (32 CBM), polysaccharide lyases (25 PL), and glycosyltransferases (4 GT). This repertoire of CAZymes was compared to other plant pathogenic *Verticillium* species (Figure 3A) and it was demonstrated that *V. nonalfalfae* had statistically more putative secreted CEs than *V. alfalfae*, in particular those involved in deacetylation of xylans and xylo-oligosaccharides. Moreover, the *V. nonalfalfae* secretome consisted of more putative secreted GHs than *V. dahliae* with major differences found in the GH3 group consisting primarily of stereochemistry-retaining β -glucosidases [51], the GH5 group of enzymes acting on β -linked oligo- and polysaccharides, and glycoconjugates [52], and the GH43 group of enzymes for the debranching and degradation of hemicellulose and pectin polymers [53]. In addition, the *V. nonalfalfae* secretome was enriched in putative secreted proteins with CBMs, when compared to *V. alfalfae*, *V. longisporum* and *V. dahlia*. This was largely due to cellulose-binding module CBM1 attached to various enzymes from families CE1, CE5, CE15, GH5, GH6, GH7, GH10, GH11, GH12, GH45, GH74, PL1, PL3 and AA9, and to some extent due to chitin-binding module CBM50 found in LysM effector proteins and subgroup C chitinases [54].

Similarity searching of *V. nonalfalfae* putative secreted proteins against peptidases in the MEROPS database revealed 12 *in planta* expressed aspartic peptidases, 2 cysteine peptidases, 27 metallopeptidases, 44 serine peptidases and 1 threonine peptidase. The highest representation of *V. nonalfalfae* putative secreted peptidases was in the M14A (carboxypeptidase A1), S08A (subtilisin Carlsberg) and A01A (pepsin A) subfamilies (Table S3). Comparison of putative secreted peptidases between plant pathogenic *Verticillium* species (Figure 3B) revealed a similar distribution of peptidases among *V. nonalfalfae*, *V. alfalfae* and *V. longisporum*, while *V. dahliae* had a statistically different distribution of metallopeptidases, cysteine and serine peptidases. Other enzymatic activities of putatively secreted *V. nonalfalfae* proteins according to the KEGG analysis can be found in Table S4.

184 Querying *Verticillium in silico* secretomes for small secreted proteins (SSPs) of less than 300 aa and
 185 small secreted cysteine rich (SSCPs) proteins with more than 5% cysteine content and at least 4 Cys
 186 residues [55] showed 5-10% lower abundance of SSPs and 2-3% fewer SSCP in the *V. nonalfalfae*
 187 secretome than in the secretomes of other plant pathogenic *Verticillium* species (Figure 3C). Since
 188 small secreted proteins are the least characterized portion of fungal secretomes and many have been
 189 shown to act as effectors, our secretome analysis further focused on filtering secreted proteins for
 190 expression *in planta* to identify CSEPs relevant to *V. nonalfalfae* infection of hop. Genome-wide
 191 transcriptome analysis of the *V. nonalfalfae* interaction with hop [45] revealed that 766 (81%)
 192 transcripts in the *V. nonalfalfae in silico* secretome were expressed in infected hop samples. They
 193 showed distinct expression patterns related to different stages of infection (6, 12, 18 and 30 dpi), hop
 194 cultivar (susceptible 'Celeia' or resistant 'Wye Target') and plant tissue (roots or shoots) (Figure S2).
 195 From this dataset, all CAZymes except CBMs were omitted from further analysis, resulting in 529
 196 putatively secreted *in planta* expressed proteins (Figure 1), of which 308 had sequence similarity to
 197 PFAM domains (Table S5). These included, among others, effector-specific PFAM domains, such as
 198 LysM effectors [27], Necrosis inducing proteins (NPP1) [56], Hce2 (Homologs of *Cladosporium*
 199 *fulvum* Ecp2 effector) effector proteins [57], Cerato-platanins [58], Cyanovirin-N lectins [59],
 200 hydrophobins [60] and CFEM (Common in Fungal Extracellular Membranes) domain containing
 201 proteins [61]. Our final dataset of CSEPs comprised a total of 263 proteins without functional PFAM
 202 domains, and proteins bearing known effector-specific PFAM domains, representing 2.8% of the
 203 putative proteome. Among them, we determined also 3 CSEPs with a nuclear localization signal
 204 (NLS), implying their activity in the plant nucleus, 3 CSEPs specific to the lethal strain of *V.*
 205 *nonalfalfae* and 69 probable effector proteins (Table S6) as predicted by EffectorP [16]. Similarity
 206 searching of CSEPs to experimentally verified pathogenicity, virulence and effector genes from
 207 fungal, oomycete and bacterial pathogens in the Pathogen-host interaction (PHI) database [62]

revealed proteins matching AVR effectors (5 hits) and known effector proteins displaying reduced virulence (11 hits) or unaffected pathogenicity (4 hits) (Table S6).

2.2 V. nonalfalfae CSEPs display distinct gene expression profiles during infection of hop

Establishing successful colonization of a host plant requires effective and timely delivery of the fungal pathogen's effectors. Using quantitative real-time PCR, the expression of the 44 top-priority CSEPs selected according to their likelihood of being effectors (see Methods for selection criteria), was investigated in root and shoot samples of Verticillium wilt susceptible ('Celeia') and resistant ('Wye Target') hop at 6, 12 and 18 days after inoculation with *V. nonalfalfae*. In a preliminary experiment, the average expression of the selected CSEPs in pooled root samples at different time points was examined (Table S7). The three highest expressed CSEPs that were also selected by Effector P prediction, and two lethal pathotype specific CSEPs, were then profiled using biological replicates (Figure 4). We included the *VnaSSP4.2* gene, encoding a small secreted protein, in the gene expression analysis as a positive control for virulence-associated *V. nonalfalfae* effector [38].

The expression levels of genes *Vna5.694*, *Vna7.443* and *Vna8.691* were greater in the roots than in the shoots of both hop varieties. Gene expression of *Vna5.694* (Figure 4A), encoding a small (81 aa) secreted cysteine-rich protein of unknown function and displaying the highest similarity to *V. longisporum* CRK15920 protein, decreased with time in the roots of susceptible hop, while its expression in the resistant hop increased. A similar trend of expression was also observed in the shoots of both hop varieties. Gene expression of *Vna7.443* (Figure 4B), which produces a secreted protein (276 aa without cysteines) with the highest similarity to *V. longisporum* CRJ82870 protein, was comparable to *Vna5.694*; it decreased in the roots of susceptible hop and peaked at 12 dpi in the roots of resistant hop. A peak of expression at 12 dpi was also observed in the shoots of susceptible hop, while its expression increased with time in the shoots of resistant hop. Expression of the *Vna8.691* gene (Figure 4C), coding for a small secreted protein (95 aa without cysteines) of unknown

function with the highest similarity to *V. longisporum* CRK10461 protein, was highest in the roots of susceptible hop at 6 dpi and then decreased with time of infection. The same trend was also observed in the shoots of susceptible hop. Expression of *Vna8.691* in the roots of resistant plants was constant, and around 200-fold higher than that in ½ CD medium, whereas no expression was detected in the shoots. Interestingly, two lethal pathotype specific genes *VnaUn.279* (Figure 4D), encoding a small secreted protein (92 aa without cysteines) with the highest similarity to *V. dahliae* VdLs.17 EGY23483 protein, and *Vna4.761* (Figure 4E), encoding a 186 aa protein with 8 cysteines and the highest similarity to *V. longisporum* CRK16219 protein, had similar gene expression patterns as the virulence-associated *V. nonalfalfae* effector *VnaSSP4.2* (Figure 4F). They were expressed only in the roots and shoots of susceptible hop (with expression levels increasing with time of infection) and barely detected in the resistant plants.

2.3 Identification of novel virulence effector of *V. nonalfalfae*

Since all five selected CSEPs were specifically expressed during plant colonization, a reverse genetics approach was used to test their contribution to the virulence of *V. nonalfalfae* in hop. Knockout mutants of *VnaUn.279* displayed only minor wilting symptoms in the susceptible hop (Figure 5), while vegetative growth, fungal morphology and sporulation were not affected. Analysis of the relative area under the disease progress curve (rAUDPC) [63] indicated that four independent *VnaUn.279* knockout mutants displayed statistically significantly lower values of rAUDPC than the wild type fungus (Figure 6A). To understand the progress of disease in time, statistical modelling was undertaken. For illustrative purposes only, disease severity index (DSI) values [42] for $\Delta VnaUn.279$ and wild type fungus were modelled by logistic growth model (Figure 6B). The variability of disease progression in individual hop plants is probably due to the specific nature of *Verticillium* colonization, in which only a few attached hyphae randomly penetrate the root intercellularly [64]. As demonstrated by the inflection point of the $\Delta VnaUn.279$ logistic curve,

development of disease symptoms was delayed for 10 days compared to the wild type fungus. Based on the asymptote values, the wilting symptoms were considerably less severe in the mutant than in the wild type fungus. Additionally, fungal biomass assessment with qPCR revealed that 32% of plants were infected with *V. nonalfalfae* $\Delta VnaUn.279$ mutants compared to at least 80% for the wild type fungus. These results indicate that deletion of *VnaUn.279* not only severely reduced *V. nonalfalfae* virulence but also significantly affected the fungal infectivity via a yet unknown mechanism.

The other tested *V. nonalfalfae* knockout mutants (*Vna5.694*, *Vna7.443* and *Vna8.691* in Figure S3) showed unaffected pathogenicity and no statistical differences in rAUDPC values relative to the wild type. Deletion of *Vna4.761* was not achieved, due to its functional redundancy, since two additional copies with over 95% sequence identity have been found after Blastn search against *V. nonalfalfae* reference genome at two different genomic locations.

3 Discussion

Fungal pathogens have evolved diverse strategies to interact with host plants and secrete various effector molecules to overcome plant defense mechanisms. A recently published genome of xylem-invading Sordariomycete fungus *Verticillium nonalfalfae* [22], a transcriptome study of infected hop [45] and obtained proteomic data of fungal growth on xylem simulating medium [44] have provided an opportunity to screen for proteins that may contribute to fungal virulence in hop. In the current study, a customized bioinformatics pipeline was designed to predict the classical *V. nonalfalfae* secretome and then to refine the secretome based on experimental data to identify candidate secreted effector proteins (CSEPs).

The relative secretome size of *V. nonalfalfae* (10.2%) conforms to the nutritional lifestyle of plant pathogens with larger secretome sizes (from 2.9% to 12.1% of the proteomes, with an average of

7.4%) and fits in the phylogenetic context with other Pezizomycotina (from 3.7% to 12.1% of the proteomes, with an average of 7.3%) [55]. The majority of proteins (69.5%) in the *V. nonalfalfae* secretome were less than 500 aa residues. In contrast to some plant pathogenic Pezizomycotina, which had remarkable 10-15% enrichment of proteins of up to 100 aa residues [55], only 1.8% of such proteins were found in the *V. nonalfalfae* secretome.

The composition of the *V. nonalfalfae* predicted secretome, rich in carbohydrate active enzymes (33%), proteases (11%), lipases/cutinases (4.6%) and oxidoreductases (4%), reflected its nutritional lifestyle as a hemibiotroph and plant vascular pathogen. Hemibiotrophic fungi undergo two phases during the infection process; an initial biotrophic phase, with characteristic expression of small secreted proteins without functional annotation (SSPs), is followed by a necrotrophic stage, which is generally associated with the expression of plant cell wall-degrading enzymes (CWDEs) [8]. Similar to *V. dahliae* and *V. alfalfae* genomes [65], the *V. nonalfalfae* genome encodes more CWDEs per number of secreted proteins than other plant pathogenic fungi [8]. Pectinases, xylanases, cellulases, glucanases, proteases, cutinases and lipases are major classes of CWDEs [66] and play important roles during plant colonization. They may facilitate penetration of the plant roots to reach the xylem vessels, degrade pectin gels and tyloses, formed in response to infection, to spread inside vessels, breakdown insoluble wall polymers to acquire nutrients and contribute to the release of survival structures from dead plant material [65]. In addition to contributing to virulence [67], some CWDEs are recognized as pathogen-associated molecular patterns (PAMPs), which provoke PAMP-triggered immunity [68,69]. On the other hand, *V. dahliae* carbohydrate-binding module family 1 domain-containing proteins may suppress glycoside hydrolase 12 protein-triggered immunity in plants [70]. As in the case of *V. dahliae* [65], the *V. nonalfalfae* predicted proteome contains numerous glycoside hydrolases and polysaccharide lyases, most of which are secreted [44]; however, only five glycosyl transferases were determined in the predicted secretome. In contrast to *V. dahliae*, there was almost

double the number of carbohydrate esterases in the *V. nonalfalfae* proteome and over half were predicted to be secreted. Other abundant proteins in the *V. nonalfalfae* predicted secretome were acid proteases, subtilisin-like proteases and zinc-dependent metalloproteases. These enzymes probably participate in amino acid acquisition, manipulation of host defenses by degradation of pathogenesis-related proteins, including plant chitinases, and act as virulence factors or as elicitors of defense responses [71,72]. A significant number of lipases, phospholipases and cutinases were determined in the *V. nonalfalfae* predicted secretome and identified in a previous proteomic study [44]. In addition to supplying energy for pathogen growth, lipid hydrolysis is crucial for the production of certain signaling molecules, such as oxylipins, which manipulate the host lipid metabolism and alter plant defense responses [73]. The role of cutinases in pathogenicity is controversial and has been associated with the dissolution of the plant cuticle during penetration, suppression of callose formation, spore attachment and surface signaling [74,75]. Another group of abundant enzymes in *V. nonalfalfae* predicted and experimentally determined secretomes were oxidoreductases, in particular FAD-dependent oxidoreductases and GMC oxidoreductases. Oxidoreductases are probably secreted for protection against host-produced reactive oxygen species, such as the generation of H₂O₂, which was detected after infection with *V. dahliae* in cotton roots [76] and in tomato plants [77]. On the other hand, fungal pathogens can actively contribute to the ROS level in host plants [78]. In *V. dahliae*, NADPH oxidase complex (Nox), composed of the catalytic subunit VdNoxB and tetraspanin VdPls1, is responsible for the production of ROS and the formation of penetration peg within the hyphopodium [64]. Moreover, VdNoxB regulates the cytoskeletal organization of the VdSep5-septin ring that separates the hyphopodium from invasive hyphae and forms a specialized fungus-host penetration interface, where small secreted proteins preferentially accumulate [79].

Since small secreted proteins (SSPs) are the least characterized fungal secreted proteins and some have been reported as effectors, we particularly focused on this group of proteins. Various criteria for

the determination of SSPs have been reported [12,80,81] but, for comparison purposes, we adopted a definition [55] that considers SSPs proteins with a mature length of ≤ 300 aa residues and proteins with a relative cysteine content of $\geq 5\%$, as well as ≥ 4 cysteine residues, to be small secreted cysteine-rich proteins (SSCP). According to these criteria, the *V. nonalfalfae* predicted secretome contains 310 SSPs (32.8% of the predicted secretome) and 46 (4.9% of the secretome) of those belong to SSCPs. These numbers are lower than the average contents of SSPs (49%) and SSCPs (6.7%) determined in fungi of class 2 secretome size (500-1100 secreted proteins) and average contents of SSPs (47%) and SSCPs (7.5%) in the Pezizomycotina group [55]. In a recent study, SSPs with a mature length of ≤ 300 aa residues were identified in 136 fungal species and compared in terms of taxa and lifestyles [82]. On average, hemibiotrophs and necrotrophs had higher proportions of secreted enzymes, while biotrophs, symbionts and certain hemibiotrophs had the most abundant SSPs. Furthermore, higher numbers of species-specific SSPs (over 100) were associated with biotrophs and symbionts than necrotrophs and saprotrophs (around 30), suggesting that these effectors coevolved with their hosts, while the range was widest for hemibiotrophs. However, no species-specific SSPs have been discovered in *V. nonalfalfae*, while 13 and 19 have been reported in *V. albo-atrum* and *V. dahliae*, respectively [82].

Further analysis of the refined *V. nonalfalfae* secretome, comprised of 263 CSEPs, focused in particular on homologs of known effectors from other plant pathogens. Searching for LysM effectors using CAZy module (CBM50) and PFAM domain PF01476 revealed that the *V. nonalfalfae* secretome contains four *in planta* expressed proteins with 2-6 LysM domains, of which Vna2.979 is an ortholog of VDAG_00902 (Table S6), a core LysM effector of *V. dahliae* [27]. The necrosis- and ethylene-inducing-like proteins (NLPs) are a group of widespread conserved effectors that can trigger immune responses and cell death [56]. Similar to other *Verticillium* spp. [19,65], NLP genes are expanded in the *V. nonalfalfae* genome, with seven genes orthologous to *V. dahliae* NLP1-9 and

having no ortholog to VdNLP6. All five NLPs with homologs in the PHI database (Table S6) were expressed in hop; the most abundant was Vna7.239 (VdNLP9), in particular in the roots of susceptible hop (Figure S2), suggesting some role in the plant colonization process. Four fungal hydrophobins, characterized by high levels of hydrophobicity and the presence of eight conserved cysteine residues [60], were found in the refined *V. nonalfalfae* secretome and had homologs in the PHI database (Table S6). Although all were expressed in hop, only expression of Vna7.87 was abundant and root-specific. Interestingly, a role in the development of microsclerotia [83] was demonstrated for type II hydrophobin VDH1 from *V. dahliae*, but it was not required for pathogenicity. Further mining of CSEPs for known effectors revealed that *in planta* expressed Vna2.8 and Vna3.54 are homologs of AVR-Pita1, a neutral zinc metalloprotease from *Magnaporthe oryzae* [84], and Vna1.1274 was similar to MgSM1, a putative small protein of the Cerato-platanin family [85]. Three *V. nonalfalfae* CSEPs, Vna10.263 and Vna5.719 with high and ubiquitous expression in hop, and Vna9.246 specifically expressed only in susceptible hop, had similarity to *Candida albicans* RBT4, secreted pathogenesis-related proteins [86], while one CSEP was an ortholog of urea amidolyase (DUR1,2), which enables utilization of urea as a sole nitrogen source [87]. Highly expressed secreted protein Vna4.130 had similarity to EMP1, extracellular matrix protein 1 from *Magnaporthe grisea*, which was required for appressorium formation and pathogenicity [88]. Vna7.617, which was putatively secreted and abundantly expressed in susceptible hop, had an ortholog in *Fusarium oxysporum* membrane mucin Msb2, which regulates invasive growth and plant infection upstream of Fmk1 MAPK [89].

The remaining *V. nonalfalfae* CSEPs (92%) were hypothetical, predicted and conserved hypothetical proteins with no functional annotation and their temporal gene expression patterns in susceptible and resistant hop were explored to provide some clues to their function (Table S7 and Figure S2). In addition to the already reported effector VnaSSP4.2 [38], several other CSEPs with distinct

expression patterns and high levels of expression were found. Among five CSEPs selected for gene functional analysis using a reverse genetics approach, four were predicted as effectors by EffectorP [16] and one had an ortholog in the PHI database, displaying sequence similarity to fungal effector LaeA, a regulator of secondary metabolism [90] and morphogenetic fungal virulence factor [91]. After comparing the pathogenicity of wild type fungus to CSEPs knockout mutants (Figure 6 and Figure S3), we discovered that the later CSEP, encoded by lethal pathotype specific gene *VnaUn.279*, is a novel virulence factor of *V. nonalfalfae*. $\Delta VnaUn.279$ mutants had diminished infectivity and exhibited severely reduced virulence in hop. Reduced virulence was also reported for a number of *LaeA* deletion mutants from human pathogen *A. fumigatus* [91], plant pathogenic fungi, including *A. flavus*, *C. heterostrophus* and several *Fusarium* species [92], as well as entomopathogenic fungus *Beauveria bassiana* [93]. Altogether, these findings justify further investigation of the biological role of *VnaUn.279* in *V. nonalfalfae* pathogenicity.

Despite the other selected CSEPs mutants not displaying any virulence associated phenotype, based on their expression profiles, they probably participate in other physiological processes during *V. nonalfalfae* infection of hop. Additionally, certain CSEPs may be recognized (and subsequently termed Avr effectors) by plant resistance proteins (R proteins), which are intracellular nucleotide-binding leucine rich repeat (NLR) receptors, via direct (receptor-mediated binding) or indirect (accessory protein-mediated) interactions, resulting in effector triggered immunity (ETI) [94,95]. To support this hypothesis, further testing of CSEPs mutants in the resistant hop cultivar is required and could result in identification of corresponding hop resistance proteins. These may then be exploited in Verticillium wilt control by introducing new genetic resistance traits into hop breeding, as already successfully implemented in certain other crops [96].

4 Conclusions

After comprehensively investigating the predicted *V. nonalfalfae* secretome using a diverse bioinformatics approaches and integrating multiple lines of evidence (genomics, transcriptomics and proteomics), several candidate secreted effector proteins were identified among protein-encoding genes. These are of high interest to scientists working on Verticillium wilt and, more generally, on pathogen effectors. Since the majority were non-annotated protein sequences, two strategies were adopted to gather clues about their function. With spatio-temporal gene expression profiling, we identified those candidate effectors that have important roles during *V. nonalfalfae* colonization of hop, while pathogenicity assays with effector knockout mutants revealed the candidates that contribute to fungal pathogenicity in hop. In conclusion, a new virulence effector of *V. nonalfalfae*, encoded by lethal-pathotype specific gene *VnaUn.279*, was identified and will be subject to future functional and structural studies.

5 Material and Methods

5.1 Microbial strains and cultivation

Sordariomycete fungus *Verticillium nonalfalfae* [1] was obtained from the Slovenian Institute of Hop Research and Brewing fungal collection. Two isolates from infected hop were used, differing in aggressiveness: lethal pathotype (isolate T2) and mild pathotype (isolate Rec) [97]. Fungal mycelium was cultured at room temperature on a half concentration of Czapek Dox broth ($\frac{1}{2}$ CD), supplemented with 1 g/L malt extract, 1 g/L peptone (all from Duchefa, The Netherlands) and 1 g/L yeast extract (Sigma Life Science, USA). For solid media, 15 g/L agar (Duchefa, The Netherlands) was added to $\frac{1}{2}$ CD with supplements. Alternatively, potato dextrose agar (PDA; Biolife Italiana Srl, Italy) or Xylem simulating medium (XSM; [98]) was used.

Escherichia coli DH5 α , used for amplification of vector constructs, was cultivated in LB medium with 50 mg/L kanamycin (Duchefa, The Netherlands) at 37°C.

Agrobacterium tumefaciens (LBA4404) transformation was performed in YM medium [99] containing 100 mg/L streptomycin and 50 mg/L kanamycin (both from Duchefa, The Netherlands) at 30°C. The co-cultivation of transformed *A. tumefaciens* and *V. nonalfalfae* was carried out on IMAS plates [99] at room temperature.

5.2 Functional annotation of *V. nonalfalfae* gene models and RNA-Seq analysis

Using a customized Genialis Platform (Genialis, Slovenia; <https://www.genialis.com/genialis-platform/>), the gene models of the *V. nonalfalfae* reference genome [22] were translated into putative proteins with the ExPASy Translate tool [100] and ORF Finder [101]. The general characteristics of putative proteins (molecular weight, number of amino acids (aa), percentage of cysteines and isoelectric point) were predicted by the ProtParam tool [100]. Functional annotation of the predicted proteins was performed with HMMER searches [102] against CAZy [103,104], Pfam [105], and Superfamily [106], as well as with BLAST searches [107] against NCBI, KOG [108], MEROPS [109] and PHI databases [62], followed by Blast2GO [110] and KEGG [111,112] analyses. The overrepresentation of GO terms in the *V. nonalfalfae* *in-silico* secretome compared to proteome was assessed using a hypergeometric distribution test (HYPGEOM.DIST function in Excel) with a *p*-value < 0.05 and FDR < 0.05.

RNA-sequencing of *V. nonalfalfae* mild and lethal pathotypes was performed by IGA Technology Service (Udine, Italy) using Illumina Genome Analyzer II. For this purpose, total RNA, enriched for the polyA mRNA fraction, was isolated in three biological replicates from fungal mycelia of mild and lethal strains grown in xylem-simulating media according to [98]. Illumina raw sequence reads were deposited at NCBI (Bioproject PRJNA283258). RNA-Seq analysis was performed using CLC Genomics Workbench tools (Qiagen, USA). Differentially expressed genes between lethal and mild

fungal pathotype were identified as those with fold change $FC \geq 1.5$ or $FC \leq -1.5$ ($p \leq 0.05$; $FDR \leq 0.05$).

From our previous RNA-Seq data of compatible and incompatible interactions between hop and *V. nonalfalfae* [45], fungal transcripts expressed at a least one time point (6, 12, 18 and 30 dpi) and one hop cultivar (susceptible 'Celeia' or resistant 'Wye Target') were filtered out and data were presented as a matrix of $\log_2$ CPM (counts per million – number of reads mapped to a gene model per million reads mapped to the library) expression values. These genes were considered as expressed *in planta*.

5.3 Secretome prediction and comparison

The *V. nonalfalfae in silico* secretome was determined using a customized Genialis Platform (Genialis, Slovenia) according to the method described in [113], which reportedly gives 83.4% accuracy for fungal secreted proteins. It combines SignalP4.1 [114], WolfPsort [47] and Phobius [46] for N-terminal signal peptide prediction, TMHMM (<http://www.cbs.dtu.dk/services/TMHMM/>) for eliminating membrane proteins (allowing one transmembrane (TM) domain in the first 60 aa) and PS-Scan [115] for removing proteins with ER targeting sequence (Prosit: PS00014). In addition, we used LOCALIZER [49] to predict effector protein localization to chloroplasts and mitochondria, while proteins with a nuclear localization signal were determined with NucPred (likelihood score >0.80) [116] and PredictNLS [117]. Localization of effector proteins to the apoplast was predicted by ApoplastP [48] based on enrichment in small amino acids and cysteines, as well as depletion in glutamic acid.

To compare the composition of the *V. nonalfalfae* secretome to other closely related plant pathogenic *Verticillium* species, protein coding sequences of *Verticillium dahliae* JR2, *Verticillium longisporum* GCA_001268145 and *Verticillium alfalfae* VaMs.102 from the Ensembl Fungi database (<http://fungi.ensembl.org/info/website/ftp/index.html>) were used and secretome predictions,

HMMER searches against CAZy database and blastp searches against the *MEROPS* database were performed using the same pipeline as for *V. nonalfalfae*. Two way ANNOVA followed by Tukey's multiple comparisons test (p -value < 0.05) in GraphPad Prism 7.03 (GraphPad Software, Inc., USA) was used to find differences between sets of fungal proteins.

5.4 Refinement of *V. nonalfalfae* secretome and selection of CSEPs

A refinement of total *V. nonalfalfae in silico* secretome (Figure 1) was done to maintain only proteins, transcripts of which were expressed *in planta* according to the RNA-Seq analysis. Proteins with carbohydrate enzymatic activities (CAZy screening) were excluded from further analysis and additional filtering was applied based on the presence of PFAM domains. CSEPs were identified as proteins having known effector-specific domains [9,12,13,16,118], NLS signal or as proteins with no PFAM domains.

In the second step, the number of secreted proteins was narrowed down using the following criteria: proteins determined in the lethal pathotype-specific region identified by comparative genomics of mild and lethal *V. nonalfalfae* strains from three geographic regions [22]; proteins differentially expressed in lethal compared to mild *V. nonalfalfae* strains grown in xylem-simulating media as determined by RNA-Seq; *V. nonalfalfae* secreted proteins analyzed by 2D-DIGE and identified by MALDI-TOF/TOF MS [44]; proteins with sequence similarity to experimentally verified pathogenicity, virulence and effector genes in the PHI (Pathogen-host Interaction) database [62] and putative effector proteins predicted by EffectorP software [16].

5.5 Quantitative real time PCR and fungal biomass quantification

Susceptible 'Celeia' and resistant 'Wye Target' hop varieties were inoculated by the root dipping method [119] with *V. nonalfalfae* spores. Total RNA was isolated from the roots and stems of infected plants (6, 12 and 18 dpi) or mock-inoculated plants using a MagMAX total RNA isolation

kit (Life Technologies, USA). The quality and quantity of RNA was assessed on an Agilent 2100 Bioanalyzer (Agilent Technologies, Germany). Total RNA was transcribed to cDNA with a High Capacity cDNA Reverse Transcription kit (Applied Biosystems, USA). Real-time PCR reactions were performed on 7500 Fast Real Time PCR Systems (Life Technologies, USA) using the FastStart SYBR Green master mix (Roche, Switzerland) and primers (Table S8) designed by Primer Express 3.0 software (Thermo Fisher Scientific, USA). For each of the 44 top priority CSEPs, gene expression was analyzed on pooled samples containing the roots of five individual plants, in two technical replicates. The highest expressed CSEPs were also analyzed in five biological and two technical replicates per sample group. The CSEPs' gene expression was calculated by the comparative C_T method [120]. The cDNA from *V. nonalfalfae* mycelium cultivated on ½ CD was used as a reference sample. *V. nonalfalfae* DNA topoisomerase (*VnaUn.148*) and splicing factor 3a2 (*Vna8.801*) were selected as the best endogenous control genes according to GeNorm analysis [121] and fungal biomass normalization. For the latter, fungal DNA was extracted from infected hop using CTAB and quantified by qPCR as described in [122]. The expression of control genes was compared to fungal biomass in infected hop using Pearson's correlation coefficient.

5.6 Construction of knockout vectors and preparation of *V. nonalfalfae* knockout mutants

CSEPs knockout mutants were prepared according to Frandsen's protocol [123]. The plasmid vector pRF-HU2 was first linearized with *Nt.BbvCI* and *PacI* (New England BioLabs, USA) and purified with Illustra GFX PCR DNA and Gel Band Purification Kit (GE Healthcare, UK). Homologous gene sequences were then amplified with PCR using a *PfuTurbo Cx* Hotstart DNA polymerase (Agilent Technologies, USA) and the following settings: 95°C for 2 minutes, 30 cycles: 95°C for 30s, 55°C for 30s, 72°C for 1 minute; 75°C for 10 minutes. PCR products were purified with Illustra GFX PCR DNA and a Gel Band Purification Kit (GE Healthcare, UK) and ligated to linearized vector pRF-

HU2 with USER enzyme (New England BioLabs, USA). Vector constructs were multiplied in *E. coli* DH5α cells and isolated with a High Pure Plasmid Isolation Kit (Roche, Life Science, USA).

V. nonalfalfae knockout mutants were generated with *Agrobacterium tumefaciens* mediated transformation (ATMT) [124] using acetosyringone (Sigma Aldrich, USA). The knockout vector constructs were electroporated with Easyject Prime (EQUIBIO, UK) into electro-competent *Agrobacterium tumefaciens* (LBA4404) cells. Positive colonies with a correct construct orientation were verified by colony PCR. Co-culture of transformed *A. tumefaciens* and *V. nonalfalfae* was carried out on IMAS media as described in [99]. Colonies were transferred on a cellophane membrane (GE Healthcare, UK) to primary and secondary ½ CD selection medium with 150 mg/L timentin (Duchefa, The Netherlands) and 75 mg/L hygromycin (Duchefa, The Netherlands). Genomic DNA was isolated according to [125] from the remaining colonies and the knockout was verified with PCR. Transformed *V. nonalfalfae* conidia were stored in 30% glycerol at -80°C until testing.

5.7 Pathogenicity evaluation of *V. nonalfalfae* knockout mutants in hop

Before pathogenicity tests were carried out, fungal growth and conidiation were inspected as described previously [126]. Ten to fifteen plants of the Verticillium wilt susceptible hop cultivar 'Celeia' were inoculated at phenological stage BBCH 12 by 10-min root dipping in a conidia suspension of *V. nonalfalfae* knockout mutants as described previously [126]. Conidia of the wild type *V. nonalfalfae* lethal pathotype served as a positive control and sterile distilled water was used as a mock control. Verticillium wilting symptoms were assessed four to five times post inoculation using a disease severity index (DSI) with a 0-5 scale [42], and rAUDPC was calculated according to [63]. After symptom assessment, fungal re-isolation test and qPCR using *V. nonalfalfae* specific primers (Table S8) were performed to confirm infection of hops.

5.8 Statistics

The R package [127] was used for the statistical analysis of the pathogenicity assay of knockout mutants. Due to the different variability of rAUDPC values for the 'isolate' groups, a non-parametric approach was pursued. A Kruskal-Wallis test was used, followed by multiple comparison test with Bonferroni correction. To understand how the time post inoculation with *V. nonalfalfae* affects hop health, a simple logistic growth model [128] was fitted to DSI values for the groups under study.

6 List of Abbreviations

AA - proteins with auxiliary activities

ATMT - *Agrobacterium tumefaciens* mediated transformation

CBM - proteins with carbohydrate-binding modules

CE - carbohydrate esterases

CSEPs - candidate secreted effector proteins

CWDEs - cell wall-degrading enzymes

DSI - disease severity index

ETI - effector triggered immunity

GH - glycoside hydrolases

GT - glycosyltransferases

KOG - EuKaryotic Orthologous Groups

NLPs - necrosis and ethylene-inducing protein (NEP-1)-like proteins

NLR - nucleotide-binding leucine rich repeat

555 Nox - NADPH oxidase complex

556 PAMPs - pathogen-associated molecular patterns

557 PHI - Pathogen-host interaction database

558 PL - polysaccharide lyases

559 rAUDPC - relative area under the disease progress curve

560 ROS – reactive oxygen species

561 SSCPs - small secreted cysteine-rich proteins

562 SSPs - small secreted proteins

563 TM - transmembrane domain

564 XSM - xylem simulating medium

565 7 Declarations

566 **Ethics approval and consent to participate:** Not applicable.

567 **Consent for publication:** Not applicable.

568 **Availability of data and materials:** All data generated or analyzed during this study are included in
569 this published article and its supplementary information files. *Verticillium nonalfalfae* genomic data
570 are available at NCBI under BioProject [PRJNA283258](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA283258), while transcriptome data of *Verticillium*
571 *nonalfalfae* interaction with hop can be retrieved from NCBI Bioproject [PRJEB14243](https://www.ncbi.nlm.nih.gov/bioproject/PRJEB14243).

572 **Competing Interests:** The authors declare that they have no competing interests.

573 **Funding:** This work was financed by the Slovenian Research Agency grants P4-0077, J4-8220 and
574 342250.

Authors' Contributions: BJ conceived and coordinated the study and SB participated in its design. JJ performed *Verticillium nonalfalfae* genome assembly and gene model predictions. SR prepared the plant and fungal material and performed hop inoculations. KM and SB performed secretome analysis and RT-qPCR experiments. MF and KM prepared ATMT knockout mutants and performed the pathogenicity assays together with SR. KK performed statistical analysis of data. KM, SB and BJ analyzed and interpreted the data and drafted the manuscript. All authors read and approved the final manuscript.

Acknowledgments: We express our sincere gratitude to Dr. Vasja Progar and the Genialis team for their help with bioinformatics. We acknowledge advice on data normalization in RT-qPCR experiments by Dr. Nataša Štajner from the University of Ljubljana. We thank Martin Cregeen for English language editing.

8 References

1. Inderbitzin P, Bostock RM, Davis RM, Usami T, Platt HW, Subbarao K V. Phylogenetics and taxonomy of the fungal vascular wilt pathogen *Verticillium*, with the descriptions of five new species. PLoS One. Public Library of Science; 2011;6:e28341.
2. Pegg GF, Brady BL. *Verticillium* Wilts. CABI Pub.; 2002.
3. EFSA PLH Panel (EFSA Panel on Plant Health). Scientific opinion on the pest categorisation of *Verticillium albo-atrum* sensu stricto Reinke and Berthold, *V. alfalfae* Inderb., HW Platt, RM Bostock, RM Davis & KV Subbarao, sp. nov., and *V. nonalfalfae* Inderb., HW Platt, RM Bostock, RM Davis & KV Subbar. EFSA J. 2014.

- 596 4. Inderbitzin P, Subbarao K V. *Verticillium* systematics and evolution: How confusion impedes
597 *Verticillium* wilt management and how to resolve it. *Phytopathology*. 2014;104:564–74.
- 598 5. Klimes A, Dobinson KF, Klosterman SJ, Thomma BPHJ. Genomics spurs rapid advances in our
599 understanding of the basic biology of vascular wilt pathogens in the genus *Verticillium*. *Annu. Rev.*
600 *Phytopathol.* 2014;
- 601 6. Daayf F. *Verticillium* wilts in crop plants: Pathogen invasion and host defence responses. *Can. J.*
602 *Plant Pathol.* 2015;37:8–20.
- 603 7. Kamoun S. The secretome of plant-associated fungi and oomycetes. *The Mycota*. Berlin,
604 Heidelberg: Springer Berlin Heidelberg; 2009. p. 173–80.
- 605 8. Lo Presti L, Lanver D, Schweizer G, Tanaka S, Liang L, Tollot M, et al. Fungal effectors and plant
606 susceptibility. *Annu. Rev. Plant Biol. Annual Reviews*; 2015;66:513–45.
- 607 9. Stergiopoulos I, de Wit PJGM. Fungal effector proteins. *Annu. Rev. Phytopathol.* 2009;47:233–63.
- 608 10. de Jonge R. In silico identification and characterization of effector catalogs. *Methods Mol. Biol.*
609 2012. p. 415–25.
- 610 11. Sperschneider J, Williams AH, Hane JK, Singh KB, Taylor JM. Evaluation of secretion
611 prediction highlights differing approaches needed for oomycete and fungal effectors. *Front. Plant Sci.*
612 *Frontiers Media SA*; 2015;6:1168.
- 613 12. Saunders DGO, Win J, Cano LM, Szabo LJ, Kamoun S, Raffaele S. Using hierarchical clustering
614 of secreted protein families to classify and rank candidate effectors of rust fungi. *PLoS One*.
615 2012;7:e29847.
- 616 13. Guyon K, Balagué C, Roby D, Raffaele S. Secretome analysis reveals effector candidates

- 617 associated with broad host range necrotrophy in the fungal plant pathogen *Sclerotinia sclerotiorum*.
- 618 BMC Genomics. 2014;15:336.
- 619 14. Sonah H, Deshmukh RK, Bélanger RR. Computational prediction of effector proteins in fungi:
620 Opportunities and challenges. Front. Plant Sci. 2016;7.
- 621 15. Gibriel HAY, Thomma BPHJ, Seidl MF. The age of effectors: Genome-based discovery and
622 applications. Phytopathology. 2016;106:1206–12.
- 623 16. Sperschneider J, Gardiner DM, Dodds PN, Tini F, Covarelli L, Singh KB, et al. EffectorP:
624 predicting fungal effector proteins from secretomes using machine learning. New Phytol.
625 2016;210:743–61.
- 626 17. Klosterman SJ, Subbarao K V, Kang S, Veronese P, Gold SE, Thomma BPHJ, et al. Comparative
627 genomics yields insights into niche adaptation of plant vascular wilt pathogens. PLoS Pathog. Public
628 Library of Science; 2011;7:e1002137.
- 629 18. De Jonge R, Bolton MD, Kombrink A, Van Den Berg GCM, Yadeta KA, Thomma BPHJ.
630 Extensive chromosomal reshuffling drives evolution of virulence in an asexual pathogen. Genome
631 Res. Cold Spring Harbor Laboratory Press; 2013;23:1271–82.
- 632 19. Seidl MF, Faino L, Shi-Kunne X, van den Berg GCM, Bolton MD, Thomma BPHJ. The genome
633 of the saprophytic fungus *Verticillium tricorpus* reveals a complex effector repertoire resembling that
634 of its pathogenic relatives. Mol. Plant-Microbe Interact. The American Phytopathological Society;
635 2015;28:362–73.
- 636 20. Faino L, Seidl MF, Datema E, van den Berg GCM, Janssen A, Wittenberg AHJ, et al. Single-
637 molecule real-time sequencing combined with optical mapping yields completely finished fungal
638 genome. MBio. American Society for Microbiology; 2015;6:e00936-15.

- 639 21. Depotter JRL, Seidl MF, van den Berg GCM, Thomma BPHJ, Wood TA. A distinct and
640 genetically diverse lineage of the hybrid fungal pathogen *Verticillium longisporum* population causes
641 stem striping in British oilseed rape. *Environ. Microbiol.* 2017;19:3997–4009.
- 642 22. Jakše J, Jelen V, Radišek S, de Jonge R, Mandelc S, Majer A, et al. Genome sequence of xylem-
643 invading *Verticillium nonalfalfae* lethal strain. *Genome Announc.* 2018;6:e01458-17.
- 644 23. Sánchez-Vallet A, Saleem-Batcha R, Kombrink A, Hansen G, Valkenburg D-J, Thomma BPHJ,
645 et al. Fungal effector Ecp6 outcompetes host immune receptor for chitin binding through intrachain
646 LysM dimerization. *Elife.* 2013;2:e00790.
- 647 24. Laugé R, Joosten MHAJ, Van den Ackerveken GFJM, Van den Broek HWJ, De Wit PJGM. The
648 In planta-produced extracellular proteins ECP1 and ECP2 of *Cladosporium fulvum* are virulence
649 factors. *Mol. Plant-Microbe Interact. The American Phytopathological Society* ; 1997;10:725–34.
- 650 25. Mentlak TA, Kombrink A, Shinya T, Ryder LS, Otomo I, Saitoh H, et al. Effector-mediated
651 suppression of chitin-triggered immunity by *Magnaporthe oryzae* is necessary for rice blast disease.
652 *Plant Cell.* 2012;24:322–35.
- 653 26. Takahara H, Hacquard S, Kombrink A, Hughes HB, Halder V, Robin GP, et al. *Colletotrichum*
654 *higginsianum* extracellular LysM proteins play dual roles in appressorial function and suppression of
655 chitin-triggered plant immunity. *New Phytol.* 2016;211:1323–37.
- 656 27. Kombrink A, Rovenich H, Shi-Kunne X, Rojas-Padilla E, van den Berg GCM, Domazakis E, et
657 al. *Verticillium dahliae* LysM effectors differentially contribute to virulence on plant hosts. *Mol.*
658 *Plant Pathol.* 2017;18:596–608.
- 659 28. de Jonge R, Peter van Esse H, Kombrink A, Shinya T, Desaki Y, Bours R, et al. Conserved
660 fungal LysM effector Ecp6 prevents chitin-triggered immunity in plants. *Science* (80-.).

- 2010;329:953–5.
29. Zhou B-J, Jia P-S, Gao F, Guo H-S. Molecular characterization and functional analysis of a Necrosis- and ethylene-Inducing, protein-encoding gene family from *Verticillium dahliae*. Mol. Plant-Microbe Interact. The American Phytopathological Society; 2012;25:964–75.
30. Santhanam P, van Esse HP, Albert I, Faino L, Nürnberger T, Thomma BPHJ. Evidence for functional diversification within a fungal NEP1-like protein family. Mol. Plant. Microbe. Interact. 2013;26:278–86.
31. de Jonge R, van Esse HP, Maruthachalam K, Bolton MD, Santhanam P, Saber MK, et al. Tomato immune receptor Ve1 recognizes effector of multiple fungal pathogens uncovered by genome and RNA sequencing. Proc. Natl. Acad. Sci. National Acad Sciences; 2012;109:5110–5.
32. Fradin EF, Abd-El-Halim A, Masini L, van den Berg GCM, Joosten MHAJ, Thomma BPHJ. Interfamily transfer of tomato Ve1 mediates *Verticillium* resistance in *Arabidopsis*. Plant Physiol. 2011;156:2255–65.
33. Fradin EF, Zhang Z, Rovenich H, Song Y, Liebrand TWH, Masini L, et al. Functional analysis of the tomato immune receptor Ve1 through domain swaps with its non-functional homolog Ve2. PLoS One. 2014;9:e88208.
34. Liu W, Zeng H, Liu Z, Yang X, Guo L, Qiu D. Mutational analysis of the *Verticillium dahliae* protein elicitor PevD1 identifies distinctive regions responsible for hypersensitive response and systemic acquired resistance in tobacco. Microbiol. Res. Elsevier GmbH.; 2014;169:476–82.
35. Wang B, Yang X, Zeng H, Liu H, Zhou T, Tan B, et al. The purification and characterization of a novel hypersensitive-like response-inducing elicitor from *Verticillium dahliae* that induces resistance responses in tobacco. Appl. Microbiol. Biotechnol. 2012;93:191–201.

- 683 36. Bu B, Qiu D, Zeng H, Guo L, Yuan J, Yang X. A fungal protein elicitor PevD1 induces
684 Verticillium wilt resistance in cotton. Plant Cell Rep. 2014;33:461–70.
- 685 37. Zhang L, Ni H, Du X, Wang S, Ma XW, Nürnberger T, et al. The *Verticillium*-specific protein
686 VdSCP7 localizes to the plant nucleus and modulates immunity to fungal infections. New Phytol.
687 2017;215:368–81.
- 688 38. Flajšman M, Mandelc S, Radišek S, Štajner N, Jakše J, Košmelj K, et al. Identification of novel
689 virulence-associated proteins secreted to xylem by *Verticillium nonalfalfae* during colonization of
690 hop plants. Mol. Plant. Microbe. Interact. 2016;29:362–73.
- 691 39. Keyworth WG. Verticillium wilt of the hop (*Humulus lupulus*). Ann. Appl. Biol. Blackwell
692 Publishing Ltd; 1942;29:346–57.
- 693 40. Sewell GWF, Wilson JF. The nature and distribution of *Verticillium albo-atrum* strains highly
694 pathogenic to the hop. Plant Pathol. Wiley Online Library; 1984;33:39–51.
- 695 41. Talboys PW. Resistance to vascular wilt fungi. Proc. R. Soc. London. Ser. B. Biol. Sci. The
696 Royal Society; 1972;181:319–32.
- 697 42. Radišek S, Jakše J, Simončič A, Javornik B. Characterization of *Verticillium albo-atrum* field
698 isolates using pathogenicity data and AFLP analysis. Plant Dis. 2003;87:633–8.
- 699 43. Radišek S, Jakše J, Javornik B. Genetic variability and virulence among *Verticillium albo-atrum*
700 isolates from hop. Eur. J. plant Pathol. Springer; 2006;116:301–14.
- 701 44. Mandelc S, Javornik B. The secretome of vascular wilt pathogen *Verticillium albo-atrum* in
702 simulated xylem fluid. Proteomics. 2015;15:787–97.
- 703 45. Progar V, Jakše J, Štajner N, Radišek S, Javornik B, Berne S. Comparative transcriptional

- 704 analysis of hop responses to infection with *Verticillium nonalfalfae*. Plant Cell Rep. Springer Berlin
705 Heidelberg; 2017;36:1599–613.
- 706 46. Käll L, Krogh A, Sonnhammer ELL. A combined transmembrane topology and signal peptide
707 prediction method. J. Mol. Biol. 2004;338:1027–36.
- 708 47. Horton P, Park K-J, Obayashi T, Fujita N, Harada H, Adams-Collier CJ, et al. WoLF PSORT:
709 protein localization predictor. Nucleic Acids Res. 2007;35:W585-7.
- 710 48. Sperschneider J, Dodds PN, Singh KB, Taylor JM. ApoplastP: prediction of effectors and plant
711 proteins in the apoplast using machine learning. doi.org. Cold Spring Harbor Laboratory;
712 2017;182428.
- 713 49. Sperschneider J, Catanzariti A-M, DeBoer K, Petre B, Gardiner DM, Singh KB, et al.
714 LOCALIZER: subcellular localization prediction of both plant and effector proteins in the plant cell.
715 Sci. Rep. Nature Publishing Group; 2017;7:44598.
- 716 50. Wilson D, Pethica R, Zhou Y, Talbot C, Vogel C, Madera M, et al. SUPERFAMILY--
717 sophisticated comparative genomics, data mining, visualization and phylogeny. Nucleic Acids Res.
718 Oxford University Press; 2009;37:D380-6.
- 719 51. Macdonald SS, Blaukopf M, Withers SG. *N*-acetylglucosaminidases from CAZy family GH3 are
720 really glycoside phosphorylases, thereby explaining their use of histidine as an acid/base catalyst in
721 place of glutamic acid. J. Biol. Chem. 2015;290:4887–95.
- 722 52. Aspeborg H, Coutinho PM, Wang Y, Brumer H, Henrissat B. Evolution, substrate specificity and
723 subfamily classification of glycoside hydrolase family 5 (GH5). BMC Evol. Biol. 2012;12:186.
- 724 53. Mewis K, Lenfant N, Lombard V, Henrissat B. Dividing the large glycoside hydrolase family 43

- 725 into subfamilies: a motivation for detailed enzyme characterization. Nojiri H, editor. Appl. Environ.
726 Microbiol. 2016;82:1686–92.
- 727 54. Akcapinar GB, Kappel L, Sezerman OU, Seidl-Seiboth V. Molecular diversity of LysM
728 carbohydrate-binding motifs in fungi. Curr. Genet. 2015;61:103–13.
- 729 55. Krijger J-J, Thon MR, Deising HB, Wiersel SG. Compositions of fungal secretomes indicate a
730 greater impact of phylogenetic history than lifestyle adaptation. BMC Genomics. 2014;15:722.
- 731 56. Gijzen M, Nürnberger T. Nep1-like proteins from plant pathogens: recruitment and
732 diversification of the NPP1 domain across taxa. Phytochemistry. 2006;67:1800–7.
- 733 57. Stergiopoulos I, Kourmpetis YAI, Slot JC, Bakker FT, De Wit PJGM, Rokas A. In silico
734 characterization and molecular evolutionary analysis of a novel superfamily of fungal effector
735 proteins. Mol. Biol. Evol. 2012;29:3371–84.
- 736 58. Baccelli I. Cerato-platanin family proteins: one function for multiple biological roles? Front.
737 Plant Sci. Frontiers Media SA; 2014;5:769.
- 738 59. Koharudin LMI, Viscomi AR, Montanini B, Kershaw MJ, Talbot NJ, Ottonello S, et al.
739 Structure-function analysis of a CVNH-LysM lectin expressed during plant infection by the rice blast
740 fungus *Magnaporthe oryzae*. Structure. NIH Public Access; 2011;19:662–74.
- 741 60. Bayry J, Aïmanianda V, Guijarro JI, Sunde M, Latgé J-P. Hydrophobins - unique fungal proteins.
742 PLoS Pathog. Public Library of Science; 2012;8:e1002700.
- 743 61. Kulkarni RD, Kelkar HS, Dean RA. An eight-cysteine-containing CFEM domain unique to a
744 group of fungal membrane proteins. Trends Biochem. Sci. 2003;28:118–21.
- 745 62. Urban M, Pant R, Raghunath A, Irvine AG, Pedro H, Hammond-Kosack KE. The Pathogen-Host

746 Interactions database (PHI-base): additions and future developments. Nucleic Acids Res.
747 2015;43:D645–55.

748 63. Simko I, Piepho H-P. The area under the disease progress stairs: calculation, advantage, and
749 application. Phytopathology. 2012;102:381–9.

750 64. Zhao Y-L, Zhou T-T, Guo H-S. Hyphopodium-specific VdNoxB/VdPls1-dependent ROS-Ca²⁺
751 signaling is required for plant infection by *Verticillium dahliae*. Wilson RA, editor. PLOS Pathog.
752 2016;12:e1005793.

753 65. Klosterman SJ, Subbarao K V, Kang S, Veronese P, Gold SE, Thomma BPHJ, et al. Comparative
754 genomics yields insights into niche adaptation of plant vascular wilt pathogens. Dangl JL, editor.
755 PLoS Pathog. Public Library of Science; 2011;7:e1002137.

756 66. Di Pietro A, Roncero MIG, Roldán MCR. From tools of survival to weapons of destruction: The
757 role of cell wall-degrading enzymes in plant infection. In: Deising HB, editor. Plant Relationships.
758 Berlin, Heidelberg: Springer Berlin Heidelberg; 2009. p. 181–200.

759 67. Kubicek CP, Starr TL, Glass NL. Plant cell wall-degrading enzymes and their secretion in plant-
760 pathogenic fungi. Annu. Rev. Phytopathol. 2014;52:427–51.

761 68. Ma Z, Song T, Zhu L, Ye W, Wang Y, Shao Y, et al. A *Phytophthora sojae* glycoside hydrolase
762 12 protein is a major virulence factor during soybean infection and is recognized as a PAMP. Plant
763 Cell. 2015;27:2057–72.

764 69. Zhang L, Kars I, Essenstam B, Liebrand TWH, Wagemakers L, Elberse J, et al. Fungal
765 Endopolygalacturonases are recognized as microbe-associated molecular patterns by the *Arabidopsis*
766 receptor-like protein RESPONSIVENESS TO BOTRYTIS POLYGALACTURONASES1. Plant
767 Physiol. American Society of Plant Biologists; 2014;164:352–64.

- 768 70. Gui Y-J, Chen J-Y, Zhang D-D, Li N-Y, Li T-G, Zhang W-Q, et al. *Verticillium dahliae*
769 manipulates plant immunity by glycoside hydrolase 12 proteins in conjunction with carbohydrate-
770 binding module 1. Environ. Microbiol. 2017;19:1914–32.
- 771 71. Jashni MK, Mehrabi R, Collemare J, Mesarich CH, de Wit PJGM. The battle in the apoplast:
772 further insights into the roles of proteases and their inhibitors in plant–pathogen interactions. Front.
773 Plant Sci. 2015;6:584.
- 774 72. Chandrasekaran M, Thangavelu B, Chun SC, Sathiyabama M. Proteases from phytopathogenic
775 fungi and their importance in phytopathogenicity. J. Gen. Plant Pathol. 2016;82:233–9.
- 776 73. Zeilinger S, Gupta VK, Dahms TES, Silva RN, Singh HB, Upadhyay RS, et al. Friends or foes?
777 Emerging insights from fungal interactions with plants. FEMS Microbiol. Rev. 2016;40.
- 778 74. Serrano M, Coluccia F, Torres M, L’Haridon F, Métraux J-P. The cuticle and plant defense to
779 pathogens. Front. Plant Sci. Frontiers; 2014;5:274.
- 780 75. Blümke A, Falter C, Herrfurth C, Sode B, Bode R, Schäfer W, et al. Secreted fungal effector
781 lipase releases free fatty acids to inhibit innate immunity-related callose formation during wheat head
782 infection. Plant Physiol. American Society of Plant Biologists; 2014;165:346–58.
- 783 76. Xie C, Wang C, Wang X, Yang X. Proteomics-based analysis reveals that *Verticillium dahliae*
784 toxin induces cell death by modifying the synthesis of host proteins. J. Gen. Plant Pathol. Springer
785 Japan; 2013;79:335–45.
- 786 77. Gayoso C, Pomar F, Novo-Uzal E, Merino F, de Ilárduya OM. The Ve-mediated resistance
787 response of the tomato to *Verticillium dahliae* involves H₂O₂, peroxidase and lignins and drives PAL
788 gene expression. BMC Plant Biol. 2010;10:232.

- 789 78. Heller J, Tudzynski P. Reactive oxygen species in phytopathogenic fungi: Signaling,
790 development, and disease. Annu. Rev. Phytopathol. 2011;49:369–90.
- 791 79. Zhou T-T, Zhao Y-L, Guo H-S, Kikuchi S, Arioka M. Secretory proteins are delivered to the
792 septin-organized penetration interface during root infection by *Verticillium dahliae*. Talbot NJ,
793 editor. PLOS Pathog. Public Library of Science; 2017;13:e1006275.
- 794 80. Ma L-J, van der Does HC, Borkovich KA, Coleman JJ, Daboussi M-J, Di Pietro A, et al.
795 Comparative genomics reveals mobile pathogenicity chromosomes in *Fusarium*. Nature.
796 2010;464:367–73.
- 797 81. Gan P, Ikeda K, Irieda H, Narusaka M, O’Connell RJ, Narusaka Y, et al. Comparative genomic
798 and transcriptomic analyses reveal the hemibiotrophic stage shift of *Colletotrichum* fungi. New
799 Phytol. 2013;197:1236–49.
- 800 82. Kim K-T, Jeon J, Choi J, Cheong K, Song H, Choi G, et al. Kingdom-wide analysis of fungal
801 small secreted proteins (SSPs) reveals their potential role in host association. Front. Plant Sci.
802 Frontiers Media SA; 2016;7:186.
- 803 83. Klimes A, Dobinson KF. A hydrophobin gene, VDH1, is involved in microsclerotial
804 development and spore viability in the plant pathogen *Verticillium dahliae*. Fungal Genet. Biol.
805 2006;43:283–94.
- 806 84. Jia Y, Zhou E, Lee S, Bianco T. Coevolutionary dynamics of rice blast resistance gene *Pi-ta* and
807 *Magnaporthe oryzae* avirulence gene *AVR-Pita 1*. Phytopathology. Phytopathology; 2016;106:676–
808 83.
- 809 85. Yang Y, Zhang H, Li G, Li W, Wang X, Song F. Ectopic expression of MgSM1, a Cerato-
810 platanin family protein from *Magnaporthe grisea* , confers broad-spectrum disease resistance in

811 *Arabidopsis*. Plant Biotechnol. J. 2009;7:763–77.

812 86. Röhm M, Lindemann E, Hiller E, Ermert D, Lemuth K, Trkulja D, et al. A family of secreted
813 pathogenesis-related proteins in *Candida albicans*. Mol. Microbiol. 2013;87:132–51.

814 87. Navarathna DHMLP, Lionakis MS, Lizak MJ, Munasinghe J, Nickerson KW, Roberts DD. Urea
815 amidolyase (DUR1,2) contributes to virulence and kidney pathogenesis of *Candida albicans*.
816 Chauhan N, editor. PLoS One. 2012;7:e48475.

817 88. Ahn N, Kim S, Choi W, Im K-H, Lee Y-H. Extracellular matrix protein gene, EMP1, is required
818 for appressorium formation and pathogenicity of the rice blast fungus, *Magnaporthe grisea*. Mol.
819 Cells. 2004;17:166–73.

820 89. Perez-Nadales E, Di Pietro A. The membrane mucin Msb2 regulates invasive growth and plant
821 infection in *Fusarium oxysporum*. Plant Cell. 2011;23:1171–85.

822 90. Bok JW, Keller NP. LaeA, a regulator of secondary metabolism in *Aspergillus* spp. Eukaryot.
823 Cell. 2004;3:527–35.

824 91. Bok JW, Balajee SA, Marr KA, Andes D, Nielsen KF, Frisvad JC, et al. LaeA, a regulator of
825 morphogenetic fungal virulence factors. Eukaryot. Cell. American Society for Microbiology (ASM);
826 2005;4:1574–82.

827 92. Sarikaya-Bayram Â, Palmer JM, Keller N, Braus GH, Bayram Â. One Juliet and four Romeos:
828 VeA and its methyltransferases. Front. Microbiol. 2015;6:1.

829 93. Qin Y, Ortiz-Urquiza A, Keyhani NO. A putative methyltransferase, mtrA, contributes to
830 development, spore viability, protein secretion and virulence in the entomopathogenic fungus
831 *Beauveria bassiana*. Microbiology. 2014;160:2526–37.

- 832 94. Dodds PN, Rathjen JP. Plant immunity: towards an integrated view of plant-pathogen
833 interactions. Nat. Rev. Genet. 2010;11:539–48.
- 834 95. Cui H, Tsuda K, Parker JE. Effector-triggered immunity: from pathogen perception to robust
835 defense. Annu. Rev. Plant Biol. 2015;66:487–511.
- 836 96. Vleeshouwers VGAA, Oliver RP. Effectors as tools in disease resistance breeding against
837 biotrophic, hemibiotrophic, and necrotrophic plant pathogens. Mol. Plant-Microbe Interact. 27:196–
838 206.
- 839 97. Radišek S, Jakše J, Javornik B. Development of pathotype-specific SCAR markers for detection
840 of *Verticillium albo-atrum* isolates from hop. Plant Dis. 2004;88:1115–22.
- 841 98. Neumann MJ, Dobinson KF. Sequence tag analysis of gene expression during pathogenic growth
842 and microsclerotia development in the vascular wilt pathogen *Verticillium dahliae*. Fungal Genet.
843 Biol. 2003;38:54–62.
- 844 99. Frandsen RJN, Frandsen M, Giese H. Plant Fungal Pathogens. 2012;835:17–45.
- 845 100. Gasteiger E, Gattiker A, Hoogland C, Ivanyi I, Appel DR, Bairoch A. ExPASy: the proteomics
846 server for in-depth protein knowledge and analysis. Nucleic Acids Res. 2003;31:3784–8.
- 847 101. Wheeler DL, Church DM, Federhen S, Lash AE, Madden TL, Pontius JU, et al. Database
848 resources of the National Center for Biotechnology. Nucleic Acids Res. 2003;31:28–33.
- 849 102. Finn RD, Clements J, Eddy SR. HMMER web server: interactive sequence similarity searching.
850 Nucleic Acids Res. 2011;39:W29–37.
- 851 103. Cantarel BL, Coutinho PM, Rancurel C, Bernard T, Lombard V, Henrissat B. The
852 Carbohydrate-Active EnZymes database (CAZy): an expert resource for glycogenomics. Nucleic

853 Acids Res. 2009;37:D233-8.

854 104. Lombard V, Golaconda Ramulu H, Drula E, Coutinho PM, Henrissat B. The carbohydrate-
855 active enzymes database (CAZy) in 2013. Nucleic Acids Res. 2014;42:D490-5.

856 105. Finn RD, Bateman A, Clements J, Coggill P, Eberhardt RY, Eddy SR, et al. Pfam: The protein
857 families database. Nucleic Acids Res. 2014;42:222–30.

858 106. Gough J, Karplus K, Hughey R, Chothia C. Assignment of homology to genome sequences
859 using a library of hidden Markov models that represent all proteins of known structure. J. Mol. Biol.
860 2001;313:903–19.

861 107. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J.
862 Mol. Biol. 1990;215:403–10.

863 108. Tatusov RL, Fedorova ND, Jackson JD, Jacobs AR, Kiryutin B, Koonin E V, et al. The COG
864 database: an updated version includes eukaryotes. BMC Bioinformatics. 2003;4:41.

865 109. Rawlings ND, Barrett AJ, Bateman A. MEROPS: the database of proteolytic enzymes, their
866 substrates and inhibitors. Nucleic Acids Res. 2012;40:D343–50.

867 110. Conesa A, Götz S, García-Gómez JM, Terol J, Talón M, Robles M. Blast2GO: a universal tool
868 for annotation, visualization and analysis in functional genomics research. Bioinformatics.
869 2005;21:3674–6.

870 111. Kanehisa M, Goto S. KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acids Res.
871 2000;28:27–30.

872 112. Kanehisa M, Goto S, Sato Y, Kawashima M, Furumichi M, Tanabe M. Data, information,
873 knowledge and principle: back to metabolism in KEGG. Nucleic Acids Res. 2014;42:D199-205.

- 874 113. Min XJ. Evaluation of computational methods for secreted protein prediction in different
875 eukaryotes. 2010;3:143–7.
- 876 114. Petersen TN, Brunak S, von Heijne G, Nielsen H. SignalP 4.0: discriminating signal peptides
877 from transmembrane regions. Nat. Methods. Nature Publishing Group, a division of Macmillan
878 Publishers Limited. All Rights Reserved.; 2011;8:785–6.
- 879 115. de Castro E, Sigrist CJA, Gattiker A, Bulliard V, Langendijk-Genevaux PS, Gasteiger E, et al.
880 ScanProsite: detection of PROSITE signature matches and ProRule-associated functional and
881 structural residues in proteins. Nucleic Acids Res. 2006;34:W362–5.
- 882 116. Brameier M, Krings A, MacCallum RM. NucPred--predicting nuclear localization of proteins.
883 Bioinformatics. 2007;23:1159–60.
- 884 117. Cokol M, Nair R, Rost B. Finding nuclear localization signals. EMBO Rep. 2000;1:411–5.
- 885 118. Heard S, Brown NA, Hammond-Kosack K. An interspecies comparative analysis of the
886 predicted secretomes of the necrotrophic plant pathogens *Sclerotinia sclerotiorum* and *Botrytis*
887 *cinerea*. PLoS One. 2015;10:e0130534.
- 888 119. Mandelc S, Timperman I, Radišek S, Devreese B, Samyn B, Javornik B. Comparative
889 proteomic profiling in compatible and incompatible interactions between hop roots and *Verticillium*
890 *albo-atrum*. Plant Physiol Biochem. 2013;68:23–31.
- 891 120. Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the comparative C(T) method. Nat.
892 Protoc. 2008;3:1101–8.
- 893 121. Schlotter YM, Veenhof EZ, Brinkhof B, Rutten VPMG, Spee B, Willemse T, et al. A GeNorm
894 algorithm-based selection of reference genes for quantitative real-time PCR in skin biopsies of

895 healthy dogs and dogs with atopic dermatitis. Vet. Immunol. Immunopathol. 2009;129:115–8.

896 122. Cregeen S, Radišek S, Mandelc S, Turk B, Štajner N, Jakše J, et al. Different gene expressions
897 of resistant and susceptible hop cultivars in response to infection with a highly aggressive strain of
898 *Verticillium albo-atrum*. Plant Mol. Biol. Rep. 2015;33:689–704.

899 123. Frandsen RJN, Andersson J a, Kristensen MB, Giese H. Efficient four fragment cloning for the
900 construction of vectors for targeted gene replacement in filamentous fungi. BMC Mol. Biol.
901 2008;9:70.

902 124. Knight CJ, Bailey AM, Foster GD. *Agrobacterium*-mediated transformation of the plant
903 pathogenic fungus *Verticillium albo-atrum*. J. Plant Pathol. 2009;91:745–50.

904 125. Möller EM, Bahnweg G, Sandermann H, Geiger HH. A simple and efficient protocol for
905 isolation of high molecular weight DNA from filamentous fungi, fruit bodies, and infected plant
906 tissues. Nucleic Acids Res. 1992;20:6115–6.

907 126. Flajšman M, Radišek S, Javornik B. Pathogenicity assay of *Verticillium nonalfalfae* on hop
908 plants. Bio-protocol. 2017;7:e2171.

909 127. R Core Team. R: A language and environment for statistical computing. Vienna, Austria; 2016.

910 128. Pinheiro JC, Bates DM. Mixed-effects models in S and S-PLUS. Springer; 2000.

911 129. Robinson MD, Oshlack A. A scaling normalization method for differential expression analysis
912 of RNA-seq data. Genome Biol. BioMed Central; 2010;11:R25.

913 **9 Figure captions**

Figure 1. Bioinformatics pipeline for secretome prediction, identification and characterization of *V. nonalfalfae* candidate secreted effector proteins (CSEPs). *V. nonalfalfae* predicted proteome was first filtered based on signal peptide, transmembrane domains and subcellular localization to determine classically secreted proteins. This total predicted secretome was then refined to proteins expressed in *planta* and carbohydrate active enzymes were removed so that only proteins with effector-specific PFAM domains, NLS signal or no PFAM domains were retained in the final dataset of 263 CSEPs. These were narrowed down to 44 top-priority candidates based on the results of RNA-Seq and 2D-DIGE analyses, sequence similarity searching to known effectors in the PHI database and EffectorP prediction.

Figure 2. Classification of the *V. nonalfalfae* predicted secreted proteins into functional groups based on the Superfamily annotation scheme. Groups were classified after [50].

Figure 3. Relative abundance of carbohydrate-active enzymes (CAZymes), peptidases and small secreted proteins within secretomes of plant pathogenic *Verticillium* species. (A) Comparison of CAZymes from different classes is presented in percentages of predicted fungal secretome. CE, carbohydrate esterase; GH, glycoside hydrolase; GT, glycosyl transferase; PL, polysaccharide lyase; CBM, proteins with carbohydrate-binding modules; AA, proteins with auxiliary activities. (B) Comparison of various classes of peptidases depicted as percentage of predicted fungal secretome. (C) Relative abundance of small secreted proteins and small secreted cysteine rich proteins in predicted fungal secretomes. SSPs, small secreted proteins with less than 300 aa; SSCPs, small secreted proteins with more than 5% cysteine content and at least 4 Cys residues [55]; *Vna*, *V. nonalfalfae*; *Va*, *V. alfalfae*; *Vd*, *V. dahliae*; *Vl*, *V. longisporum*.

Figure 4. Gene expression profiles of selected *V. nonalfalfae* CSEPs in the roots and shoots of infected hop. FC, fold change in gene expression was determined by quantitative real-time PCR using topoisomerase and splicing factor 3a2 as endogenous controls and fungal samples grown on ½

CD medium as a reference. Means $\pm$ SEM (n=5) are presented. Statistical significance was determined with the t-test using the Holm-Sidak approach, with $\alpha = 5\%$. 'Celeia', *Verticillium* wilt susceptible hop; 'Wye Target', *Verticillium* wilt resistant hop; dpi, days post inoculation

Figure 5. Pathogenicity testing in susceptible hop 'Celeia'. Pictures were taken 31 days post inoculation of hop roots with *V. nonalfalfae* conidia suspension. (A) mock-inoculated plants; (B) plants inoculated with wild type *V. nonalfalfae* T2 strain; (C and D), plants inoculated with *V. nonalfalfae* $\Delta VnaUn.279$ mutant strains (two replicates).

Figure 6. *V. nonalfalfae* knockout mutant $\Delta VnaUn.279$ showed severely reduced virulence. (A) Relative area under the disease progress curve (rAUDPC) is presented for knockout mutant $\Delta VnaUn.279$ and for wild type (wt) *V. nonalfalfae*. Darker dots depict double values. A Kruskal-Wallis test followed by multiple comparison test resulted in two groups: *a* for $\Delta VnaUn.279$ and *b* for wild type. (B) Disease severity index (DSI) values were fitted by a simple logistic growth model for mutant $\Delta VnaUn.279$ and for wild type *V. nonalfalfae* hop isolates. The upper horizontal line is the asymptote (black for $\Delta VnaUn.279$, red for wt); the vertical lines show the inflection points (black for $\Delta VnaUn.279$, red for wt) at which the predicted DSI is one half of the asymptote.

10 Additional files

Additional file 1. An Excel file containing 8 tables, each in a separate worksheet.

Table S1. *V. nonalfalfae* secretome predicted using Genialis bioinformatics platform. Gene IDs marked in bold represent 263 proteins in the CSEPs catalogue. 2D-DIGE, proteins secreted by mild and lethal strains of *V. nonalfalfae* growing in xylem simulating medium (XSM) were analysed by 2D-DIGE and identified by MALDI-TOF/TOF MS [44]; RNA-Seq, differential fungal gene expression (fold change, $FC \geq 1.5$ or $FC \leq -1.5$) of *V. nonalfalfae* lethal versus mild fungal pathotype growing in XSM; LOCALIZER, a tool for subcellular localization prediction of both plant and

effector proteins in the plant cell [49]; ApoplastP, a tool for prediction of effectors and plant proteins in the apoplast using machine learning [48].

Table S2. KOG functional annotation of *V. nonalfalfae* predicted secretome. KOG, a database of euKaryotic Orthologous Groups from NCBI that allows identification of ortholog and paralog proteins [108].

Table S3. *MEROPS* classification of predicted *V. nonalfalfae* secretome. *MEROPS*, a database of peptidases and the proteins that inhibit them [109].

Table S4. Assignment of KEGG accessions to the predicted *V. nonalfalfae* secreted proteins. KEGG, the Kyoto Encyclopedia of Genes and Genomes is a database resource for understanding high-level functions and utilities of the biological systems [111].

Table S5. List of PFAM domains identified in the predicted *V. nonalfalfae* secretome. PFAM, a collection of protein families, represented by multiple sequence alignments and hidden Markov models (HMMs) [105].

Table S6. List of *V. nonalfalfae* CSEPs, including EffectorP prediction and similarity to effectors in the PHI database. EffectorP, a machine learning prediction program for fungal effectors [16]; PHI, Pathogen-Host Interaction database, which contains expertly curated molecular and biological information on genes proven to affect the outcome of pathogen-host interactions [62].

Table S7. Bulk expression of best ranked *V. nonalfalfae* candidate effector genes in hop roots. Gene expression was calculated by the comparative C_T method [120]. Hop plants were inoculated by the root dipping method with *V. nonalfalfae* conidia and sampled at 6, 12 and 18 days post inoculation. Analyzed samples contained the roots of five individual plants from either susceptible hop variety 'Celeia' (CE) or resistant 'Wye Target' (WT). cDNA from *V. nonalfalfae* mycelium cultivated on ½ Czapek Dox (CD) medium was used as a reference sample. *V. nonalfalfae* DNA topoisomerase (*VnaUn.148*) and splicing factor 3a2 (*Vna8.801*) were used as endogenous control genes. Numbers

985 represent log₂ fold changes in the expression of genes in infected plants at indicated time points
 986 compared to gene expression in ½ CD medium. *na*, not available

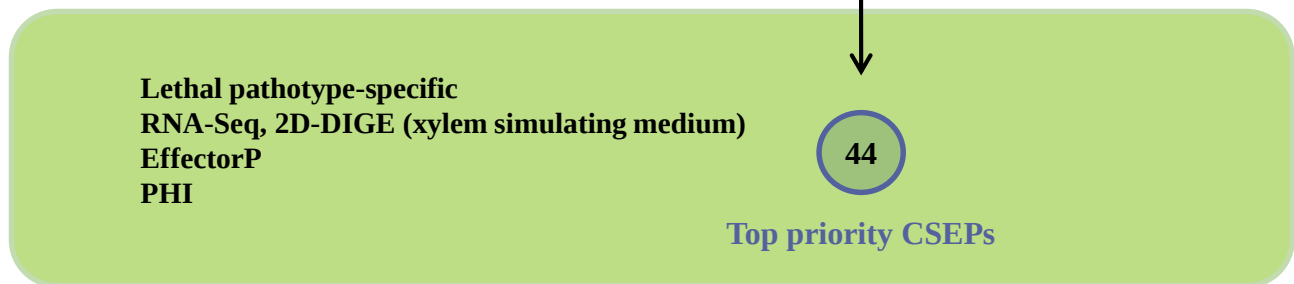
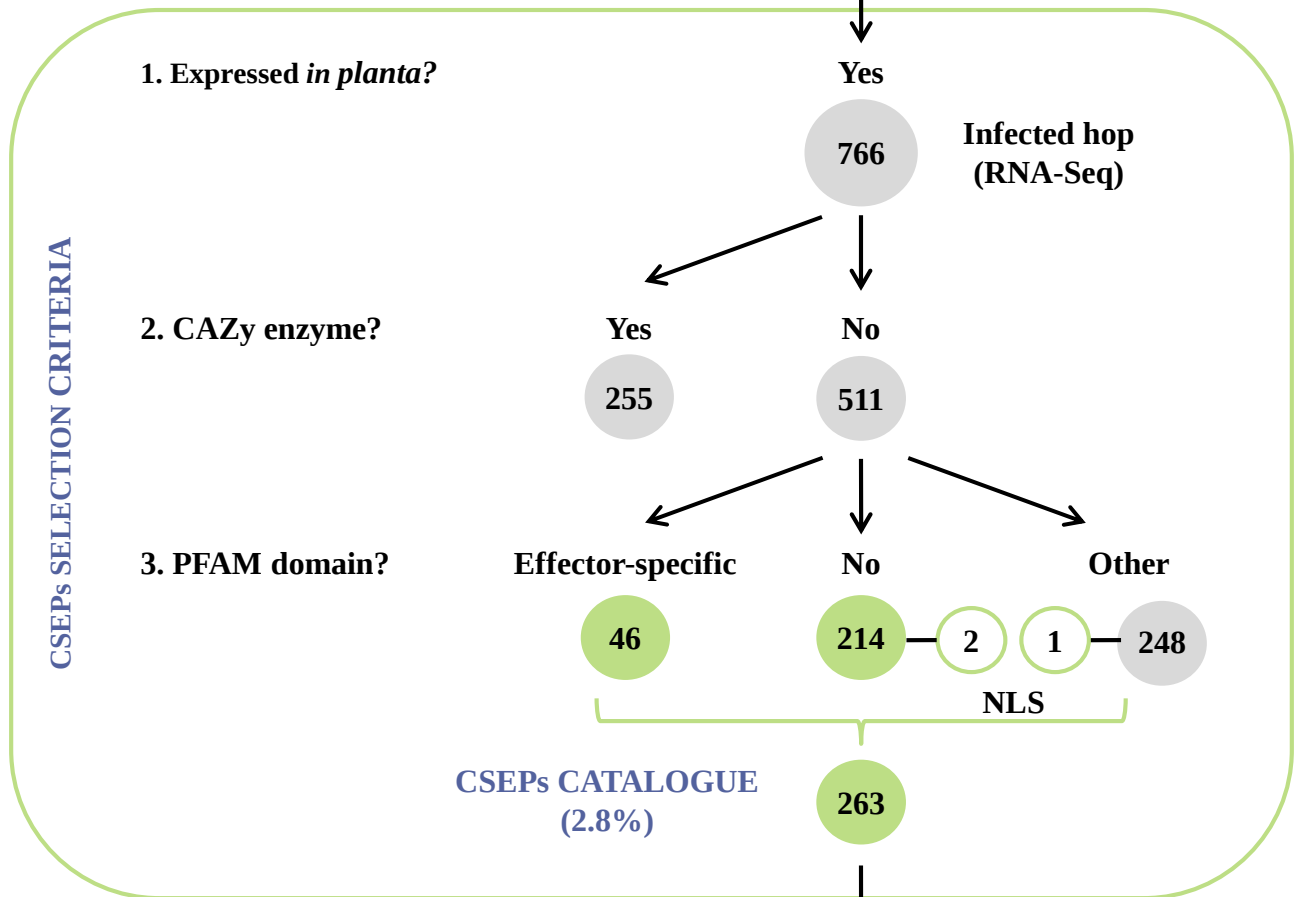
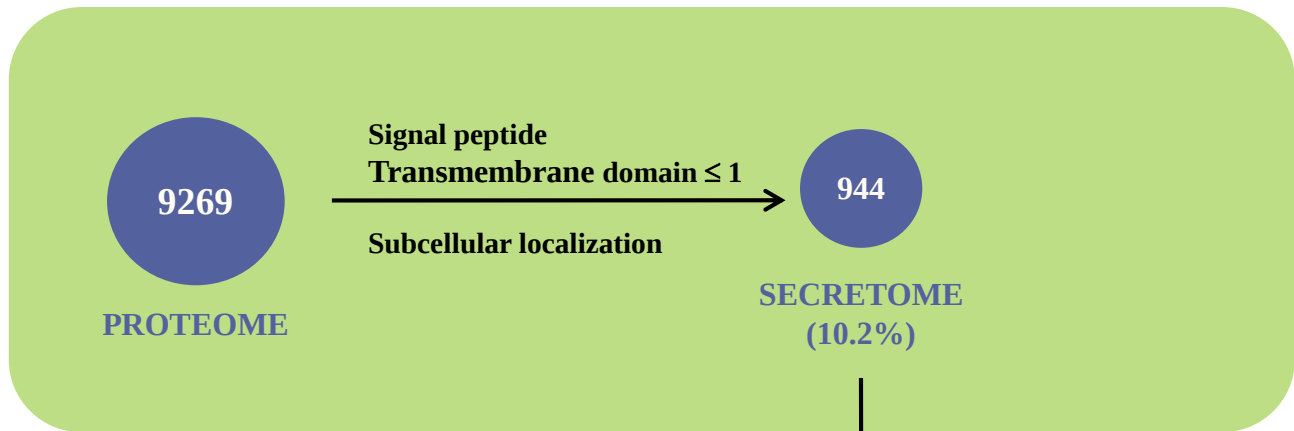
987 Table S8. List of primers used in this study. ^a, oligonucleotide designed according to template [123];
 988 ^b, oligonucleotide that amplifies the promoter region of the target gene; ^c, oligonucleotide that
 989 amplifies the terminator region of the target gene; ^d, oligonucleotide that amplifies the target gene for
 990 knockout (KO) selection; ^e, oligonucleotide that amplifies genomic and vector sequences for KO
 991 selection; ^f, *V. nonalfalfae* lethal pathotype-specific marker [97].

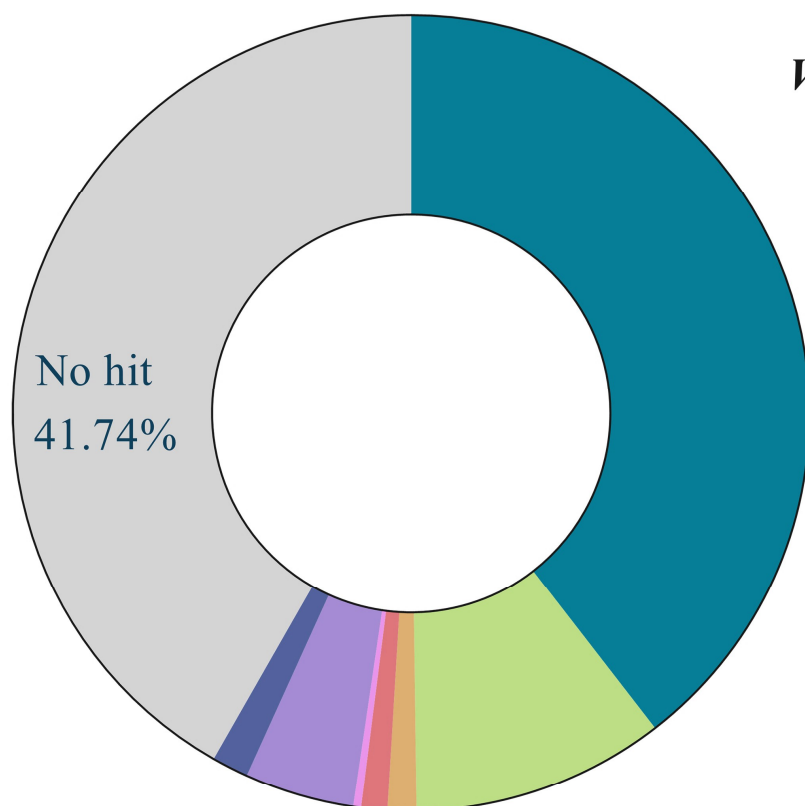
992 Additional file 2. A figure in PDF format. Enrichment of carbohydrate metabolic processes and
 993 peptidase activity in the predicted *V. nonalfalfae* secretome compared to proteome. GO terms for
 994 biological process (A) and molecular function (B) are presented at GO level 4. Relative abundance of
 995 gene ontology (GO) terms determined by Blast2GO is expressed in percentages of predicted fungal
 996 secretome and proteome, respectively.

997 Additional file 3. A figure in PDF format. A heatmap displaying the expression patterns of *V.*
 998 *nonalfalfae* genes during infection of susceptible 'Celeia' and resistant hop 'Wye Target'. Fungal
 999 transcripts were first identified by mapping of reads with at least 90% sequence identity and 90%
 1000 sequence coverage to the *V. nonalfalfae* reference genome [22] using CLC Workbench.
 1001 Normalization by trimmed mean of M values (TMM) [129] was performed to eliminate composition
 1002 biases between libraries. Read counts were converted into log₂-counts-per-million (logCPM) values
 1003 and a cutoff of CPM >1 was chosen. Color scale bar represents the logCPM values, with darker red
 1004 color meaning higher expression values.

1005 Additional file 4. A figure in PDF format. Some *V. nonalfalfae* CSEPs deletion mutants exhibit
 1006 unaffected pathogenicity. Relative area under the disease progress curve (rAUDPC) is presented for
 1007 the three *V. nonalfalfae* deletion mutants (each in four replicates) and the wild type fungus. Depicted

1008 are mean values $\pm$ SEM (n=10). Analysis of variance was first performed by Levene's test, followed
 1009 by Dunnett's test to compare each treatment (knockout) with a single control (wild type); however,
 1010 no statistical differences were found at α level of 5%.





V. nonalfalfae secretome: 944 proteins

Metabolism (39.51%)

Polysaccharide metabolism and transport	91
Carbohydrate metabolism and transport	62
Redox	38
Secondary metabolism	27
Transferases	8
Coenzyme metabolism and transport	5
Cell envelope metabolism and transport	4
Nitrogen metabolism and transport	4
Amino acids metabolism and transport	1
Nucleotide metabolism and transport	2
E- transfer	1
Other enzymes	131

Processes IC (10.27%)

Proteases	72
Ion metabolism and transport	17
Phospholipid metabolism and transport	7
Transport	1

Processes EC (1.17%)

Cell adhesion	9
Toxins/defense	1
Blood clotting	1

Information (1.06%)

DNA replication/repair	9
RNA processing	1

Regulation (0.32%)

DNA-binding	2
Signal transduction	1

General (4.45%)

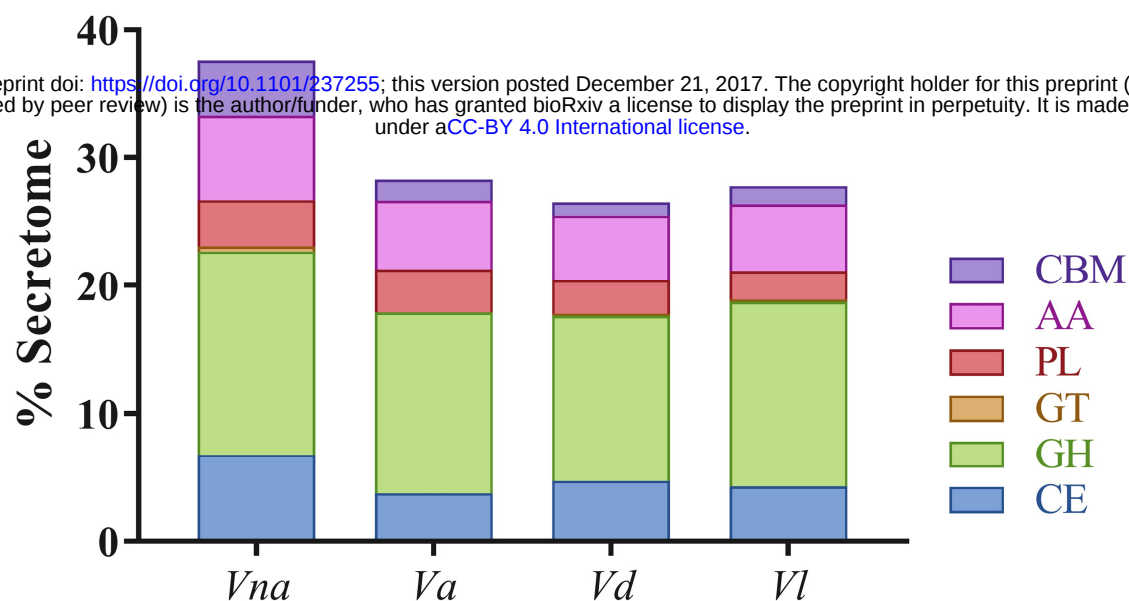
Small molecule binding	34
Protein interaction	3
General	3
Ligand binding	2

Other (1.48%)

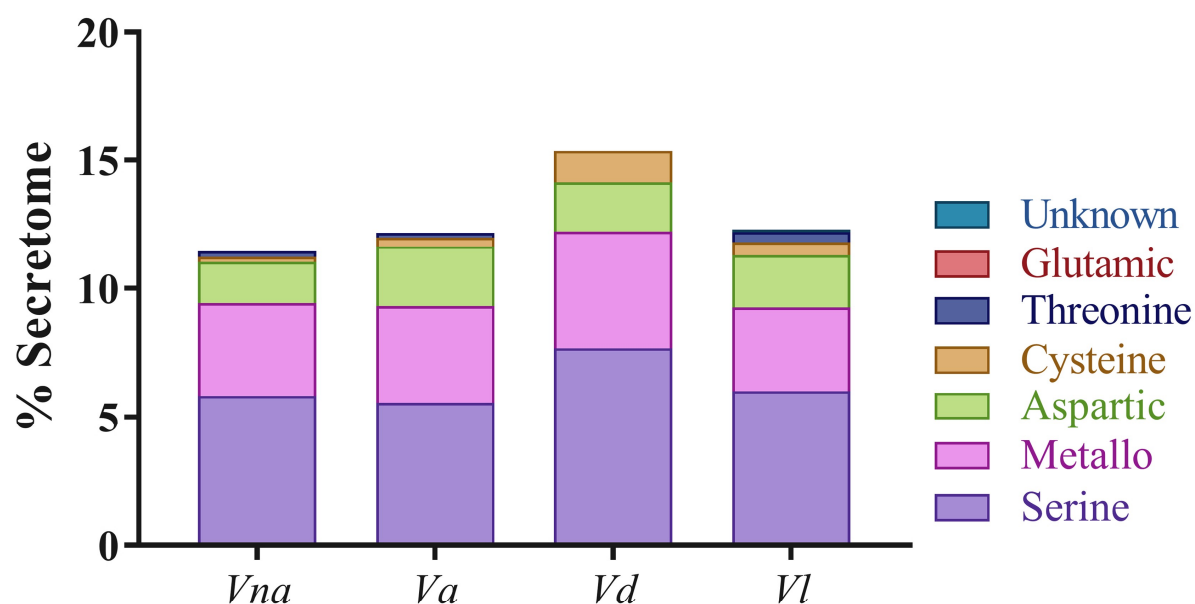
Unknown function	11
Viral proteins	2

(A)

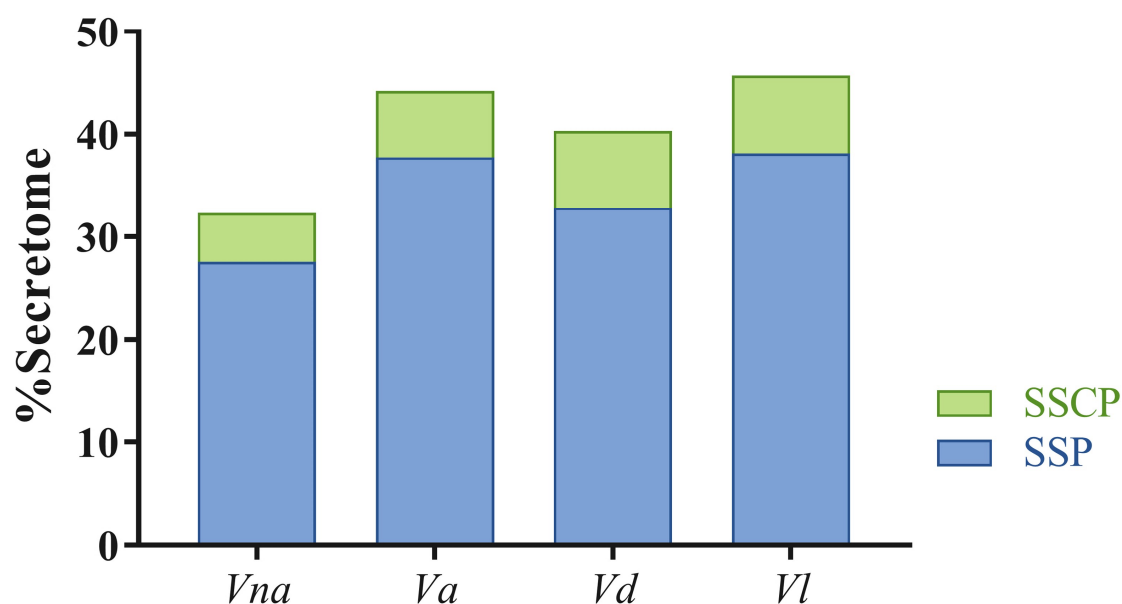
bioRxiv preprint doi: <https://doi.org/10.1101/237255>; this version posted December 21, 2017. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.

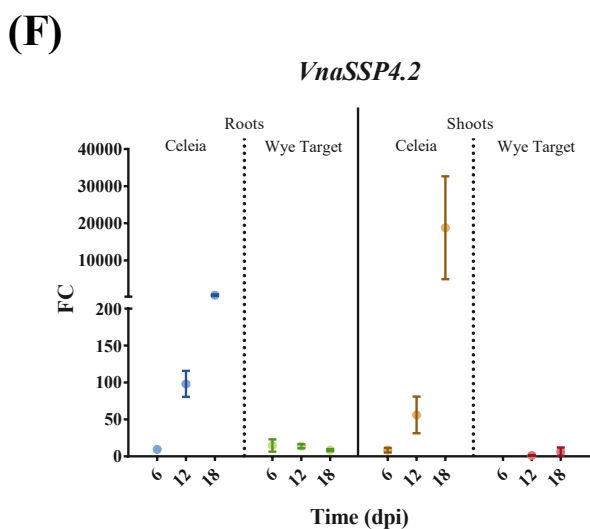
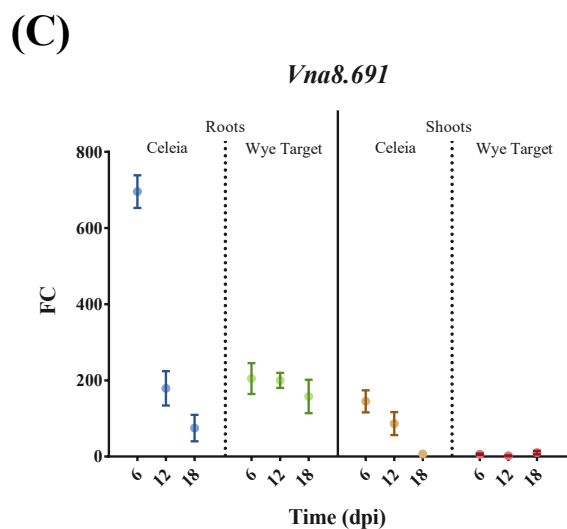
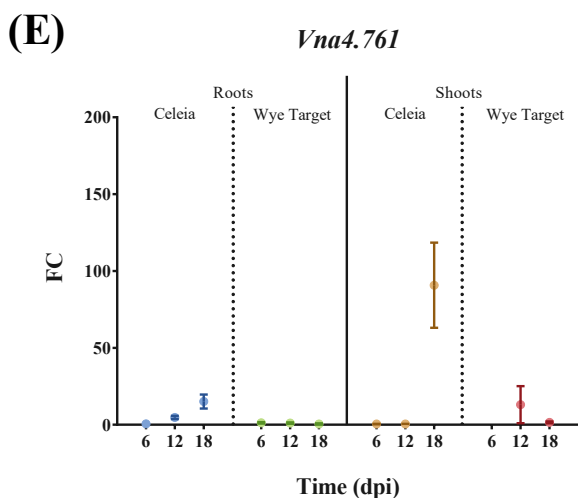
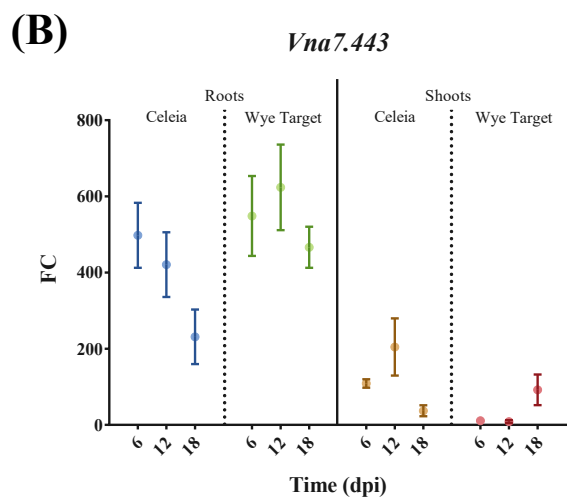
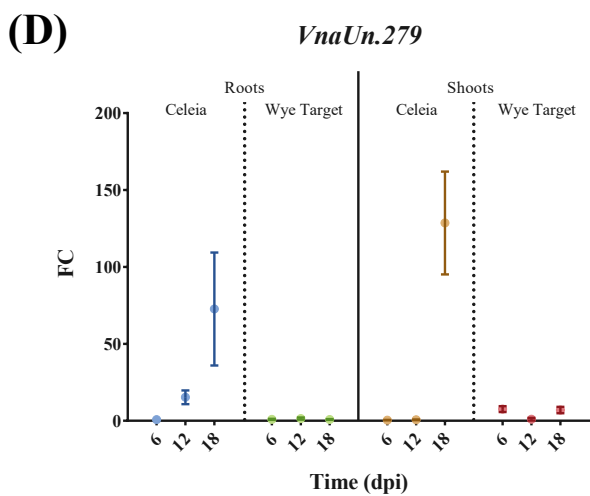
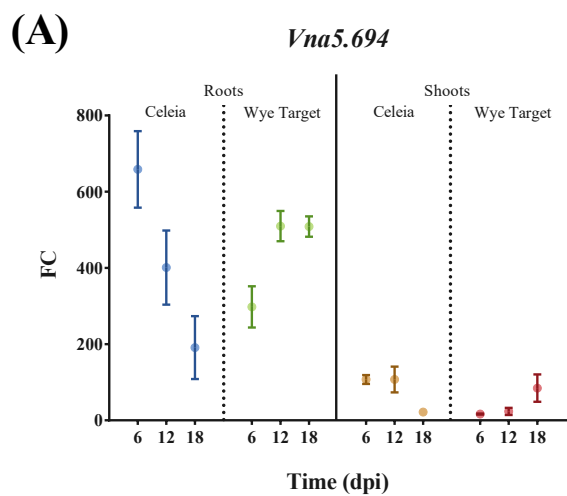


(B)



(C)





A



B



C

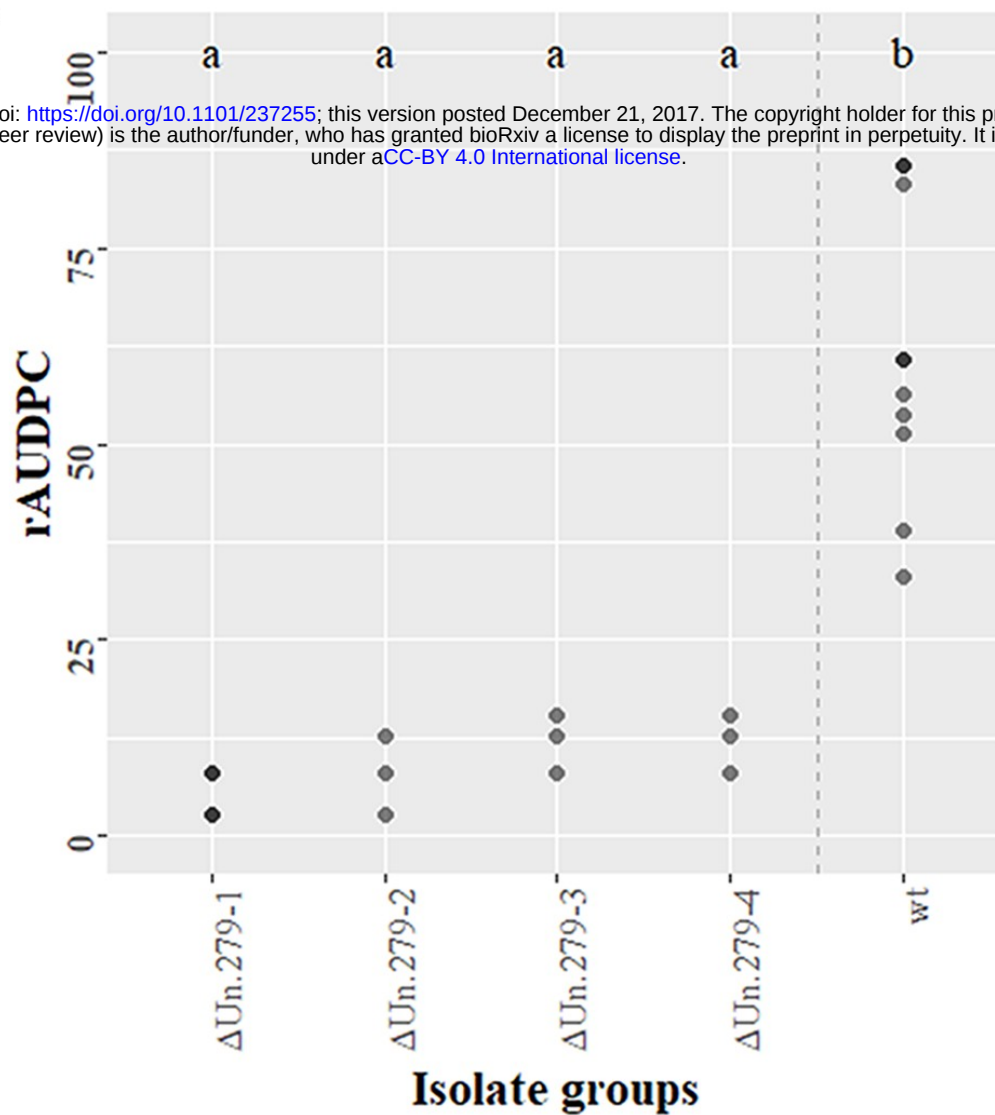


D



A

bioRxiv preprint doi: <https://doi.org/10.1101/237255>; this version posted December 21, 2017. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.



B

