1	Title: Effector prediction and characterization in the oomycete pathogen
2	Bremia lactucae reveal host-recognized WY domain proteins that lack the
3	canonical RXLR motif
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5	Short title: Effector prediction and characterization in the oomycete pathogen
6	Bremia lactucae
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24 Abstract

25 Pathogens infecting plants and animals use a diverse arsenal of effector proteins to 26 suppress the host immune system and promote infection. Identification of effectors 27 in pathogen genomes is foundational to understanding mechanisms of pathogenesis, 28 for monitoring field pathogen populations, and for breeding disease resistance. We 29 identified candidate effectors from the lettuce downy mildew pathogen, Bremia 30 *lactucae*, using comparative genomics and bioinformatics to search for the WY 31 domain. This conserved structural element is found in *Phytophthora* effectors and 32 some other oomycete pathogens; it has been implicated in the immune-suppressing 33 function of these effectors as well as their recognition by host resistance proteins. We 34 predicted 54 WY domain containing proteins in isolate SF5 of *B. lactucae* that have 35 substantial variation in both sequence and domain architecture. These candidate 36 effectors exhibit several characteristics of pathogen effectors, including an N-37 terminal signal peptide, lineage specificity, and expression during infection. 38 Unexpectedly, only a minority of *B. lactucae* WY effectors contain the canonical N-39 terminal RXLR motif, which is a conserved feature in the majority of cytoplasmic 40 effectors reported in *Phytophthora* spp. Functional analysis effectors containing WY 41 domains revealed eleven out of 21 that triggered necrosis, which is characteristic of 42 the immune response on wild accessions and domesticated lettuce lines containing 43 resistance genes. Only two of the eleven recognized effectors contained a canonical 44 RXLR motif, suggesting that there has been an evolutionary divergence in sequence 45 motifs between genera; this has major consequences for robust effector prediction in 46 oomycete pathogens.

47 Author Summary

There is a microscopic battle that takes place at the molecular level during infection 48 49 of plants and animals by pathogens. Some of the weapons that pathogens battle with 50 are known as "effectors," which are secreted proteins that enter host cells to alter 51 physiology and suppress the immune system. Effectors can also be a liability for plant 52 pathogens because plants have evolved ways to recognize these effectors, triggering 53 a defense response leading to localized cell death, which prevents the spread of the 54 pathogen. Here we used computer models to predict effectors from the genome 55 of Bremia lactucae, the causal agent of lettuce downy mildew. Three effectors were 56 demonstrated to suppress the basal immune system of lettuce. Eleven effectors were 57 recognized by one or more resistant lines of lettuce. In addition to contributing to our 58 understanding of the mechanisms of pathogenesis, this study of effectors is useful 59 for breeding disease resistant lettuce, decreasing agricultural reliance on fungicides.

60

61 Introduction

62 The phylum Oomycota includes some of the most devastating pathogens of 63 both plants and animals [1]. Although oomycetes resemble fungi in their filamentous 64 growth and infection structures, they are more closely related to brown algae than to 65 fungi [2]. Notable oomycetes include the plant pathogens causing late blight of potato 66 (*Phytophthora infestans*) [3], sudden oak death (*Phytophthora ramorum*) [4], root rot 67 (*Phytophthora* [5] and *Pythium* spp. [6,7]), white blister rust of *Brassica* spp. (*Albugo* 68 spp.) [8], and downy mildews (e.g. *Bremia, Peronospora, Plasmopara* spp.) [9], as well 69 as several important animal pathogens infecting fish (Saprolegnia spp.), shellfish 70 (Aphanomyces astaci), and mammals, including humans (Pythium insidiosum) [1,10]. 71 Many types of plant and animal pathogens, including the oomycetes, secrete proteins 72 known as effectors to promote virulence by manipulating the physiology of the host 73 cells and by suppressing the host immune system [11]. In plants, effectors are also 74 determinants of resistance or susceptibility through their direct or indirect 75 interactions with cognate nucleotide-binding leucine rich repeat proteins (NLRs) 76 encoded by resistance genes [12]. Effectors are secreted from the pathogen and may 77 act extracellularly or they may be translocated into the cytoplasm [11]. One class of 78 translocated effectors from plant pathogenic oomycetes of the class Peronosporales, 79 which includes *Phytophthora* and the downy mildews, are the RXLR effectors, so 80 called for their N-terminal motif usually consisting of arginine, followed by any amino 81 acid, then followed by leucine and arginine. RXLR effectors also contain an N-terminal 82 signal peptide, which designates them for extracellular transport by way of the 83 endoplasmic reticulum and Golgi apparatus [13]. The RXLR motif is often associated 84 with a downstream EER motif, both of which have been associated with secretion 85 and/or translocation of effectors into the plant cell [13.14]. For some RXLR effectors. 86 such as Avr3a, the RXLR motif has been shown to be cleaved just prior to the EER 87 sequence and therefore plays a role in secretion rather than uptake into the host cell 88 [15]. The RXLR motif is similar in sequence to the PEXEL motif (RXLX[ED0]) of the 89 distantly related malaria pathogen (*Plasmodium falciparum*) [16] and the TEXEL 90 motif of *Toxoplasma gondii* (RRLXX) [17], both of which are required for proteolytic 91 modification in the endoplasmic reticulum and destine effector proteins for 92 specialized export out of the cell [18].

93 The downy mildews and the related *Phytophthora* species have different 94 lifestyles (obligate biotrophy vs. facultative hemibiotrophy) [19]; however, their 95 effectors share similar features. Many effectors in the Peronosporales have RXLR and 96 EER motifs; although several alternative sequences to RXLR have been found in 97 downy mildews, including RVRN (ATR5 from Hyaloperonospora arabidopsidis) [20], 98 QXLR (*Pseudoperonospora cubensis*) [21], GKLR (*Bremia lactucae*) [22,23], and RXLK 99 (*Plasmopara halstedii*) [24]. The C-terminal effector domains of RXLR effectors from 100 *Phytophthora* and downy mildews also share some common sequence motifs and 101 structural features, such as the 24 to 30 amino acid W, Y, and L motifs, which were 102 first identified bioinformatically [25]. Structural analysis on four different RXLR 103 effectors from *Phytophthora infestans* (Avr3a and PexRD2), *P. capsici* (Avr3a11), and 104 the downy mildew pathogen *H. arabidopsidis* (ATR1) revealed that the W and Y motifs 105 form an alpha-helical fold that may play a role in protein-protein interactions [26– 106 29]. This effector-associated fold, termed the WY domain after its conserved 107 tryptophan and tyrosine residues, is structurally highly conserved between effectors 108 from multiple Peronosporales species, despite sharing less than 20% sequence 109 similarity across the whole domain [26][30]. The WY domain appears to be specific 110 to the Peronosporales and was predicted to be present in nearly half of the RXLR 111 effectors of *P. infestans* and a fourth of the RXLR effectors in *H. arabidopsidis* [26].

Functional studies of WY domain containing proteins have indicated that certain residues in the WY domain are essential for the immune suppressing functions of *P. sojae* Avr1b [31], *P. infestans* Avr3a [32], and *P. infestans* PexRD2 [33]. Furthermore, mutation of two conserved leucines in the WY domain of PexRD2

116 disrupted interaction with its target, MAPKKKE, consistent with this domain being 117 important for protein-protein interactions [33]. WY domain containing proteins 118 appear to interact with a variety of host targets, with Avr3a from *P. infestans* targeting 119 the E3 ligase CMPG1 [32], P. infestans PexRD54 targeting potato autophagy-related 120 protein ATG8 [34], PsAvh240 from *P. sojae* targeting an aspartic protease [35], and 121 PSR2 from *P. sojae* and *P. infestans* suppressing host RNA silencing through 122 interactions with dsRNA-binding protein DRB4 [36–38]. Mutation analysis of the 123 regions encoding the seven individual WY domains of PSR2 demonstrated differential 124 contributions of each domain to virulence of *P. sojae*, suggesting that the WY domain 125 may act as a module during effector evolution [30]. In addition to its roles in immune 126 suppression, the WY domain has been shown to be important for immune recognition 127 of the effector by nucleotide binding-leucine rich repeat (NB-LRR) resistance proteins 128 [39].

129 Effector annotation in oomycete genomes has often relied on sequence 130 similarity to known effectors or on prediction of conserved motifs, such as the RXLR 131 motif, or in the case of Crinklers, the LXLFLAK motif [3]. Due to the short length and 132 degeneracy of the RXLR sequence, the motif occurs frequently by chance; therefore, 133 there is a high false positive rate (>50%) using string-based searches [25]. HMM-134 based searches have much lower false positive rates, but the false negative rate may 135 be higher if the genome of interest has diverged significantly from the species used to 136 build the HMM. Downy mildews have a narrow host range and pathogenicity-related 137 proteins are likely to show high lineage-specificity due to co-evolution with their 138 hosts. There is already evidence that downy mildew effectors show divergence from

the canonical RXLR motif [20,23,24], thus complementary approaches for effector prediction that utilize other conserved features, such as the WY domain, are necessary to fully characterize the repertoire. In addition, due to the importance of the WY domain in effector function [31–33], WY domain containing proteins may be better candidates for effectors than those containing only the RXLR motif.

144 To identify a more complete repertoire of candidate effectors in the reference 145 genome of *B. lactucae* and to test whether the WY domain is informative for 146 predicting effectors in the Peronosporales, we searched for this domain using an 147 HMM built from sequences of the WY domain in three *Phytophthora* species [27]. This 148 revealed additional effector candidates that had not been found using an RXLR-based 149 search; similar results were also obtained for other downy mildews and well-studied 150 *Phytophthora* species. These predicted WY proteins from *B. lactucae* had other 151 signatures of oomycete effectors, such as presence of a secretion signal, N-terminal 152 intrinsic disorder, lineage specificity, and expression during infection. A subset of the 153 predicted WY effectors suppressed the host immune system, while others elicited 154 programmed cell death in specific genotypes of lettuce, indicative of recognition by 155 host resistance proteins. Therefore, searches for the WY domains are highly useful for 156 identification of downy mildew effectors lacking the RXLR motif, which was 157 previously considered canonical for effectors of the Peronosporales.

158

159 **Results**

160 Our HMM search initially identified 59 candidate WY effectors encoded by the
161 gene models predicted in the *B. lactucae* SF5 assembly [40]; however, three pairs of

162 genes appeared to be allelic based on sequence similarity and read depth, leaving a 163 total of 55 non-redundant genes encoding candidate effectors (Figure 1A; 164 Supplemental Table 1). Signal peptides were predicted for 43 of these 55 proteins, of 165 which two proteins had a predicted transmembrane helix outside of the signal 166 peptide (Figure 1A). Several predicted WY proteins seemed to be missing their start 167 codons due to N-terminal truncation when compared to close relatives. One of these 168 predicted proteins, BLN06, was found to be missing a significant portion of the N-169 terminal sequence in SF5 compared to BL24, which was the source isolate for the 170 cloning of *BLN06* reported in Pelgrom *et al.* [41].

171 The N-terminal sequences of the 55 predicted WY effectors were examined for 172 the RXLR motif using both HMMs and string searches. One protein was predicted to 173 have an RXLR motif by the HMM and an additional 10 proteins were identified by a 174 string search for [RQGH]xLR or RXL[QKG], while 33 proteins were predicted to have 175 an EER motif using a string search for [DE][DE][EK] (Figure 1A). To find divergent 176 RXLR-like motifs in the WY effector candidates, we searched for a highly degenerate 177 pattern based on mutational studies of the RXLR motif [35] and natural RXLR variants 178 reported for other downy mildews [5–9]. This revealed an additional 38 proteins with 179 an RXLR-like motif within the first 60 amino acids after the signal peptide matching 180 the pattern [RKHG0][X]{0,1}[LMYFIV][RNK], many of which also had an EER motif 181 (Figure 1A) (Supplemental Table 1). Therefore, the majority of candidate WY 182 effectors in *B. lactucae* have a non-canonical RXLR motif. In the C-terminal domain, 183 the WY effector candidates had one to seven WY domains per protein, with a diversity 184 of domain architectures (Figure 1). The WY domain from *B. lactucae* showed high 185 conservation of the characteristic conserved tryptophan (W) residue but appeared to 186 show equal preference for tyrosine (Y) or phenylalanine (F) for the second 187 characteristic residue of the domain (Fig. 1B). To investigate whether or not there are 188 WY effector candidates that are lacking the canonical RXLR motif in other oomycetes, 189 the predicted open reading frames (ORFs) from several published oomycete genomes 190 were surveyed for the RXLR motif and the WY domain. Approximately half of the WY 191 proteins predicted in seven downy mildew species lacked the RXLR motif; in six 192 *Phytophthora* spp., the majority of predicted WY proteins had RXLR motifs, but 193 between 9–21% of secreted WY proteins did not contain this motif (Fig. 2). 194 Consequently, the repertoire of candidate WY effectors in other oomycetes may be 195 heavily under-reported.

196 Intrinsic disorder had previously been reported to be a characteristic of the N-197 terminus of oomycete effectors containing the RXLR motif [42]. Therefore, we 198 investigated whether the predicted degree of structural disorder was a characteristic 199 of candidate WY effectors lacking a canonical RXLR. The predicted levels of intrinsic 200 disorder were calculated for proteins containing RXLR motifs, for proteins containing 201 WY domains but no RXLR motif, as well as for the entire predicted secretome for 202 comparison. Proteins containing RXLR and/or WY domains had higher levels of 203 intrinsic disorder at their N-termini after the highly ordered signal peptide than the 204 entire set of secreted proteins (Fig. 3). Proteins containing a WY domain but lacking 205 a canonical RXLR motif had on average more disordered N-termini than effectors that 206 had RXLR but not a WY domain. The WY domain containing region had higher levels 207 of predicted structure than the RXLR-containing proteins that lacked a WY domain and the secreted proteins as a whole (Fig. 3), consistent with the WY domains forming
an α-helix bundled structure [27]. This predicted pattern of high intrinsic disorder at
the N-terminus and high structure towards the C-terminus was consistently observed
in all six downy mildews and six *Phytophthora* species analyzed (Suppl. Fig. 2).
Therefore, a high level of intrinsic disorder is a consistent characteristic of the Ntermini of oomycete effectors, regardless of whether they have a canonical RXLR
motif; the functional significance of this remains to be investigated.

To evaluate lineage specificity of effectors due to co-evolution of pathogens with their hosts, we used BLAST to identify orthologs in other oomycete species. All of the 39 predicted secreted candidate WY effectors of *B. lactucae* had little sequence similarity to sequences in other genomes with the best BLASTP hit being only 46% identity with a protein from *P. infestans* (Fig. 4). Most of the proteins had best-hit identities between 20 to 30%, which is similar to the level of amino acid conservation between WY domains in different effectors [27].

A time-course RNA-seq experiment of lettuce seedlings infected with *B. lactucae* isolate SF5 was analyzed to investigate the expression of the WY effector candidates during infection. Expression was detected for all candidate WY encoding genes. The four mostly highly expressed WY containing effectors lacked a canonical RXLR motif (Fig. 5).

To test whether *B. lactucae* WY effectors were recognized by the host immune system, 21 randomly-selected WY effectors that differed in their number of WY domains were chosen to be screened against lettuce germplasm. Genes were cloned from amplified genomic DNA to attempt to capture both alleles of each effector from

the heterozygous isolate SF5 [40]. In order to capture additional allelic diversity,
some effectors were also cloned from the heterokaryotic isolate C82P24 [40]. Two
alleles were obtained for many effectors; in addition, chimeric sequences appeared to
be obtained for several genes. Clones of all unique sequences for each effector were
retained because they could be informative for dissecting the sequence basis of host
recognition. This resulted in multiple distinct clones (wildtype alleles plus chimeric
sequences) of some effectors.

238 We screened 215 different accessions of wild and cultivated lettuce (Suppl. 239 File 2) using Agrobacterium-mediated transient expression for the elicitation of cell 240 death for their reactions to clones representing the 21 WY candidate effectors. These 241 accessions collectively express the majority of the known *Dm* genes as well as new 242 resistance factors [36]. Eleven of the 21 WY effectors were recognized by one or more 243 accessions (Fig. 6). Alleles of the same effector showed similar reactions except for 244 three genes (Fig. 6). The truncated version of BLN06 cloned from SF5 did not trigger 245 cell death in LS102, NunDm17, or RYZ2146 (Suppl. Figure 3) in contrast to BLN06 246 cloned from BL24 [41], suggesting that the recognition of this effector by these 247 genotypes may be determined by the N-terminal region of the effector after the 248 secretion signal.

The lines ViAE and ViCQ, which have introgressions from the wild lettuce species *L. virosa* [43], recognized five effectors: BLN08, BSW03, BSW04m, BSW04p, and BSW14, as well as the chimeric sequence BSW04m/p (Fig. 6A). The *L. virosa* accessions that were the resistance donors for ViAE and ViCQ also recognized these five effectors, but not the chimeric sequence BSW04m/p (Fig. 6B). Many *L. sativa*

254 cultivars and a few genotypes of *L. saliana* and *L. serriola* were observed to have 255 necrosis or yellowing in response to the chimeric effector BSW04m/p, suggestive of 256 a non-specific reaction to this unnatural protein. Some of the genotypes that 257 recognized BSW04m/p also recognized BSW04p, but not the paralog BSW04m. Three 258 of these genotypes, Capitan, Ninja, and Femke, share the resistance gene Dm11; 259 therefore, we tested two additional cultivars, Fila and Mondian, which also contain 260 Dm11 for recognition of BSW04p. These varieties also recognized BSW04m/p and 261 BSW04p, but not BSW04m. Due to recognition of BSW04p by multiple varieties that 262 contain *Dm11*, BSW04p is a candidate for the protein encoded by the *Avr11* gene. 263 BLN08, BSW03, BSW04m, and BSW14 are candidate avirulence proteins for which 264 cognate *Dm* genes have vet to be identified.

265 Bioinformatic prediction of subcellular localization using NucPred [44] 266 suggested that BSW04p had a C-terminal nuclear localization signal (score 0.79). To 267 investigate whether BSW04p was nuclear localized, N-terminal yellow fluorescent 268 protein (YFP) fusions were expressed in lettuce using *Agrobacterium*-mediated 269 transient assays. We also made N-terminal YFP fusions of two other effectors. BLN08 270 and BSW03, which lacked predicted nuclear localization signals. BSW04p was 271 localized to the nucleus as predicted, while BLN08 and BSW03 were localized to the 272 cytoplasm and/or periplasm (Fig. 7). Therefore, predictions of subcellular 273 localization were accurate for these three effectors and may indicate the cellular 274 location of their targets during infection.

We also tested candidate effectors for their ability to suppress PAMP-triggered
immunity (PTI). Twenty-one effectors were transiently expressed in *Nicotiana*

benthamiana and the level of reactive oxygen species (ROS) production induced by
flg22 was measured. Three effectors significantly suppressed ROS production to a
similar extent as two known bacterial suppressors of PTI (Fig. 8). The level of
induction of ROS by flg22 was significantly higher with some effectors; however,
there was no induction of ROS in the absence of flg22. Therefore, in this assay, at least
three effectors suppress PTI and some may actually increase PTI responsiveness.

283

284 **Discussion**

285 Effectors play a critical role in interactions between pathogens and their hosts. 286 Accurate prediction and annotation of effector repertoires is foundational for 287 functional genomics studies of pathogens. Due to their economic importance in 288 agriculture, an increasing number of *Phytophthora* and downy mildew pathogens are 289 being sequenced. Prior to this study, the majority of cloned avirulence genes have 290 encoded proteins with a RXLR motif [45]. This was not surprising considering that 291 most of these studies used the RXLR motif as their primary, or only, criterion to 292 identify candidate effectors. Our results demonstrate that there are many candidate 293 effectors containing the effector-related WY domain that lack the canonical RXLR 294 motif, especially in the downy mildews, but also in *Phytophthora* spp. In *B. lactucae*, 295 we have shown that these non-RXLR, WY domain containing effectors show 296 characteristics of RXLR effectors, such as N-terminal intrinsic disorder, lineage 297 specificity, and expression during infection. Furthermore, some of these proteins can 298 act as suppressors of the host immune system, while others trigger the hypersensitive 299 response in resistant host cultivars. Thus, despite lacking the canonical RXLR-motif, these WY domain containing proteins have both virulence and avirulence activity.
Consequently, numerous candidate effector genes are likely to have been missed in
the genomic analysis of other species within the Peronosporales. Bioinformatics
pipelines for predicting effectors in both *Phytophthora* spp. and downy mildew
pathogens should include an HMM for the WY domain because it may be more
informative of function and result in fewer false positives than string searches for the
RXLR motif.

307 Many WY effectors were initially predicted to not have a signal peptide and 308 therefore not to be secreted; this could have been due to misannotation or reflect 309 biological reality. Several had a predicted signal peptide downstream of the ATG in 310 the gene model; these signal peptides started with a methionine, which was 311 supported as the correct start in manual curation using RNAseq data. Therefore, 312 studies that rely on the presence of a signal peptide when using predicted ORFs may 313 be missing true effectors with misannotated start codons. Some WY effectors had 314 clearly lost the signal peptide due to N-terminal deletion; this was observed by 315 comparisons within effector families. The truncated effectors are unlikely to be 316 functional because they are missing the secretion signal and would not therefore be 317 secreted from the cell. Signal peptide loss may be an evolutionary strategy for the 318 pathogen to evade recognition of effectors.

Although very few of the WY effectors from *B. lactucae* contained the canonical RXLR motif, nearly all of them contained a degenerate RXLR motif. However, the degenerate RXLR regular expression should not be used on its own to search for genes encoding RXLR proteins due to the high false positive rate (>50%). Functional studies

323 are needed to ascertain which of these degenerate motifs can function similarly to the 324 canonical RXLR motif in protein secretion [15]. Despite lacking the canonical RXLR 325 motif, the WY effectors had other features similar to RXLR proteins such as high N-326 terminal intrinsic disorder and the presence an EER motif. N-terminal intrinsic 327 disorder has been predicted for RXLR effectors of *Phytophthora* and is also a common 328 feature of bacterial effectors [46]. The biological significance of these features 329 remains to be studied, particularly whether these intrinsically disordered domains 330 are important in post-translational modification or protein-protein interactions as is 331 the case for intrinsically disordered regions in other organisms [47,48].

332 The number of WY domains per protein varied considerably, from one to 333 seven in *B. lactucae*. Many pathogen effectors exhibit rapid evolution and divergence 334 due to selective pressure of evolving host targets and host resistance proteins [25]. 335 Duplication of domains may allow for evolution of novel effector functions or for 336 evasion of host recognition while retaining function [30]. Variation in the number of 337 repeated domains is reminiscent of NB-LRR proteins, which recognize effectors 338 (and/or effector activity) and also have leucine-rich repeated domains [49,50]. 339 Duplication of regions encoding WY domains may have happened within a gene 340 through replication errors or between different genes through illegitimate 341 recombination. The genomic sequences of the repeats will be analyzed in multiple 342 isolates to reveal the origins of these duplications and domain expansion or 343 contraction.

Analysis of the amino acid sequences of the WY domain in *B. lactucae* showed
that it may be better considered as a W[Y/F] domain, due to the equal preference for

346 phenylalanine and tyrosine for the second characteristic residue. This substitution 347 has a BLOSUM score of 3, indicating that it is fairly common. Both tyrosine and 348 phenylalanine are aromatic amino acids; however, the hydroxyl group on tyrosine 349 makes it slightly bulkier, more polar, and introduces a potential phosphorylation site. 350 Variation in this region is not uncommon in other oomycete species: the fifth of the 351 seven domains in Psr2 of *P. sojae* has an F instead of a Y and is similar in structure to 352 the single domain of ATR1 in *H. arabidopsidis* that has a cysteine at the Y position [30]. 353 Further structural characterization is needed to reveal whether these substitutions 354 alter protein structure and their biological function.

355 In oomycetes, WY domain containing effectors have been shown to have 356 several functions including PTI suppression [51]. At least three WY effectors from B. 357 *lactucae* were able to suppress the host immune system by interfering with pathogen-358 triggered production of ROS. Suppression of host defenses is critical to the survival of 359 *B. lactucae* and therefore it is not surprising that multiple effectors target the basal 360 immune system. Identification of the host targets of these effectors will determine 361 which steps in the signal transduction cascade are modulated by each effector or may 362 reveal candidates for susceptibility genes in the host that are required for successful 363 proliferation of *B. lactucae*.

Effectors are powerful tools for the discovery and characterization of host resistance genes [52]. Eleven of the *B. lactucae* effectors tested were recognized by one or more lettuce lines. Their cognate R genes will be identified using mapping of segregating F_{2:3} and recombinant inbred line populations. Recognition of four *B. lactucae* effectors (BLG01, BLN08, BLR31, and BLR38) has been successfully mapped

369 [23,41,53]. BLG01 and BLN08 have been shown to be recognized broadly by *L. saligna*370 [23,53]. Our study confirms the results for BLN08 and revealed BSW14 as an
additional effector recognized by *L. saligna*. BLN08 and BSW14 share little sequence
372 similarity (besides containing WY domains). Non-host resistance in *L. saligna* is
373 clearly complex [54], but these results show that it is mediated in part by recognition
374 of multiple effectors.

375 Not all RXLR candidate effectors have WY domains and the presence of a WY 376 domain is not required for the avirulence activity of all effectors; for example, the *B*. 377 lactucae effectors BLG01 and BLG03 do not contain WY domains, yet they elicit an 378 immune response in lettuce [23]. Structural elucidation of an RXLR effector lacking 379 the WY motif, H. arabidopsidis ATR13, revealed that it contained a helical fold that 380 was distinct from the WY fold [55]. It would be informative to determine and compare 381 the protein structures of additional RXLR effectors to determine whether there are 382 other conserved C-terminal domains that may be involved in effector function in a 383 similar way as the WY domain.

384 Both RXLR and WY effectors provide tools for monitoring pathogen 385 populations and effector-driven resistance breeding. Analysis of diverse, global 386 isolates will allow the characterization of individual effector repertoires as well as the 387 development of the pan-repertoire for a whole pathogen species. Effectors also will 388 be highly instrumental in cloning their cognate resistance genes as well as the 389 identification of effector targets in the host. In addition, screens for resistance using 390 transient expression of individual effectors will allow the pyramiding of resistance 391 genes with different specificities that will maximize the evolutionary hurdle for the

392 pathogen to become virulent. Ultimately, knowledge of effector repertoires will allow
393 data-driven deployment of resistance genes leading to more durable disease
394 resistance [56].

395

396 Materials and Methods

397

398 Effector prediction

399 To search for the WY domain, a Hidden Markov Model (HMM) was built using 400 HMMer v3.1 [57] based on the inferred amino acid sequences of WY domains from *P*. 401 infestans, P. sojae, and P. ramorum. These sequences were obtained from the 402 supplemental material of Boutemy et al. [27] who predicted genes encoding WY 403 domains based on motif searches of candidate RXLR effectors and identified a 49 404 amino acid long motif that spanned the WY domain in the crystal structures of 405 Avr3a11 and PexRD2 [26]. Candidate WY effectors were predicted using this HMM to 406 search translated predicted ORF sequences (>80 amino acids) as well as gene models 407 in the draft genome of *B. lactucae* isolate SF5 [40]. Sequences with a positive HMM bit 408 score were considered to be putative WY domain effectors as in [27]. Signal peptide 409 prediction was performed on candidate WY effectors using SignalP v 4.1 [58] and 410 PhobiusSP [59]. Default settings were altered for SignalP v 4.1 to have sensitivity 411 similar to SignalP v 3.0. Output was compared between SignalP v 4.1 sensitive and the 412 combination of SignalP4.0 + SignalP 3.0, and identical results were obtained from the 413 two methods. Gene models were more accurate for predicting signal peptides than 414 translated ORFs — in part due to misannotated start codons upstream of the probable

415 true start codon and signal peptides in the ORFs. However, on several occasions the 416 gene model was missing a signal peptide found in the ORF – these gene models were 417 manually updated to reflect this likely true start site. SignalP v 4.1 in sensitive mode 418 was better able to predict signal peptides in proteins with misannotated start codons 419 compared to SignalP v 4.0 or v 5.0.

420

421 RXLR prediction

422 A combination of string searches and HMMs were used to search for the RXLR motif 423 in oomycete predicted WY proteins. The following strings were used based on 424 variants observed in downy mildews: [RQGH]xLR or RXL[QKG] for RXLR and 425 [DE][DE][KR] for EER. The Whisson et al. HMM [14] was also tested, although this did 426 not reveal any more RXLR effector candidates. To search for additional non-canonical 427 RXLR motifs in WY candidates, а highly degenerate string of 428 [RKHGQ][X]{0,1}[LMYFIV][RNK] was used.

429

430 Estimation of false positive rates for effector prediction

We determined false positive rates for each motif by analyzing multiple permutations of the non-redundant secretome using the same pipeline as described above for effector prediction. At least ten random permutations of the sequence space were created using the MEME fasta-shuffle-letters program (with a kmer size of 1) using peptides starting after the cleavage site identified by SignalP v4.1. The false positive rate for each motif was estimated as the average frequency of detection in the permutated sequences divided by the observed frequency in the original sequences.

- 438 The estimated false positive rate for the RXLR and WY motif searches are given in
- 439 Table 1.
- 440

441 Table 1. False positive rates for RXLR and WY HMM and string searches across

442 **14 oomycete genomes**

		False Positive	
		Rate (Average	
Motif	String/HMM	+SD)	Range
RXLR	RXLR	$45.2\%\pm3.4\%$	32-54%
RXLR	[RQGH]XLR or RXL[QKG]	56%±2%	50-64%
RXLR	Whisson HMM [14]	$1.5\%\pm0.8\%$	0.1-3.3%
EER	[DE][DE][KR]	$49.4\%\pm2.6\%$	41-62%
RXLR-EER	RXLR + [DE][DE][KR]	$9.4\%\pm3.0\%$	4-16%
Degenerate RXLR	[RKHGQ][X]{0,1}[LMYFIV][RNK]	$74\%{\pm}0.8\%$	71-77%
WY	Boutemy HMM [27]	$0.05\% \pm 0.09\%$	0-0.3%

443

444 Intrinsic Disorder

445 PONDR VSL2 [60] was used to calculate levels of intrinsic disorder for groups 446 of candidate effectors, grouped by the presence of an RXLR-like motif, EER-like motif, 447 and/or WY domain. PONDR VSL2 is a meta-predictor of protein disorder that utilizes 448 neural networks and amino acid context to give a weighted disorder score at each 449 amino acid position. The average sequence disorder was calculated at each amino 450 acid position in each group of effectors and aligned on the first amino acid. Plots were 451 generated from the average positional disorder scores calculated from all peptides in 452 a group. The entire non-redundant, predicted secretome was used as a reference for 453 comparison.

455 Sequence comparison and determination of lineage specificity

To generate a neighbor-joining tree of the WY effector candidates, whole 456 457 protein amino acid alignments were performed using MUSCLE 3.8.425 [61] 458 implemented in Geneious 11.0.5 (http://www.geneious.com) with default settings. A 459 UPGMA tree was built in Geneious with bootstrap resampling (100 replicates). The 460 resulting tree was annotated using the Interactive Tree of Life [62] and stylistically 461 refined in the GNU Image Manipulation Program (http://www.gimp.org). To build a 462 sequence logo for the WY domain from *B. lactucae*, a multiple sequence alignment of 463 WY domains was built using MUSCLE 3.8.425 [61], was manually corrected for 464 alignment errors. and sequence logo generated the using Weblogo 465 (http://weblogo.berkeley.edu/logo.cgi).

466 To determine if the candidate WY proteins are unique to *B. lactucae*, BLASTp 467 [63] was used to search for orthologs in other oomycete species. BLASTp-based 468 sequence comparisons with an e-value threshold of 0.01 were performed against the 469 following oomycete genomes: Albugo laibachii [8], H. arabidopsidis [64], P. capsici 470 [65], P. infestans [3], Phytophthora parasitica [66], P. ramorum [5], P. sojae [5], 471 Pythium ultimum [7], Saprolegnia parasitica [67], Pseudoperonospora cubensis [21], 472 Plasmopara halstedii [24], Plasmopara viticola [L-7-2 [68], INRA-PV221 [69], 473 Peronospora tabacina [70], and Peronospora effusa [71].

474

475 RNA-seq analysis

476 Messenger RNA was isolated from cotyledons of lettuce cv. Cobham Green
477 infected with isolate SF5 of *B. lactucae* with Dynabeads[™] mRNA DIRECT[™]

478 Purification kit (Thermo Fisher Scientific, Waltham, MA) per manufacturer 479 recommendations for plant tissue. Library construction was done following the 480 protocol of Zhong *et al.* [72]. The resulting libraries were sequenced in single-end 481 mode in a HiSeq 3000 at the UC Davis DNA Technologies Core 482 (http://dnatech.genomecenter.ucdavis.edu/). The quality of the libraries was 483 assessed using FastQC V0.11.2 [73]. Bacterial and human contaminants were filtered 484 with BWA-MEM [74] mapping against custom references of microbial and human 485 databases. The remaining reads were mapped to a joint reference made up of the L. 486 *sativa* cv. Salinas (GenBank: GCF_002870075.1) and *B. lactucae* isolate SF5 (GenBank: 487 GCA 004359215.1) reference assemblies using STAR v2.6.0 [75]. STAR was run with 488 the options "--sjdbOverhang 99 --sjdbGTFtagExonParentGene Parent --quantMode 489 GeneCounts". The strand specific read counts for each replicate were calculated from 490 the reads per gene table output by STAR. Reads counts were normalized for gene 491 length by dividing the read counts by the length of the gene it was mapped to (Reads 492 Per Kilobase: RPK). The total RPK of the *B. lactucae* portion of each replicate was 493 calculated and used to divide the RPK to calculate the transcripts per million (TPM) 494 for each gene. A subset of the genes that encode putative WY domains were taken 495 from this total, the average TPM for three replicates of each time point was calculated 496 (excluding replicate 2 at 72 hours, due to low coverage) and the log10(1+average 497 TPM) was calculated for each time point. These values were plotted using Heatmap2 498 [76] from the package gplots, clustering of putative effectors by expression was 499 performed with hclust [77].

501 Gateway cloning

502 Effector candidates without their signal peptide were cloned into the 503 pEarleyGate100 (pEG100) vector for plant expression [78] using Gateway cloning 504 (Thermo Fisher Scientific). Candidate effector genes were amplified from genomic 505 DNA isolated from spores of *B. lactucae* isolate SF5 or C82P24 using Phusion High 506 Fidelity Polymerase (Thermo Fisher Scientific) and primers (Suppl. File 3) that 507 amplified each ORF after the predicted signal peptide cleavage site. The Kozak 508 sequence (ACCATG) was added to the forward primer for correct translational 509 initiation. PCR products were purified using polyethylene glycol (PEG) precipitation 510 to remove primers and primer-dimers, recombined into pDONR207 using BP Clonase 511 II (Thermo Fisher Scientific), and transformed into chemically competent Escherichia 512 *coli* DH10B cells. The resulting entry clones were sequenced using primers designed 513 to pDONR207 to confirm gene identity and identify alleles. The entry clones were 514 then recombined into pEG100 using LR Clonase II and transformed into E. coli to 515 generate expression clones. The expression clones were then transformed into 516 *Aarobacterium tumefaciens* strain C58rif⁺ using electroporation. Kanamycin-resistant 517 colonies were confirmed for the transgene using gene-specific primers.

518

519 Agrobacterium-mediated transient assays

Agroinfiltration and transient expression experiments were performed using conditions optimized for lettuce [79]. *A. tumefaciens* was grown overnight from glycerol stocks and resuspended in 10 mM MgCl₂ to OD=0.3–0.5. The youngest fully expanded leaves of 3 to 4-week-old greenhouse grown lettuce plants were infiltrated

524 with the *A. tumefaciens* cultures using a needless syringe. Leaves were examined four 525 to five days post-infiltration for signs of macroscopic cell death, indicative of immune 526 recognition by host resistance proteins. A. tumefaciens containing pEG100 empty 527 vector or pEG100:GFP were used as negative controls; A. tumefaciens containing T-528 DNAs that expressed HopM1 or AvrPto, which are *Pseudomonas syringae* effectors 529 known to elicit cell death in lettuce [80], were used as positive controls. To control 530 for false negatives, leaves that did not show cell death in response to the positive 531 control HopM1 were excluded from the analysis. To control for false positives, leaves 532 that showed cell death in response to the empty vector or GFP control were also 533 excluded from the analysis. In addition, any cell death observed in the initial 534 screening was confirmed by repeating infiltrations on at least four plants for effectors 535 found to cause cell death.

536

537 Subcellular localization prediction and characterization

538 Nuclear localization was predicted using NucPred [44] and nuclear 539 localization motifs predicted using LOCALIZER [81]. For localization experiments, 540 effectors were cloned without their signal peptide into pEG104 [31] as described 541 above resulting in an N-terminal YFP fusion. Three to five days post-Agroinfiltration, 542 lettuce leaves expressing N-terminal YFP fusions of effectors were examined for 543 subcellular localization. Nuclear staining was performed by incubating cut leaf tissue 544 in 18 nM DAPI in water for at least five minutes. Imaging was performed on a Zeiss 545 LSM 710 laser scanning confocal microscope with a 40x objective lens.

546

547 PTI suppression assay

548 Effectors were expressed in five-week-old plants of *N. benthamiana* using 549 Agroinfiltration as described above. Two days post-infiltration, two leaf discs (3.8 550 mm) were taken from each infiltration site away from the leaf veins using a cork 551 borer. Leaf discs were floated abaxial side up in 200 µL of distilled water in a 96-well 552 white assay plate and incubated at room temperature for 24 hrs. To measure 553 suppression of ROS production [82], the water was removed and 100 μ L of assay 554 solution was added to the leaf discs, which contained 17 µg/mL luminol (Sigma-555 Aldrich, St Louis, MO) and $10 \,\mu\text{g/mL}$ horseradish peroxidase type 6A (Sigma-Aldrich). 556 One of the two paired leaf discs was exposed to 100 nM of flg22 peptide (flg+) and the 557 other was not exposed to any elicitor as a control for endogenous ROS production. 558 The known bacterial suppressors of PTI, HopS2, and HopX1 [83], were used as 559 positive controls for suppression of ROS production; GFP was used as the negative 560 control. Luminescence was measured on a FilterMax F5 Plate Reader (Molecular 561 Devices, San Jose, CA) promptly after adding in the assay solution and was measured 562 every two minutes for a total of 42 minutes. Each assay was replicated sixteen times.

563

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effector proteins are available in Supplementary data. The all data for interactions
between effectors and individual lines of *Lactuca* spp. are available at
http://bremia.ucdavis.edu/BIL/BIL interaction.php.

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- 884
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- 888

889 Figure legends

890

891	Figure 1. WY effector candidates from <i>B. lactucae</i> isolate SF5. (A) UPGMA
892	consensus tree of the 59 predicted WY effectors from <i>B. lactucae</i> isolate SF5 based
893	on whole protein amino acid sequence alignment using MUSCLE. Sequences that
894	appear to be allelic are indicated with asterisks next to the sequence name.
895	Bootstrap values and branch lengths are given for closely related proteins. Signal
896	peptides are shown as circles (red circle for SP; white circle for no SP). RXLR motifs
897	are shown as triangles (black triangle for RXLR ([RQGH]xLR or RXL[QKG]); grey
898	triangle for degenerate RXLR ([RKHGQ][X]{0,1}[LMYFIV][RNK]), white triangle for
899	no RXLR-like sequence) followed by an inverted triangle for EER motifs (yellow
900	triangle for [DE][DE][RK], white triangle for no EER-like sequence). WY domain
901	architecture is shown using blue rectangles, with a black line representing the total
902	length of the protein and rectangle position representing the location of the WY
903	motif as predicted by HMMer 3.0. (B) Sequence logo for the WY domain from <i>B</i> .
904	<i>lactucae</i> built from a multiple sequence alignment of WY domains predicted by
905	HMMer 3.0.

906

907 Figure 2. Distribution of predicted secreted WY effectors with or without RXLR 908 and/or EER motifs in the genomes of downy mildew pathogens and

909 *Phytophthora* species. HMMer was used to search predicted secretomes for each

910 species for the WY domain. For the WY domaining containing proteins, the presence

911	of RXLR and EER were determined by searching for [RQGH]XLR or RXL[QKG] within
912	the first 60 amino acids after the signal peptide and [DE][DE][KR] within the first
913	100 amino acids after the signal peptide. For the non-WY domain containing
914	proteins, the RXLR and EER was determined by searching for a strict "RXLR" with
915	[DE][DE][KR], within the first 60 and 100 amino acids, respectively, plus proteins
916	found by searching for the RXLR-EER domain using the RXLR-EER HMM from [14].
917	
918	Figure 3. Intrinsic disorder in the first 150 N-terminal amino acid sequences
918 919	Figure 3. Intrinsic disorder in the first 150 N-terminal amino acid sequences of RXLR and/or WY containing candidate effectors in <i>B. lactucae</i> . Proteins were
919	of RXLR and/or WY containing candidate effectors in <i>B. lactucae</i> . Proteins were
919 920	of RXLR and/or WY containing candidate effectors in <i>B. lactucae</i> . Proteins were categorized as WY with no RXLR (yellow), WYs with RXLR (black), RXLR+EER (with
919 920 921	of RXLR and/or WY containing candidate effectors in <i>B. lactucae</i> . Proteins were categorized as WY with no RXLR (yellow), WYs with RXLR (black), RXLR+EER (with or without WY, purple), and the total predicted secretome (blue). RXLR and EER

925 Figure 4. Lineage specificity of *B. lactucae* candidate secreted WY effectors.

926 Secreted WY proteins predicted in isolate SF5 of *B. lactucae* were used as a query for

927 BLASTp against other oomycete translated ORFs. Isolate C82P24 was queried for *B*.

928 *lactucae.* The best BLAST hit percent identity for each protein was calculated. The

box plot shows the distribution of the best BLAST hits per *B. lactucae* WY protein

930 from each species, with each dot representing an individual data point. No hits were

931 observed to Albugo laibachii, Saprolegnia parasitica, or Pythium ultimum.

932

933 Figure 5. RNA-seq expression levels during infection for the *B. lactucae* WY

effector candidates. The transcripts per million (TPM) of each WY-encoding gene
was normalized by the total number of reads per kilobase assigned to *B. lactucae*from RNAseq analysis of infected cotyledons over a seven-day time-course. This

937 value was log10(+1) transformed to enable visualization. The four highest

938 expressed WY encoding genes do not have recognizable RxLR translocation

- 939 associated motifs.
- 940

941 Figure 6. Results of the screen for recognition of *B. lactucae* WY effector by

942 **diverse genotypes of lettuce.** Candidate effectors were expressed in lettuce leaves

943 using Agroinfiltration and leaves were scored for their reaction four to seven days

944 post-infiltration. Qualitative scores were converted into numeric scores for data

analysis: 0 = no reaction (green), 1 = mild chlorosis (blue-green), 2 = chlorosis

946 (yellow), 3 = mild necrosis or mixture of chlorosis and necrosis (pink), and 4 = full

947 necrosis (magenta). The figure shows average scores for each effector on each

948 genotype; only accessions and effectors that had a necrotic or chlorotic response are

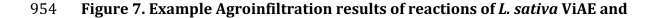
949 shown. Scores for all accessions screened and sample sizes can be found at

950 http://bremia.ucdavis.edu/BIL/BIL_interaction.php. Two groups of 28 and 11

951 accessions of *L. saligna* that had identical reactions are shown together; the

952 individual accessions making up each group are shown in Supplementary Table 2.

953



955 ViCQ and their progenitor R gene donors L. virosa LS238 and LS241 to several

- 956 *B. lactucae* effectors. Photos are representative of a typical leaf. *B. lactucae*
- 957 candidate effectors either elicited necrosis (brown areas) indicative of an immune
- 958 recognition response (magenta text) or did not elicit a response (aqua text). GFP
- and HopM1 were used as negative and positive controls for necrosis, respectively.
- 960 Leaves were collected five days post-infiltration and the first column for each
- 961 accession shows the uncleared leaf tissue; the second column shows leaf tissue
- 962 cleared in ethanol.
- 963

964 Figure 8. Subcellular localization of three *B. lactucae* WY effectors in lettuce.

- 965 Confocal images of lettuce expressing YFP fusion proteins. For YFP-SW4, split
- 966 images of (A) DAPI, (B) Chloroplast autofluorescence, (C) YFP, and (D) Merged show
- 967 nuclear localization. For YFP-SW1 (E) and YFP-SW3 (F), merged images of DAPI
- 968 (blue), chlorophyll autofluorescence (red), and YFP (yellow) show cytoplasmic or
- 969 periplasmic localization.
- 970
- 971

972 Figure 9. Effect of *B. lactucae* candidate WY effectors on flg22-triggered

973 immune response. Luminescence plots measuring effect of candidate effectors on

- 974 flg22-triggered PTI induction in *N. benthamiana* leaf discs. (A) Luminescence
- 975 averaged across biological replicates (14–24 reps each) for each effector over time.
- 976 (B) Bar plots of the average area under the curve (AUC) for luminescence over 40

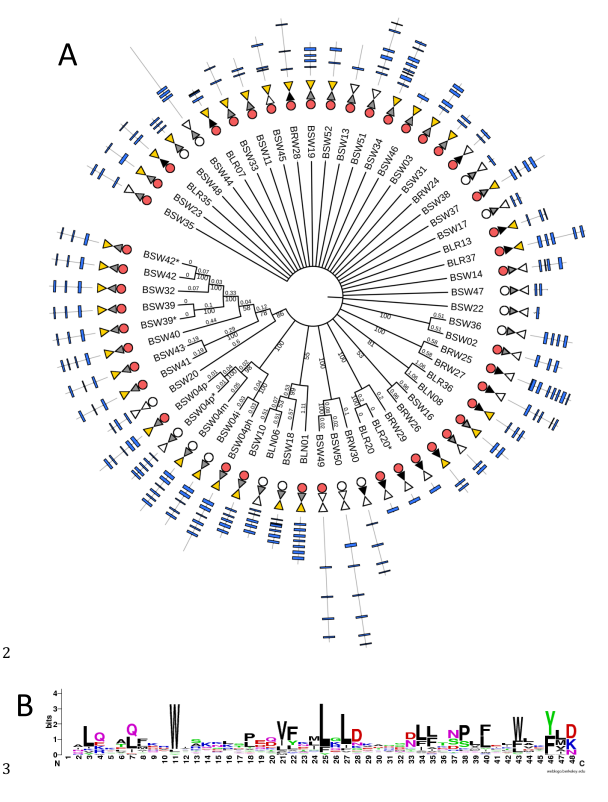
- 977 minutes for each effector (N=14–24), with flg22 (red) or without flg22 (aqua).
- 978 Asterisks indicate averages in the flg22+ condition that are statistically different
- from the average of GFP in the flg22+ condition (t-test, p < 0.05).

980

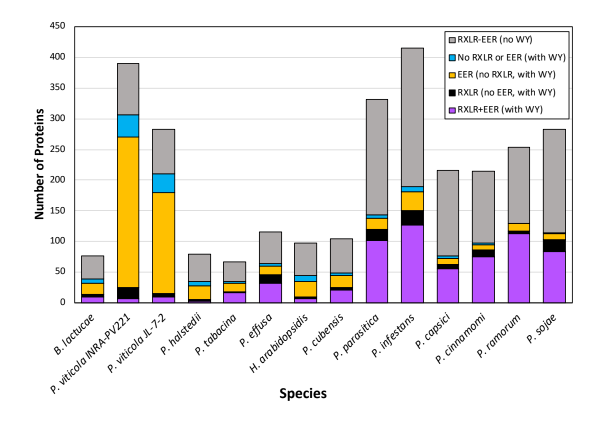
981 List of Supplemental Materials:

982 **Supplemental Table 1.** RXLR-like sequences observed within 100 amino acids of 983 the N-terminus of *B. lactucae* WY proteins, sorted by first amino acid. 984 985 Supplemental Figure 1. The number of RXLR-like motifs detected in the first 100 986 amino acids of the N-terminus of *B. lactucae* WY effectors. 987 988 **Supplemental Figure 2.** Intrinsic disorder of first 150 amino acids of RXLR and WY 989 containing candidate effectors mined from other oomycete genomes. 990 991 **Supplemental Figure 3.** Agroinfiltration results from BLN06-SF5 and sequence 992 comparison between BLN06 between SF5 and BL24. 993 994 **Supplemental File 1.** Sequences and NCBI reference numbers of the 59 WY 995 proteins predicted from the *B. lactucae* SF5 genome assembly. 996 997 **Supplemental File 2.** Lettuce genotypes tested for recognition of *B. lactucae* 998 effectors. 999 1000 **Supplemental File 3.** Primers used in Gateway Cloning of the *B. lactucae* effectors. 1001 1002

1 Figures and Tables

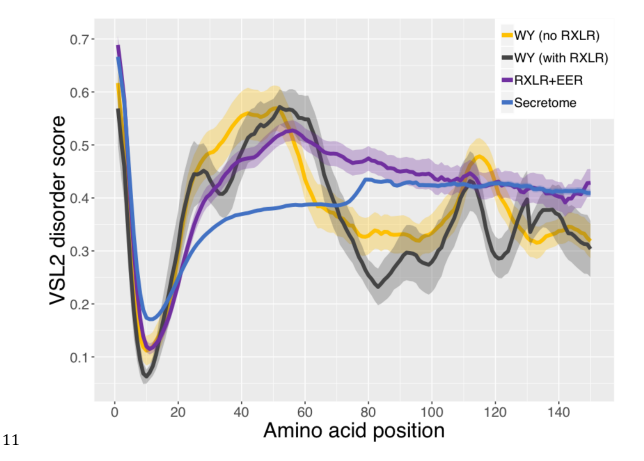


4 Figure 1. WY effector candidates from *B. lactucae* isolate SF5.



6

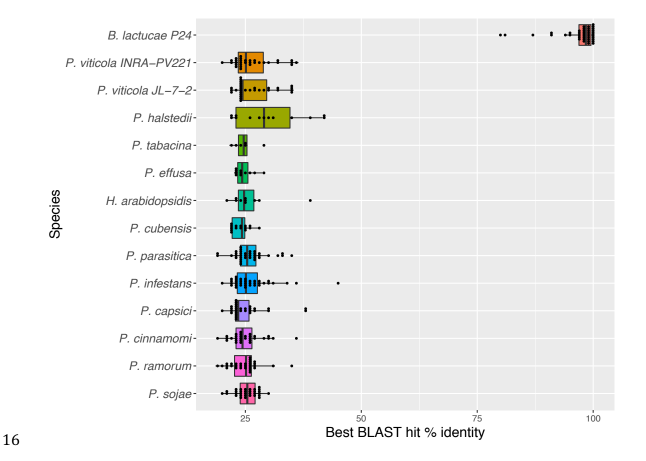
- 7 Figure 2. Distribution of predicted secreted WY effectors with or without RXLR
- 8 and/or EER motifs in the genomes of downy mildew pathogens and
- 9 Phytophthora species.



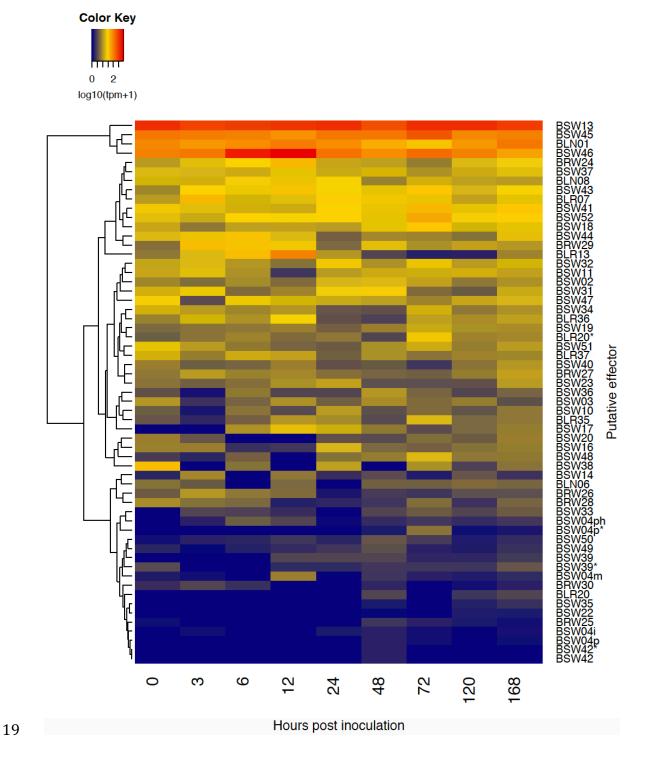
12 Figure 3. Intrinsic disorder in the first 150 N-terminal amino acid sequences

13 of RXLR and/or WY containing candidate effectors in *B. lactucae*.

14



17 Figure 4. Lineage specificity of *B. lactucae* candidate secreted WY effectors.



20 Figure 5. RNA-seq expression levels during infection for the *B. lactucae* WY

21 effector candidates.

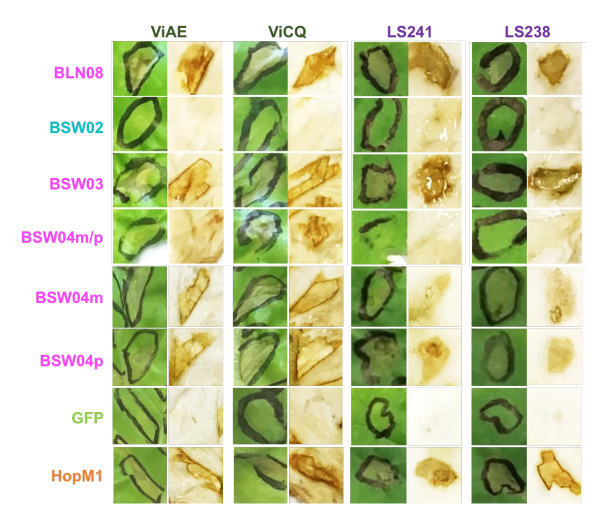
		ß	9	90	8	03	BSW04m	BSW04m/p	04p	11a	11b	13	14	19		trol
Genotype	species	BLR13	BLR36	3LN06	3LN08	BSW03	MS:	NS!	BSW04p	BSW11a	BSW11b	BSW13	BSW14	3SW19	GFP	contro
CGN14312	indica	1.3	3.0	3.3						1.5	1.5	1.5	2.8	1.0	0.8	3.9
CGN14312	indica	0.8	1.5	0.0	0.5	0.4	1.5	2.3	1.0	1.0	1.3	1.5	0.6	0.0	0.4	3.8
CGN5282	saligna	0.0	0.0	0.0	2.5	0.0	0.0	1.5	0.0	0.0	0.0	4.0	4.0	0.0	0.0	4.0
CGN5309	saligna	0.0	2.8	0.0	3.8	0.3	0.6	0.3	1.0	0.0	0.0	4.0	4.0	0.0	0.7	4.0
PI491000	saligna	0.0	0.0	0.0	4.0	0.0	0.0	0.0	0.0	0.3	0.0	1.5	1.6	0.0	0.0	3.9
CGN5329	saligna	0.0	0.0	0.0	2.4	0.1	0.9	1.0	1.8	0.0	0.0	0.0	2.8	2.5	0.3	3.9
CGN5330	saligna	0.0	0.0	0.8	3.3	0.0	0.6	1.0	3.1	0.0	0.0	0.0	3.5	0.0	0.1	4.0
CGN5322	saligna	0.0	0.0	0.0	3.5	0.2	0.1	1.0	2.4	0.0	0.0	0.0	4.0	0.0	0.1	4.0
PI491208	saligna	0.0	0.0	0.0	3.6	0.6	0.6	2.0	3.6	0.0	0.0	0.0	4.0	0.0	0.7	4.0
CGN5318	saligna	0.0	0.0	0.0	3.5	0.8	0.3	2.0	0.3	0.0	0.0	0.0	4.0	0.0	0.1	3.7
PI491207	saligna	0.0	0.0	0.0	4.0	1.1	2.0	2.0	0.5	0.0	0.0	0.0	4.0	0.0	0.2	4.0
25 acc.	saligna	0.0	0.2	0.0	3.0	0.1	0.1	0.3	0.1	0.0	0.0	0.0	3.5	0.1	0.1	3.9
11 acc.	saligna	0.0	2.9	0.0	3.6	0.4	0.3	0.5	0.3	0.0	0.0	0.1	4.0	0.0	0.2	4.0
CGN5309	saligna	0.0	2.5	0.0	3.7	0.3	0.6	0.3	1.0	0.0	0.0	4.0	4.0	0.0	0.3	4.0
CGN5314	saligna	0.0	3.3	0.0	3.8	0.8	0.8	0.8	1.5	0.0	0.0	0.0	4.0	0.0	0.4	4.0
CGN5315	saligna	0.0	1.8	0.0	3.9	0.3	1.0	0.8	1.9	0.0	0.0	0.0	4.0	0.0	0.0	4.0
Blonde	sativa	0.5	0.5	0.0	0.0	0.5	0.5	1.0	0.0	2.0	2.7	0.5	0.0	0.0	0.2	4.0
Capitan	sativa	0.0	0.0	0.0	0.6	0.2	0.2	3.3	3.5	0.0	0.0	0.0	0.0	0.0	0.2	3.8
Capsule	sativa	0.0	0.0	0.0						0.0	0.0	0.0	4.0	0.0	0.0	4.0
Femke	sativa	0.0	0.0	0.0	0.0	0.0	2.4	3.8	3.0	0.0	0.0	0.0	0.0	0.0	0.0	4.0
Fenston	sativa	0.0	0.0	1.0	0.0	0.6	1.3	1.5	2.8	0.0	0.0	0.0	0.0	0.0	0.4	3.6
FrRsal-1	sativa	0.0	1.5	0.0	0.0	0.0	0.1	3.4	0.0	0.0	0.0	0.0	0.0	0.0	0.0	3.8
Ninja	sativa	0.0	1.2	0.0	0.0	0.0	0.2	2.0	3.4	0.0	0.6	0.0	0.0	0.0	0.2	4.0
Pennlake	sativa	0.0	0.0	0.0	0.0	0.0	0.0	2.1	0.0	0.4	0.0	0.0	0.0	0.0	0.0	3.5
PI491226	sativa	0.0	0.0	0.0	3.7	0.6	0.5	1.0	1.5	0.0	0.0	0.0	4.0	0.0	0.1	3.9
RYZ2164	sativa	0.0	0.0	0.0	0.0	0.0	0.0	3.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	4.0
Salvius	sativa	0.0	0.0	0.0	0.0	0.0	0.0	2.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	3.6
UCDM14	sativa	0.0	0.0	0.0	0.1	0.1	0.0	0.4	0.0	3.1	0.0	0.0	0.2	0.0	0.4	3.8
Versaï	sativa	0.0	0.0	0.4	0.0	0.0	0.7		0.0	3.2	2.3	0.0	0.0	0.0	0.0	4.0
ViAE	sativa	0.0	0.0	0.0	3.4	3.5	3.9	2.0	3.8	0.0	0.0	0.0	2.8	0.0	0.1	3.7
ViCQ	sativa	1.0	0.0	0.0	3.9	2.8	3.8	3.3	3.4	0.0	0.0	0.0	3.2	0.0	0.1	3.8
Arm999	serriola	0.0	0.0	0.0	0.0	0.0	3.6	4.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	4.0
CGN14280	serriola	2.2	0.0	0.5	0.3	0.0	0.0	0.3	0.4	0.0	0.0	0.0	0.0	0.0	0.3	3.3
ISR-380	serriola	0.5	0.5	1.0	0.0	1.0	0.5	0.5	0.5	0.5		0.3	1.0	2.5	0.4	4.0
LS102	serriola	0.0	0.0	0.0	0.0	0.3	0.0	0.0	0.0	0.0	0.0	1.0	2.3	1.3	0.2	4.0
PI491108	serriola	0.0	0.0	0.0	2.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	3.3	0.0	0.0	4.0
PI491178	serriola	0.0	0.0	0.0		0.0	0.0	0.0	0.0	4.0	0.0	0.0	0.0	0.0	0.0	4.0
CGN14305	virosa	0.0	0.0	0.0	1.5	0.7	0.9	1.5	1.4	0.0	0.0	0.0	2.3	0.0	0.4	3.6
CGN4683	virosa				3.5	0.0	0.0		0.0						0.0	4.0
CGN5333	virosa	0.0	0.0	0.5	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	2.3	1.5	0.0	3.9
CGN9365	virosa	0.0	0.0	0.0	0.2	0.0	0.8	0.0	0.0	0.0	0.0	0.0	3.8	0.0	0.0	4.0
LS238	virosa	0.0	0.0		1.8	2.0	3.9	0.0	3.6	0.0	0.0	0.0	2.0	0.0	0.0	3.4
LS241	virosa	0.0	0.0	0.0	2.5	2.3	3.9	0.0	3.2	0.0	0.0	0.0	3.5	0.0	0.0	3.1

Color KeyScoreNo reaction0Mild chlorosis1Chlorosis2Mild Necrosis3Necrosis4No data-

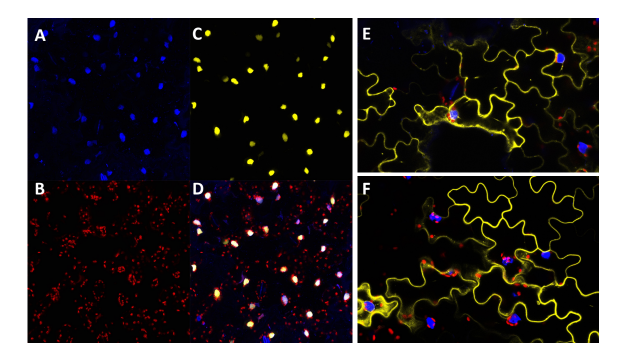
22

23 Figure 6. Results of the screen for recognition of *B. lactucae* WY effector by

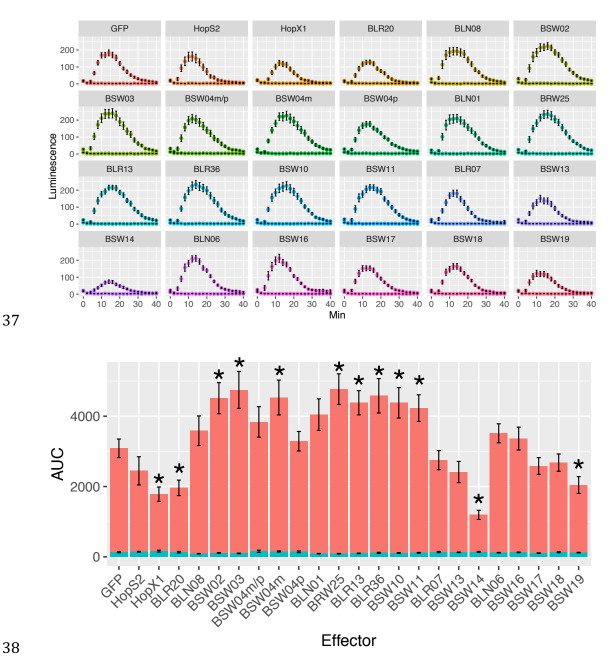
24 diverse genotypes of lettuce.

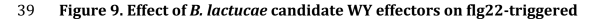


- 26 Figure 7. Example Agroinfiltration results of reactions of *L. sativa* ViAE and
- 27 ViCQ and their respective progenitor R gene donors *L. virosa* LS241 and LS238
- 28 to several *B. lactucae* effectors.
- 29
- 30



32 Figure 8. Subcellular localization of three *B. lactucae* WY effectors in lettuce.





immune response.