Secretome analysis reveals effector candidates associated with broad host range necrotrophy in the fungal plant pathogen *Sclerotinia sclerotiorum*

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http://www.biomedcentral.com/1471-2164/15/336
Secretome analysis reveals effector candidates associated with broad host range necrotrophy in the fungal plant pathogen *Sclerotinia sclerotiorum*

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**Abstract**

**Background:** The white mold fungus *Sclerotinia sclerotiorum* is a devastating necrotrophic plant pathogen with a remarkably broad host range. The interaction of necrotrophs with their hosts is more complex than initially thought, and still poorly understood.

**Results:** We combined bioinformatics approaches to determine the repertoire of *S. sclerotiorum* effector candidates and conducted detailed sequence and expression analyses on selected candidates. We identified 486 *S. sclerotiorum* secreted protein genes expressed *in planta*, many of which have no predicted enzymatic activity and may be involved in the interaction between the fungus and its hosts. We focused on those showing (i) protein domains and motifs found in known fungal effectors, (ii) signatures of positive selection, (iii) recent gene duplication, or (iv) being *S. sclerotiorum*-specific. We identified 78 effector candidates based on these properties. We analyzed the expression pattern of 16 representative effector candidate genes on four host plants and revealed diverse expression patterns.

**Conclusions:** These results reveal diverse predicted functions and expression patterns in the repertoire of *S. sclerotiorum* effector candidates. They will facilitate the functional analysis of fungal pathogenicity determinants and should prove useful in the search for plant quantitative disease resistance components active against the white mold.

**Keywords:** *Sclerotinia sclerotiorum*. Effectors, Gene expression, Secretome, Necrotrophic fungal, Pathogen, *Arabidopsis thaliana*

**Background**

The white mold fungus *Sclerotinia sclerotiorum* (Lib.) de Bary is a cosmopolitan necrotrophic pathogen infecting over 400 plant species. It is among the most devastating pathogens of soybean, rapeseed and sunflower, causing several hundred million dollar losses annually at the pre- and postharvest stages [1]. *S. sclerotiorum* host range is remarkably broad, with fruit and vegetable productions also being severely impacted [2]. *S. sclerotiorum* and its close relative the grey mould fungus *Botrytis cinerea* are among the few fungal pathogens considered as typical necrotrophs. As such, they derive energy to complete their life cycle mostly from dead plant cells, as opposed to biotrophs that feed on living plant cells.

There is now ample evidence that biotrophic and hemibiotrophic fungi secrete specialized effector proteins manipulating host cell physiology to obtain nutrients, suppress plant defense and ultimately promote infection [3]. Effectors may also trigger plant defense responses, leading to plant resistance, when recognized directly or indirectly by the plant immune system in a gene-for-gene relationship. This results from a co-evolutionary arms race between pathogen effectors, their plant targets, and components of the plant immune system [4]. Necrotrophs have long been considered as less adapted, secreting mostly degrading enzymes and toxins that unspecifically trigger programmed cell death (PCD) and dismantle plant cells.
However, host specific necrotrophs such as *Cochliobolus victoriae* secrete effector proteins translocated into plant cells that interact with specific corresponding host proteins to facilitate disease progression [5,6]. This involves the activation of plant PCD instead of its suppression as in the case of infection by biotrophic pathogens. *S. sclerotiorum* also produces the non-proteic pathogenicity determinant oxalic acid. This molecule induces the synthesis of reactive oxygen species (ROS) and triggers plant PCD late during infection, but has the opposite effect, suppressing ROS burst and PCD induction, at the early stages of infection [7]. The SSITL secreted integrin-like protein of *S. sclerotiorum* promotes virulence and delays the activation of plant defense responses, supporting the view that *S. sclerotiorum* secretes effectors to finely manipulate plant physiology [8]. In addition, enzymes secreted by necrotrophs can act as virulence factors independently of their catalytic activity [9]. Effector repertoires vary considerably, notably according to pathogens lifestyle [10], and it is becoming clear that interactions between necrotrophs and their host plants are considerably more complex and subtle than previously considered. What is the effector candidate repertoire associated with broad host range necrotrophy remains unclear. As a first step towards elucidating the molecular bases for colonization by *S. sclerotiorum*, its repertoire of effector candidates needs to be determined.

The recent release of genome sequences for a number of plant pathogenic fungi facilitated the search for effector candidates (ECs) at the genome level [11]. Nevertheless, considering that pathogen effector repertoires are typically lineage-specific, the identification of effectors remains challenging [4]. The analysis of *S. sclerotiorum* genome sequence uncovered sets of genes associated with the manipulation of redox status, including enzymes of OA biosynthesis, the degradation of plant cell wall, and 603 secreted proteins with other functions [12]. Known hallmarks of fungal effectors such as the presence of signal peptides and absence of transmembrane domains, small size and amino-acid composition generally produce lists of hundreds of potential effectors in a given pathogen. Therefore, more sophisticated approaches are required to pinpoint the most relevant ECs for the promotion of infection in *S. sclerotiorum* secretome.

A limited number of known fungal effector families show conservation at the sequence level or similar predicted functions. This is notably the case for the toxin and cell death elicitor proteins of the Necrosis and ethylene-inducing Like Proteins (NLPS), the ceratopterin, cyanovirin-N homology (CVNH) and ECP2 families [13-16]. The growing number of characterized fungal effectors suggests conservation at the biochemical function level in the overall effector repertoire of several fungal pathogens. The ability to bind chitin or other cell wall oligosaccharides, masking the presence of the pathogen or dampening damage-induced plant responses, is a feature common to effectors from multiple fungal pathogens [17-20]. Fungal effectors harboring a protease inhibitor activity are also common [21-24]. The biochemical activity of a few other fungal effectors such as *M. oryzae* Fungaly-sin metalloprotease AvrPita [25], *U. maydis* choromase mutant cmu1 [26] and peroxidase inhibitor PEP1 [27] may also be part of the arsenal of effector functions in multiple fungal lineages. This hypothesis suggests that thorough annotation of protein domains and prediction of biochemical function of secreted proteins may prove useful to identify novel effectors in *S. sclerotiorum*.

However, a majority of effectors do not show significant similarity to known sequences in other organisms nor obvious protein domains. Yet other genomic characteristics may help identify EC genes. The rapid evolution of effector genes allows the fungii to overcome selection pressures induced by resistant plant cultivars. A high ratio of non-synonymous over synonymous substitutions (Ka/Ks) in alleles from related strains is a frequently used proxy for inferring fast gene evolution and the action of positive selection [28]. This approach has been used to reveal ECs in several filamentous plant pathogen lineages [29-34]. Positive selection has been detected in *B. cinerea* genome [35] suggesting that it may be used to mine *S. sclerotiorum* genome for ECs. Second, gene duplication is another hallmark of several known fungal effector genes, such as the ToxB host specific toxin of *Pyrenophora tritici-repentis* [36,37]. Third, genomic regions with high repeat and transposable element content are enriched in effector genes in several lineages of plant pathogens [37-39] suggesting that genome architecture analysis can assist in the search for EC genes. Finally, effectors can alter host cell function by mimicking plant peptides [40]. These ECs likely elude functional annotation on the basis of primary amino-acid sequence, but may be revealed using three-dimensional structure prediction.

As opposed to Oomycete pathogen genomes in which many effector genes can be identified through conserved sequence motifs [29,41], the use of conserved sequence motifs, such as the Y/F/WxX motif [42], has proven limited in revealing fungal effectors across lineages. However, the presence of a signal peptide directing protein secretion and gene expression in planta are relatively universal properties of effectors that can be exploited as first filters to narrow down the list of effector candidates in fungal genomes.

*S. sclerotiorum* effector proteins would be useful as probes to search for resistance components in plants and to design strategies for inhibiting infections by this devastating but poorly characterized pathogen. In this study, we report a diverse repertoire of *S. sclerotiorum* effector candidates revealed by an in depth analysis of its predicted secretome. We combined refined secretome
annotation, phylogeny, selection and gene duplication analyses, and three-dimensional structure prediction to identify 78 ECs. Among those, we highlight a predicted subtilisin inhibitor, a xylanase, a duplicated gene of unknown function and three toxin mimics as high priority candidates for functional studies. We analyzed in planta expression pattern for 16 EC genes and revealed host-blind and host-regulated ECs.

Results
Definition and annotation of *S. sclerotiorum* secretome
In the original analysis of *S. sclerotiorum* genome, secreted proteins were predicted using SignalP, TargetP and TMHMM, and annotated using Interproscan [12]. To identify and explore candidate effectors in the genome of *S. sclerotiorum*, we built a bioinformatics workflow exploiting genomic features typical of known filamentous plant pathogen effectors. We first refined the prediction of secreted proteins combining predictions from SignalP2 and 4 to identify 1070 proteins with a secretion signal. To evaluate the sensitivity of this prediction, we applied the same methodology on a list of 1985 eukaryotic secreted proteins validated experimentally from the SPdb [43]. We retrieved 1971 proteins predicted as secreted, corresponding to a true positive rate of 99.34%. Next, we removed 172 proteins predicted by TMHMM to harbor transmembrane helices, then 153 proteins predicted by GPIsom to harbor glycosphatidylinositol anchor motifs, that likely represent surface proteins rather than secreted effectors. This resulted in a list of 745 predicted secreted proteins. To increase the likelihood of identifying genes relevant for infection, we selected those for which there is evidence for expression in planta based on publicly available EST and microarrays data. We considered genes expressed, even if not induced, during interaction with rapeseed, sunflower or tomato (see Methods). Although this filtering pipeline likely excluded relevant effector candidates, it resulted in a total of 486 genes encoding predicted secreted proteins expressed in planta (SPEPs, Figure 1a, Additional file 1: Table S1).

Next, we performed three different effector-oriented analyses on the 486 SPEP genes. First, we used Blast2GO, PFAM domain and nuclear localization signal (NLS) searches to annotate 326 SPEPs (Figure 1b). We built a database of known fungal effectors and explored the literature to select 31 *S. sclerotiorum* ECs among annotated SPEP genes. Second, we defined clusters of orthologous genes (COGs) between *S. sclerotiorum* and *B. cinerea* predicted genes using Inparanoid (Figure 1c). A total of 197 SPEP genes grouped in COGs. We aligned *S. sclerotiorum* and *B. cinerea* orthologs for these 197 gene pairs and calculated ratios of non synonymous over synonymous substitutions (Ka/Ks) to identify five ECs with signature of positive selection. The 289 SPEP genes with no ortholog...
in B. cinerea were grouped into clusters based on sequence similarity to identify 29 ECs distributed in 24 families containing genes duplicated in S. sclerotiorum. Finally, we analyzed the taxonomic distribution of SPEP genes across the kingdom Fungi using BlastP searches against a database of 14 complete genomes representative of all major fungal lineages. This identified 70 S. sclerotiorum-specific SPEP genes, most of which had no annotation (Figure 1d). We used protein structure prediction and pattern and fold recognition searches to identify 17 ECs analogous to known protein fold encoded by S. sclerotiorum-specific SPEP genes.

Using effector-oriented analyses, we identified four lists of ECs, containing a total of 78 EC genes (four being common to two lists). We could not predict any enzymatic activity encoded by 33% of the SPEP genes (160), suggesting that S. sclerotiorum effector repertoire encodes diverse functions that are not restricted to plant cell degrading enzymes. Besides, 59.5% of the SPEP genes (289) did not cluster in B. cinerea COGs, revealing a relatively high degree of divergence from this closely related fungal pathogen.

**Sclerotinia effector candidates showing conserved domains**

Some fungal effectors show conserved protein domains and biochemical functions that can be identified at the protein sequence level. In a first approach to identify ECs, we used (i) PFAM annotations, (ii) nuclear localization signal (NLS) predictions, and (iii) BlastP searches against known fungal effectors. We found 326 annotated SPEP genes among which we selected 13 as effector candidates based on the presence of specific PFAM domains (Table 1). Indeed, six predicted SPEPs (SS1G_00642, SS1G_02014, SS1G_04786, SS1G_09392, SS1G_12336, SS1G_14184) included chitin-recognition or chitin-binding domains (PF00187, PF03067) and one SPEP (SS1G_12509) contained LysM domains (PF01476). SS1G_07836 was annotated as a concanavalin A-like lectin/glucanase that binds to complex carbohydrates, and harbors a Peptidase A4 (PF01828) with a typical lectin fold. SS1G_08698 contains a Ricin type beta-trefoil lectin domain (PF00652) typical of ricin-like toxins. These predicted SPEPs are relevant ECs considering that chitin- and carbohydrate-binding activity has been reported for several fungal effector families. We also selected three predicted SPEPs (SS1G_01593, SS1G_03282, SS1G_12605) with protease inhibitor domains (PF05992), another activity found in known fungal effectors. We identified two predicted SPEPs (SS1G_03611 and SS1G_13935) with a CFEM cystein-rich fungal effector motif (PF05730). Second, we selected another 11 SPEP genes based on the presence of at least one predicted NLS in their sequence (Table 1). The co-occurrence of a secretion signal and a NLS in encoded proteins suggest that they may be active in the plant nucleus. Consistent with a nuclear activity, SS1G_01866 and SS1G_05895 also harbored transcription factor domains. Third, we selected 7 predicted SPEPs that were retrieved based on sequence similarity to known effectors or ECs from other plant pathogenic fungi (Table 1). We recovered the two necrosis and ethylene inducing peptides SsNEP1 and SsNEP2 (SS1G_03080 and SS1G_11912) described in [44] sharing ~40% identity with various NEP-like proteins, and harboring a characteristic PFAM domain (PF05630). We selected SS1G_00849 for sharing 59% identity with Colletotrichum hingginsiam effector candidate 91 (CHEC91, [45]). There was no protein domain identified in this protein but three-dimensional structure prediction indicates that it is analogous to Alternaria alternata AltA-1 allergen. We selected SS1G_02904 for sharing 35% identity with C. hingginsiam effector candidate 80 (CHEC80, [45]). It features a Cyanovirin-N homology domain (PF13639) that corresponds to a carbohydrate-binding module [15,46]. We identified SS1G_08858 showing 25% identity with M. oryzae effector AvrPita (MGG_15730, [25]) and harboring a M35 metalloprotease domain (PF02102). Finally we selected SS1G_10096 as similar (>60% identity) to pathogen-associated molecular patterns (PAMP) with a cerato-platanin domain (PF07249) [14,45,47,48]. These 31 ECs identified based on protein annotations showed that the activity of S. sclerotiorum secretome may not be limited to typical cell death elicitors but is predicted to cover a wide range of functions known for fungal effectors, including chitin binding, proteases and protease inhibitors.

**A family of subtilisin-inhibitor effector candidates conserved in Ascomycetes**

Effectors with a protease inhibitor activity have been described in fungal plant pathogens with a biotrophic phase of infection [22,49]. We identified three S. sclerotiorum candidate effectors (SS1G_01593, SS1G_03282 and SS1G_12605, Table 1) with a serine protease inhibitor 19 domain (PF05992) that corresponds to the propeptide inhibitor domain of subtilisins. To document the taxonomic distribution of SS1G_01593 homologs in fungi, we performed a BlastP search against the predicted proteome of 234 fungal species. Using signal peptide predictions, we identified 99 secreted homologs across 97 species. Secreted homologs of SS1G_01593 are restricted to Ascomycetes and found in all Leotiomycete species considered (Figure 2a). At the sequence level, homologs of SS1G_01593 are clearly distinct from C. fulvum Avr2, U. maydis Pit2, Uromyces fabae RTP1p and M. lini AvrP123, suggesting that SS1G_01593 family represent a distinct class of protease inhibitor effectors. To get insights into SS1G_01593 function and evolution, we predicted the 3D structure of the protein and mapped residue conservation among the 99 secreted homologs on this structure. SS1G_01593
| Protein ID  | Annotation                        | PFAM domain(s)                                                                 | Length | Comments                                                                 |
|------------|-----------------------------------|--------------------------------------------------------------------------------|--------|--------------------------------------------------------------------------|
| SSIG_00642 | Chitin binding protein            | Chitin recognition protein (PF00187.14)                                        | 563    |                                                                          |
| SSIG_02014 | Chitin binding protein            | Chitin binding domain (PF03067)                                                 | 426    |                                                                          |
| SSIG_04786 | Chitin binding protein            | Chitin recognition protein (PF00187.14)                                        | 399    |                                                                          |
| SSIG_09392 | Starch binding domain containing  | Chitin binding domain (PF03067)                                                 | 398    |                                                                          |
| SSIG_12336 | Chitin binding protein            | Chitin binding domain (PF03067)                                                 | 294    |                                                                          |
| SSIG_14184 | Agglutinin isolectin 3-like       | Chitin recognition protein (PF00187.14)                                        | 245    |                                                                          |
| SSIG_12509 | LysM domain protein               | LysM (PF01476)                                                                  | 447    |                                                                          |
| SSIG_07836 | Concanavalin A lectin glucanase   | Peptidase_A4 (PF01828)                                                          | 252    |                                                                          |
| SSIG_08698 | Ricin-type toxin                  | Ricin-type beta-trefoil lectin domain (PF00652)                                 | 409    |                                                                          |
| SSIG_01593 | Serine protease inhibitor         | Peptidase inhibitor I9 (PF05922)                                               | 95     |                                                                          |
| SSIG_03282 | Serine protease inhibitor         | Peptidase inhibitor I9 (PF05922)                                               | 522    |                                                                          |
| SSIG_12605 | Alkaline serine protease alp1     | Peptidase inhibitor I9 (PF05922)                                               | 400    |                                                                          |
| SSIG_03611 | Cystein-rich protein              | CFEM (PF05730)                                                                  | 119    |                                                                          |
| SSIG_13935 | Cystein-rich protein              | CFEM (PF05730)                                                                  | 529    |                                                                          |

**Table 1 List of 31 S. sclerotiorum effector candidates selected based on their annotation**

**SPEPs containing PFAM domains found in fungal effectors**

| Protein ID  | Annotation                        | PFAM domain(s)                                                                 | Length | Comments                                                                 |
|------------|-----------------------------------|--------------------------------------------------------------------------------|--------|--------------------------------------------------------------------------|
| SSIG_00642 | Chitin binding protein            | Chitin recognition protein (PF00187.14)                                        | 563    |                                                                          |
| SSIG_02014 | Chitin binding protein            | Chitin binding domain (PF03067)                                                 | 426    |                                                                          |
| SSIG_04786 | Chitin binding protein            | Chitin recognition protein (PF00187.14)                                        | 399    |                                                                          |
| SSIG_09392 | Starch binding domain containing  | Chitin binding domain (PF03067)                                                 | 398    |                                                                          |
| SSIG_12336 | Chitin binding protein            | Chitin binding domain (PF03067)                                                 | 294    |                                                                          |
| SSIG_14184 | Agglutinin isolectin 3-like       | Chitin recognition protein (PF00187.14)                                        | 245    |                                                                          |
| SSIG_12509 | LysM domain protein               | LysM (PF01476)                                                                  | 447    |                                                                          |
| SSIG_07836 | Concanavalin A lectin glucanase   | Peptidase_A4 (PF01828)                                                          | 252    |                                                                          |
| SSIG_08698 | Ricin-type toxin                  | Ricin-type beta-trefoil lectin domain (PF00652)                                 | 409    |                                                                          |
| SSIG_01593 | Serine protease inhibitor         | Peptidase inhibitor I9 (PF05922)                                               | 95     |                                                                          |
| SSIG_03282 | Serine protease inhibitor         | Peptidase inhibitor I9 (PF05922)                                               | 522    |                                                                          |
| SSIG_12605 | Alkaline serine protease alp1     | Peptidase inhibitor I9 (PF05922)                                               | 400    |                                                                          |
| SSIG_03611 | Cystein-rich protein              | CFEM (PF05730)                                                                  | 119    |                                                                          |
| SSIG_13935 | Cystein-rich protein              | CFEM (PF05730)                                                                  | 529    |                                                                          |

**SPEPs containing nuclear localization signal (NLS)**

| Protein ID  | Annotation                        | PFAM domain(s)                                                                 | Length | Comments                                                                 |
|------------|-----------------------------------|--------------------------------------------------------------------------------|--------|--------------------------------------------------------------------------|
| SSIG_01866 | Ring-7 protein                    | PHD-like zinc-binding domain (PF13771); Ring finger domain (PF13639)          | 425    | NLS (118–141)                                                            |
| SSIG_03146 | -                                 | -                                                                              | 193    | NLS (147–155)                                                            |
| SSIG_04309 | -                                 | -                                                                              | 177    | NLS (82–87)                                                              |
| SSIG_05895 | Zinc finger CCCH-type domain      | Zinc finger C-x8-C-x5-C-x3-H type (PF00642)                                    | 310    | NLS (256–291)                                                            |
| SSIG_09383 | -                                 | -                                                                              | 212    | NLS (143–150)                                                            |
| SSIG_06787 | -                                 | -                                                                              | 430    | NLS (189–250; 340–349)                                                   |
| SSIG_06890 | -                                 | -                                                                              | 284    | NLS (66–120; 143–167; 225–245)                                            |
| SSIG_07404 | -                                 | DUF3108 (PF11306)                                                              | 282    | NLS (65–79; 118–138; 223–233)                                            |
| SSIG_09050 | -                                 | -                                                                              | 454    | NLS (410–413)                                                            |
| SSIG_11108 | Ribosomal protein s17             | -                                                                              | 373    | NLS (338–348)                                                            |
| SSIG_13142 | -                                 | -                                                                              | 131    | NLS (89–99)                                                              |

**SPEPs showing homology to fungal effectors or fungal effector candidates**

| Protein ID  | Annotation                        | PFAM domain(s)                                                                 | Length | Comments                                                                 |
|------------|-----------------------------------|--------------------------------------------------------------------------------|--------|--------------------------------------------------------------------------|
| SSIG_03080 | SnNEP1                            | Necrosis inducing protein (PF05630)                                            | 246    | Homolog to B. cinerea NEP1 (Staats et al., 2007); P. Sojae NIP (Qutob et al., 2002); studied in (Bashi et al., 2011) |
| SSIG_11912 | SnNEP2                            | Necrosis inducing protein (PF05630)                                            | 245    | Homolog to B. cinerea NEP1 (Staats et al., 2007); P. Sojae NIP (Qutob et al., 2002); studied in (Bashi et al., 2011) |
| SSIG_00849 | AltA-1 allergen analog            | -                                                                              | 152    | Homolog to C. hingginsianum HE651255; CHEC91 (Kleemann et al., 2012)       |
| SSIG_02904 | CVNH protein                      | CyanolVrin-N Homology domain (PF08881)                                         | 169    | Homolog to C. hingginsianum HE651243; CHEC80 (Kleemann et al., 2012)       |
| SSIG_08858 | Deuterolysin metalloprotease      | Deuterolysin metalloprotease (M3S family (PF02102)                              | 354    | Homolog to M. oryzae AvPta (MGG_15370 - Otsch et al., 2000)               |
Sclerotinia effector candidates showing high Ka/Ks ratios

According to the red queen hypothesis, fungal pathogen effectors are predicted to evolve rapidly. Signatures of natural selection were used to identify effectors and elicitors of immune responses in bacterial, oomycete and fungal plant pathogens [29,35,50,51]. To detect signatures of natural selection in SPEP genes, we calculated Ka/Ks ratios using Yn00 [28] on alignments between *S. sclerotiorum* and *B. cinerea* 197 core ortholog pairs. We obtained Ka/Ks values ranging from 0.009 to 6, with a median value of ~0.1. Notably 5 genes (2.5%) showed Ka/Ks > 1, suggesting positive selection (Figure 3a, Table 2). To identify codon sites under positive selection in the five SPEP genes showing Ka/Ks > 1, we used a Bayesian inference approach on alignments of *S. sclerotiorum* 1980, *B. cinerea* B05.10 and *B. cinerea* t4 orthologs. We could not detect sites with Ka/Ks > 1 in SS1G_04551 and SS1G_07158 using this dataset. We detected 7, 50 and 20 sites with Ka/Ks > 1 in SS1G_07749, SS1G_10165 and SS1G_10617 mature proteins respectively. The p-value for positive selection using M8 and M8e models was 8.2e-2, 3.8e-4 and 3.0e-1 for SS1G_07749, SS1G_10165 and SS1G_10617 respectively (Table 2). Interestingly, SS1G_07749 encodes a putative xylanase that may function as a virulence factor such as xylanases from other fungi [9,52]. To get insights on the constraints shaping the evolution of SS1G_07749 xylanase, we predicted the 3D structure of SS1G_07749 mature protein and mapped the local Ka/Ks ratios on this structure (Figure 3b). All seven selected sites correspond to surface-exposed residues with three residing at the surface potentially interacting with plant xylanase inhibitors (R36, D74, Q76), three at the surface of the region corresponding to *B. cinerea* Xyn11 necrotizing peptide (N174, L189, S200) and one (L43) modifying the surface of the substrate-binding pocket. These findings are consistent with adaptation to host and the possible involvement of SS1G_07749 in virulence.

**Sclerotinia effector candidates encoded by recently duplicated genes**

Several effectors of filamentous plant pathogens evolved through gene duplication followed by rapid diversification
To identify SPEP genes that underwent duplications since the divergence of *S. sclerotiorum* with *B. cinerea*, we grouped proteins from their whole predicted proteomes based on sequence similarity using Markov clustering, and selected groups matching all three criterions: (i) containing at least 3 proteins, (ii) containing at least one *S. sclerotiorum* SPEP gene and (iii) containing more than 50% of proteins from *S. sclerotiorum*. We obtained 24 such groups including a total of 71 *S. sclerotiorum* SPEP genes. Among which 29 *S. sclerotiorum* SPEP genes contained only secreted proteins and probably carry effector-specific functions. Group 182 and 167 contained SPEP genes identified as ECs based on the presence of CFEM and LysM domains respectively (Table 1). SPEP genes from group 51, 80 and 204 had no conserved domain or homology to proteins of known function. In some pathogen genomes, the expansion of effector gene families is associated with the proximity to transposable elements [4]. To determine the repeat environment of *S. sclerotiorum* ECs, we annotated repeats in *S. sclerotiorum* genome, and calculated for all genes the distance to the nearest repeat. Median distance to repeats is 7.89 Kbp across *S. sclerotiorum* genome and 3.56 Kbp for the 29 ECs that underwent recent duplications. Nine duplicated ECs are located less than 2Kbp apart from a retrotransposon and may have undergone duplication due to transposable element activity (Figure 4b, Table 3).

Cluster051 is remarkable for containing only genes coding for secreted proteins, with only one from *B. cinerea* and 6 from *S. sclerotiorum*, such as the SPEP gene SS1G_13371 (Table 3). To get support for the duplication of SS1G_13371 gene ancestor in *S. sclerotiorum* lineage, we constructed a phylogenetic tree of SS1G_13371 homologs. A total of 58 homologs in 25 fungal species could be retrieved from the JGI database covering 238 complete fungal genomes. We selected the 20 closest homologs to build a parsimony tree based on a 90 amino-acid alignment (Figure 4c). For the 6 *S. sclerotiorum* genes of cluster 051, the phylogeny revealed clustering based on paralogy rather than orthology, suggesting that this family expanded after the separation of the 25 species analyzed. Gene duplication in this family may have allowed increased accumulation of the corresponding protein, neo-functionization in some paralogs, or differential regulation.

*S. sclerotiorum*-specific effector candidates with toxin structural analogs

The emergence of virulence is frequently associated with a high rate of mutation, gene gain and gene loss in effector genes. This evolutionary pattern results in a discontinuous taxonomic distribution for effector genes [4,55]. To identify genes showing a discontinuous taxonomic distribution among *S. sclerotiorum* SPEP genes, we looked for homologs of all 486 *S. sclerotiorum* SPEP genes in the complete genome of 13 fungal pathogens with a necrotrophic phase of infection covering all major
fungal lineages. As expected, the number of SPEP homologs identified in a given genome decreased with phylogenetic distance to *B. cinerea*, and for which the best BlastP hit is not a specific, absent in the 13 other fungal species considered (Figure 5a). Forty-six SPEP genes had homologs in all 14 fungal species monitored, and 70 were specific, absent in the 13 other fungal species considered (Figure 5b). Among those, only six (8.6%) had weak matches to PFAM domains (0.004 < e-value < 0.74) other than domains of unknown function. We could not obtain information about the function of the 64 remaining SPEP genes based on annotation or homology.

To get insights into the putative function of *S. sclerotiorum*-specific predicted SPEPs with no annotation, we performed 3D protein structure predictions followed by fold-recognition searches with an aim to identify structural analogs of known function. We focused on 17 SPEP genes encoding mature proteins of at least 50 amino-acids and for which the best BlastP hit is not a *B. cinerea* protein (Table 4). We closely examined the predicted structures of three SPEPs in comparison with their structural analogs. SS1G_09512 had analogy to the lectin domain of lectolinysin, a toxin forming pores in cell membranes regulated by sugar-binding through a lectin domain [56] (Figure 6a). SS1G_12769 had analogy to the hookworm saposin-like protein Na-SLP-1, a toxin forming pores in membranes through lipid-binding activity [57] (Figure 6b). SS1G_13235 has analogy to the C-terminal domain of death-associated protein 5 (DAP5), a member of the eukaryotic translation initiation factor eIF4G protein family (Figure 6c). Cleavage of DAP5 by caspases at its C-terminus induces apoptosis [58]. SS1G_07354 also had eIF4G protein as closest analog (Table 4). We propose that these EC are toxin analogs that may have emerged through convergent evolution and could contribute to *S. sclerotiorum* virulence.

### S. sclerotiorum effector candidate genes show diverse patterns of expression in planta

To test whether ECs identified in this work could play a role during interaction with host plants, we monitored the expression pattern after plant inoculation by quantitative RT-PCR for 16 EC genes, representative of the four lists of ECs identified through our bioinformatics analyses. For this, we inoculated four different host and model plants, including Tomato, *Nicotiana benthamiana*, *Arabidopsis thaliana* resistant accession Rubezhnoe and susceptible accession Shahdara, and harvested samples at 6, 24 and 48 hours post-inoculation (hpi). *S. sclerotiorum* mycelium grown in *vitro* was used as a reference. We included the *S. sclerotiorum* ubiquitin-conjugating enzyme gene SS1G_11173 as non-induced control gene and the cerato-platanin gene SS1G_10096 as an *in planta*-induced control [14]. Eight effector candidate genes showed in planta induction ≥2 on all four host plants tested. One gene (SS1G_07295) did not show in planta induction ≥2 fold on any plant. Using hierarchical clustering, we grouped effector candidate genes into five clusters according to their expression pattern (Figure 7a). Genes in cluster I show strong induction *in planta* over at least two time points; genes in cluster II showed strong and early

| Table 2 List of five *S. sclerotiorum* effector candidates selected based on Ka/Ks > 1 in pairwise comparisons with their *B. cinerea* orthologs |
|------------------|--------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Protein ID       | Annotation          | PFAM domains    | Length | Ka/Ks vs BC1T | Ka/Ks vs BcT4 | Ka/Ks > 1 sites | Prob            |
| SS1G_04551       | Pectinesterase      | Pectinesterase  | 308    | 2             | max Ka/Ks = 0.93 | NA              |
| SS1G_07158       | -                   | DUF1374         | 328    | 6             | max Ka/Ks = 0.4 | NA              |
| SS1G_07749       | Xylanase            | Glycoside hydrolase family 11 | 200    | 4             | R36, L43, D74, Q76, S128, N174, S200 | 0.08224 |
| SS1G_10165       | Pectinesterase      | Pectinesterase  | 310    | 2             | 0.061669 | P22, K31, T34, A36, S54, A63, S85, S86, G88, S89, Q95, A118, D146, I165, F191, D208, P211, S212, T213, L218, 226I, 230A, 233S, 236A, 237G, 238T, S246, V253, M258, S259, N260, 261 V, N263, V269, S274, P275, N276, 278Q, H285, A286, A290, H301, S302, P306, S310, N316, K318, S319, S324 | 0.00038 |
| SS1G_10617       | Glycoside hydrolase family 15 protein | Glycoside hydrolase family 15 (PF07232.16); Carbohydrate-binding module family 20; Carbohydrate-binding module family 25 (PF03423.8) | 628    | 2             | 0.078172 | N53, R56, M81, S591, N128, S313, S361, C383, R409, S416, N514, Y526, F537, V549, K586, S605, S605, S617, Q639 | 0.2987 |

Site specific Ka/Ks ratio estimated using Bayesian inference based on the M8 model of Yang et al., 2000. Only sites with Ka/Ks > 1 in the mature protein sequence are reported. P-value for positive selection estimated using a likelihood ratio test based on the comparison of twice the log likelihood difference obtained with the M8 and M8a null model with a chi-square probability table of degree of freedom 1. BC1T, *Botrytis cinerea* B.05 genome; BcT4, *Botrytis cinerea* T4 genome; ND, not determined.
induction in *N. benthamiana*, but only moderate and late induction on other host plants; genes in cluster III showed stronger induction on *A. thaliana* Shahdara accession; genes in cluster IV showed late (48 hpi) induction, stronger on *N. benthamiana* and tomato; and genes in cluster V showed moderate induction at early time points (≤24 hpi). The cerato-platanin gene *SS1G_10096* grouped in cluster IV.

The set of 16 ECs analyzed presented diverse expression patterns. We observed peaks of expression on *N. benthamiana* at 6, 24 or 48 hpi for *SS1G_08858*, *SS1G_07749* and *SS1G_00849* respectively illustrating the diversity of induction kinetics. On *A. thaliana* Shahdara accession, peaks of expression occurred at 6, 24 or 48 hpi for *SS1G_01593*, *SS1G_06213* and *SS1G_10096* respectively (Figure 7b). At 24 hpi, extensive cell death was visible on leaves of *A. thaliana* accession Shahdara whereas only limited cell death symptoms were visible on *N. benthamiana* (Additional file 2: Figure S1), suggesting that the activation of host cell death is not the only determinant of *S. sclerotiorum* EC induction. We observed a consistent 2- to 4-fold induction between 6 and 24 hpi for *SS1G_06213* on all four host plants tested. By contrast, *SS1G_08858* was induced >4-fold at 6 hpi on *N. benthamiana*, at 24 hpi on tomato, and at 48 hpi on *A. thaliana* (Figure 7c). This result suggests that *S. sclerotiorum* possess effector genes that are regulated independently of the host being colonized and others that are differentially regulated in a host-dependent manner. Furthermore, *SS1G_13371* was induced >2 fold during infection of *A. thaliana* resistant accession Rubezhnoe, but not during infection of the susceptible accession Shahdara. Conversely, *SS1G_00849* was induced >8 fold during infection of the susceptible accession Shahdara, but only ~4 fold during infection of the resistant accession Rubezhnoe (Figure 7d). This data points towards a versatile repertoire of effector candidates the expression of which can be modulated according to the nature of the host plant being colonized.

**Discussion**

In their global analysis of *S. sclerotiorum* genome, Amselem et al. [12] identified 603 genes encoding non-CAZYme, non-peptidase secreted proteins. These secretome genes did not appear significantly enriched in genes induced *in planta*. In this study, we combined multiple bioinformatics approaches to identify a total of 745 predicted secreted proteins, among which 486 with experimental evidence for expression *in planta* (SPEPs). The predicted SPEPs include SnNEP1 and SnITL1 that have proposed to be *S. sclerotiorum* virulence factors [8,44]. Since we have chosen to focus the search for effector candidates on these 486 SPEP genes, we have deliberately ignored genes expressed *in planta* for which experimental evidence is lacking, and enzymes that contribute to the biosynthesis of secondary metabolites as virulence determinants. It is therefore expected that the diversity of *S. sclerotiorum* virulence factors exceeds that of the candidate effectors presented here. Sequence similarity to known fungal effectors is a powerful method to
uncover effector families conserved across species [16], that allowed us to identify S. sclerotiorum homologs of B. cinerea NEP1 [44] and Spl1 [14], M. oryzae MGG_15370 and C. hingginsianum CHEC91 and CHEC80. To complement this approach, we used PFAM domain and NLS motif searches to reveal additional effector candidates. We identified putative chitin-binding proteins, putative protease inhibitors, cystein-rich proteins and putative nuclear localized proteins. Effectors with chitin binding activity such as C. fulvum Ecp6, M. oryzae Slp1 and

| Group   | N° Ss - Bc* | S. sclerotiorum SPEPs | Other genes in group SPEPs | SPEPs annotation          | Nearest repeat to SPEPs (distance Kb) |
|---------|-------------|-----------------------|---------------------------|--------------------------|--------------------------------------|
| Cluster012 | 7-6        | SS1G_01081            | BC1G_01968, SS1G_09509, BC1G_01095, SS1G_00547, BC1G_09386, SS1G_09141, SS1G_00588, BC1G_02407, SS1G_06186, BC1G_12856, SS1G_02784, BC1G_13021 | Catalase                  | SSrRNA_AN (41.6)                 |
| Cluster022 | 6-4        | SS1G_04468            | SS1G_04513, SS1G_09104, SS1G_09671, BC1G_12617, BC1G_02926, SS1G_09338, BC1G_00394, SS1G_14236, BC1G_00455 | Glycoside hydrolase family 47 protein | Helitron-2_PSt (47.3)            |
| Cluster035 | 5-4        | SS1G_10949            | BC1G_01945, SS1G_12508, SS1G_12999, BC1G_02687, BC1G_11888, BC1G_10788, SS1G_01984, SS1G_14293 | Glucose oxidase           | BOTY_LTR (0.8)                   |
| Cluster047 | 4-3        | SS1G_11700            | SS1G_08020, BC1G_05350, BC1G_01594, SS1G_11304, BC1G_11407, SS1G_05897 | Glycoside hydrolase family 18 protein | BOTY_LTR (1.0)                   |
| Cluster051 | 6-1        | SS1G_13371            | BC1G_04114, SS1G_12365, SS1G_03721, SS1G_14379, SS1G_11693, SS1G_10104 | -                         | Tad1-14_BG (1.1)                 |
| Cluster057 | 4-2        | SS1G_05454            | BC1G_01964, SS1G_12510, SS1G_00677, SS1G_00773, BC1G_00533 | Glycosyl hydrolases family 18 protein | BOTY_LTR (0.9)                   |
| Cluster080 | 3-2        | SS1G_05073            | BC1G_10397, SS1G_10773, BC1G_07160, SS1G_13589 | -                         | BOTY_LTR (5.8)                   |
| Cluster094 | 3-2        | SS1G_09630            | SS1G_03681, SS1G_10564, BC1G_10623, BC1G_03527 | Pyrroline-5-carboxylate reductase | BOTY_LTR (3.6)                   |
| Cluster167 | 2-1        | SS1G_13935            | BC1G_12793, SS1G_13934 | CFM domain containing    | BOTY_LTR (9.6)                   |
| Cluster169 | 2-1        | SS1G_02369            | SS1G_00501, BC1G_00594 | Xyloglucan-specific endo-beta-glucanase A | BOTY_LTR (0.5)                  |
| Cluster170 | 2-1        | SS1G_04264, SS1G_12024| BC1G_15278 | Cell wall glucanase      | BOTY_LTR (13.0) – Gypsy-31_ADe-I (35.9) |
| Cluster171 | 2-1        | SS1G_13501            | BC1G_06328, SS1G_03093 | Bacterial alpha-L-rhamnosidase domain protein | Harbinger-5_PSt (1.8) |
| Cluster175 | 2-1        | SS1G_10902, SS1G_03618| BC1G_03590 | Endo-beta-xylanase       | BOTY_LTR (3.9) - BOTY_LTR (11.4) |
| Cluster176 | 2-1        | SS1G_04098, SS1G_09225| BC1G_01026 | Tripeptidyl peptidase sed3 | BOTY_LTR (1.1) - Mariner-3_AF (5.9) |
| Cluster181 | 2-1        | SS1G_07498, SS1G_01811| BC1G_06353 | Glucose-methanol-choline oxidoreductase | BOTY_LTR (18.2) - BOTY_LTR (0.6) |
| Cluster182 | 3-1        | SS1G_12509            | SS1G_00772, SS1G_05453 | LysM domain protein       | Mariner-3_AF (8.1)               |
| Cluster186 | 2-1        | SS1G_04200            | SS1G_02425, SS1G_01334 | Alpha-mannosidase family protein | BOTY_LTR (2.2) |
| Cluster187 | 2-1        | SS1G_04207            | BC1G_00240, SS1G_04205 | Polygalacturonase         | BOTY_LTR (3.4) |
| Cluster196 | 2-1        | SS1G_05273            | SS1G_11068, BC1G_09997 | Amidase family protein    | BOTY_LTR (28.8) |
| Cluster203 | 2-1        | SS1G_03160            | SS1G_00233, BC1G_07555 | Autophagy related lipase  | BOTY_LTR (1.2) |
| Cluster204 | 2-1        | SS1G_08110, SS1G_12361| BC1G_12374 | -                       | SSrNA_AN (2.3) – Tad1-14_BG (1.8) |
| Cluster206 | 2-1        | SS1G_08104            | SS1G_09795, BC1G_12379 | Acetyl xylan esterase     | SSrNA_AN (6.2) |
| Cluster214 | 2-1        | SS1G_00446            | BC1G_00660, SS1G_05782 | Heterokaryon incompatibility Het-c domain protein | SINE3-2_AO (5.4) |
| Cluster220 | 2-1        | SS1G_05461            | BC1G_06146, SS1G_13907 | Thioesterase-like domain protein | TGATGAA)n (0.3) |

Repbase nomenclature for transposons used here: BOTY_LTR, Botrytis cinerea gypsy-type retrotransposon; Gypsy-31_ADe-I, internal portion of retrotransposon GYPSY31; Harbinger-5_PSt, Harbinger-type DNA transposon; Mariner-3_AF, Mariner DNA transposons; SINE3-2_AO, SINE3 nonautonomous non-LTR retrotransposon; Tad1-14_BG, Tad1 Non-LTR retrotransposon from barley powdery mildew. Retrotransposons are indicated in bold. * Number of genes from S. sclerotiorum - B. cinerea.

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http://www.biomedcentral.com/1471-2164/15/336
M. graminicola Mg3LysM function in suppressing plant immunity [18,20,59]. Similarly, SsITL1 (SS1G_14133) integrin-like secreted protein suppresses plant jasmonic acid and ethylene signaling pathways and enhances susceptibility [8]. These findings suggest that *S. sclerotiorum* secretes proteins able to suppress plant immunity.

The comparative analysis of *Fusarium graminearum* secretome and genomes of other ascomycetes revealed a high level of conservation with only 25 *F. graminearum* specific out of 574 secreted proteins [60]. The taxonomic distribution of *S. sclerotiorum* SPEP homologues analyzed in this work supports the conservation of more than 50% of SPEP genes across ascomycetes. As proposed by Brown et al. [60], these core SPEP genes may support *S. sclerotiorum* epiphytic growth and highlight important distinctions between multiple phases in infection by this fungus [61]. Nevertheless it also revealed 70 SPEP genes (14%) specific to *S. sclerotiorum*, many of which are unannotated proteins. The systematic prediction of their 3D structure allowed identifying putative structural analogs of some predicted SPEPs and suggests that they may carry unique functions to assist *S. sclerotiorum* pathogenicity. It will be interesting to take advantage of these predictions to test the biological function of these effector candidates and confront them to experimentally determined structures. Furthermore, in spite of the limited sequence diversity included in the dataset analyzed here, we were able to detect signatures of positive selection in five *S. sclerotiorum* SPEP genes (2.5% of genes analyzed). Similar frequency (3.2%, 21 out of 642 genes) has been reported in *B. cinerea* [35]. In the future, an in depth exploration of sequence diversity in *S. sclerotiorum* should allow to reveal more sites subjected to selection. SPEP genes for which positive selection has been detected encode cell wall degrading enzymes, including SS1G_07749 encoding a putative xylanase. This protein is related to *B. cinerea* Xyn11 considered as a Pathogen Associated Molecular Pattern (PAMP) [9]. The detection of positive selection in SS1G_07749 is therefore consistent with the hypothesis that PAMPs may be characterized by signatures of positive selection in a background of strong negative selection [50]. It may therefore be hypothesized that a subset of *S. sclerotiorum* critical secreted enzymes are engaged in an evolutionary arms race with plant pattern recognition receptors, driving opposing forces of natural selection *S. sclerotiorum* effector genes.

Since plant inhibitors are known for many fungal cell wall degrading enzymes, it is also possible that an evolutionary arms race with plant inhibitors drives the evolution of some *S. sclerotiorum* effector candidates [53,62,63]. Remarkably, we also identified 14.4% of species-specific SPEP genes. The extent to which evolutionary constraints imposed by a broad host range contributes to diversification

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**Figure 5 Taxonomic distribution of *S. sclerotiorum* SPEP genes across 14 fully sequenced fungal pathogen genomes.**

(a) Bar chart showing the number of *S. sclerotiorum* SPEP genes conserved along a phylogeny of fungal pathogens. Conservation was determined based on BlastP searches as described in the methods.

(b) Distribution of *S. sclerotiorum* SPEP genes according to the number of species in which they are not conserved. *S. sclerotiorum* SPEP genes conserved in a given species but not in *B. cinerea* are shown in red.
in the effector candidate repertoire of *S. sclerotiorum* remains to be determined. Detailed functional analysis of effector gene alleles and their plant targets will be needed to address this question.

Although *S. sclerotiorum* is considered as a typical necrotroph, there is evidence that it colonizes plant tissues through multiple phases involving important transcriptional and physiological reprogramming [61]. Consistent with this model, the phytotoxin oxalic acid dampens plant immune responses at the initial stages of infection and later enhances programmed cell death [7].

In this work, we report the sequential transcriptional activation of *S. sclerotiorum* candidate effector genes. A >2 fold induction was measured at 6 hpi for several effector candidate genes, whereas no necrotic symptoms are visible at this time, except on *A. thaliana* Shahdara accession. This suggests that the sequential secretion of effectors is required for the efficient induction of host cell death by *S. sclerotiorum*, or that some secreted proteins could contribute to *S. sclerotiorum* virulence independently of host cell death activation. By comparing the expression pattern of selected *Blumeria graminis* f. sp. *hordei* genes grown on barley and on *A. thaliana*, Hacquard et al. [64] concluded that very divergent hosts do not significantly alter the fungal gene expression program. The expression pattern of some *S. sclerotiorum* ECs is indeed independent on the host plant being colonized (e.g. SS1G_06213). Nevertheless, other ECs showed differential regulation in a host-dependent manner (e.g. SS1G_08858, SS1G_13371). The transcriptional activation of distinct set of effectors depending on the host being colonized has also been reported for the generalist root endophyte *Piriformospora indica* in barley and *A. thaliana* [65]. The growing number of transcriptomic studies on various pathosystems should help determine whether host-dependent modulation of effector gene expression differs according to pathogens lifestyle or host range. We speculate that the white mold fungus benefits from a versatile repertoire of secreted proteins with diverse functions, evolution and expression patterns, to successfully infect a wide range of host plants. A systematic characterization of *S. sclerotiorum* transcriptome on multiple hosts and the functional analysis of differentially regulated effector genes should prove useful to decipher the molecular determinants of quantitative disease resistance and host range in this fungal pathogen.

| Length | C-score | TM score | Analog model | Selected analog description |
|--------|---------|----------|--------------|----------------------------|
| SS1G_00780 | 115 | -3.14 | 0.610 | 2PFV |
| SS1G_01817 | 116 | -2.53 | 0.503 | 2RSM |
| SS1G_03830 | 155 | -4.41 | 0.460 | 3GH1 |
| SS1G_03878 | 132 | -3.79 | 0.645 | 1NI3 |
| SS1G_04309 | 177 | -4.63 | 0.461 | 2QPD |
| SS1G_06504 | 93 | -2.38 | 0.503 | 4HO1 |
| SS1G_07543 | 273 | -4.27 | 0.706 | 2FXS |
| SS1G_07543 | 273 | -4.27 | 0.706 | 2FXS |
| SS1G_07650 | 141 | -3.96 | 0.526 | 1XYL |
| SS1G_09512 | 164 | -3.65 | 0.546 | 3LE0 |
| SS1G_10915 | 136 | -4.12 | 0.512 | 3M1C |
| SS1G_11461 | 114 | -3.2 | 0.526 | 2HP3 |
| SS1G_12769 | 88 | -2.32 | 0.585 | 3S63 |
| SS1G_13016 | 124 | -2.76 | 0.555 | 3APO |
| SS1G_13142 | 131 | -4.25 | 0.506 | 1DCU |
| SS1G_13235 | 114 | -3.55 | 0.585 | 3D3M |
| SS1G_14000 | 159 | -4.43 | 0.578 | 2C9K |

C-score is a confidence score for estimating the quality of models predicted by I-TASSER ranging from -5 (low confidence) to 2 (high confidence). TM score is a quality score for the superimposition of 3D models calculated with TM-align with values in [0,1]. A TM-score >0.5 generally corresponds to the same fold in SCOP/CATH.
Conclusions
In this work, we explored systematically the diversity of candidate virulence genes in the necrotrophic fungal pathogen *S. sclerotiorum* using *in silico* structure and evolution analyses. We report the identification of 486 *S. sclerotiorum* secreted proteins expressed *in planta*, including 78 ECs. We have analyzed *in planta* expression for a representative subset of 16 ECs, highlighting diverse predicted functions and expression patterns. This study reveals that besides plant degrading enzymes, *S. sclerotiorum* genome encodes numerous predicted secreted proteins that may be involved in the interaction between the fungus and its host plants. It will facilitate future investigation on their relevance in the infection process and sheds new light on the underestimated complexity of host colonization by necrotrophic plant pathogens.

Methods

Secretome prediction and annotation
We used complete genome and predicted proteomes of *Sclerotinia sclerotiorum* strain 1980 v.2, *Botrytis cinerea* strain b05.10 v.1 and strain t4 v.1 described in [12]. The presence of secretion signals was predicted with SignalP v.2 and v.4 [66,67], transmembrane helices and GPI anchor sequence were predicted with TMHMM [68] and GPIsom [69] respectively. For the identification of genes expressed *in planta*, microarrays data for gene induction fold at 2 days post inoculation on sunflower cotyledons and Expressed Sequence Tags (ESTs) from [12] were used. ESTs were assigned to the *S. sclerotiorum* predicted transcript giving the lowest e-value in a BLASTN search. Genes were considered expressed *in planta* when either (i) showing induction fold \( \geq 1 \) during sunflower infection in microarrays data or (ii) being assigned at least one EST in either infection cushion, infected *B. napus* or infected tomato library. *S. sclerotiorum* predicted proteins were annotated using Blast2GO [70], PFAM [71] and NLStradamus [72]. Predicted proteins shorter than 40 amino-acids were excluded from the analysis. PFAM domains were annotated using HMMER3 searches against the PFAM 26.0 database [71]. We defined non-annotated predicted SPEPs as having no hit to PFAM_A with e-value <0.1. For the identification proteins similar to known fungal effectors, BlastP searches against a local database of 191 effectors with an e-value cutoff of \( 1 \times 10^{-3} \).

Definition of ortholog clusters and natural selection analysis
Core ortholog groups (COGs) between *S. sclerotiorum* 1980 and *B. cinerea* b05.10 or *B. cinerea* t4 proteomes were identified using Inparanoid7 [73] with the following parameters: score cutoff 40 bits; sequence overlap cutoff 0.5; group merging cutoff 0.5; scoring matrix BLOSUM62. COGs in which a length difference >10 amino-acids existed between *S. sclerotiorum* and *B. cinerea* were discarded. Pairwise ortholog alignments were generated using the needleall program from the EMBOSS package using the following parameters: gapopen 50.0; gapextend 0.2; minscore 100.0; aformat3 MARKX3. Needleall output files were parsed into .axt alignments used as input in Ka/Ks calculator2 [74]. Ka/Ks ratios were
Figure 7 (See legend on next page.)
calculated for all COG pairs using Yn00 method [28]. The identification of codon sites under positive selection was achieved through Bayesian inference using the Selecton2.2 server [75] with the “Positive selection enabled (M8, beta + w > =1)” evolutionary model with 8 categories, on alignments of S. sclerotiorum 1980, B. cinerea b05.10 and B. cinerea t4 orthologs.

Protein structure modeling and analysis
Protein structure modeling was performed with the I-TASSER server [76] and rendered using UCSF Chimera [77]. Site-specific alignment consensus and Ka/Ks ratios were mapped onto protein models using the ‘define attribute’ function in UCSF Chimera. Moving average over a 3 amino-acid window of the percentage consensus in a 99 homologous protein alignment was used to characterize conservation in SS1G_01593 family. Structural analogs were identified using the TM-align program in I-TASSER.

Taxonomic distribution and phylogenetic analyses
Fungal taxonomy trees are based on [16]. The presence of SPEP homologs in 234 fungal species was assessed using BlastP searches against JGI fungi Gene Catalog Proteins [78] with an e-value cutoff of 1e-5 without low complexity filter. Among retrieved homologs, proteins that had no signal peptide detected with SignalP4 or SignalP2 were discarded. For the global analysis of taxonomic distribution of SPEP genes, the predicted proteomes of Neurospora crassa, Magnaporthe oryzae, Verticillium dahlia, Fusarium oxysporum, Stagonospora nodorum, Pyrenophora triticum-repens, Alternaria brassicicola, Leptosphaeria maculans, Mycosphaerella graminicola, Aspergillus flavus, Cryptococcus neoformans and Rhizopus oryzae were used in local BlastP searches with e-value cutoff 1. For each SPEP gene, BlastP scores for all hits in a given species were summed up, and SPEP genes were considered as absent if total score is <2. Phylogenetic trees for SS1G_13371 family was generated using the parsimony method with 100 bootstrap replicates with the Extended Majority rule, as implemented in the protpars and consense programs of the Phylip 3.67 package [79].

Sequence-based clustering and genome distribution of duplicated genes
S. sclerotiorum 1980 and B. cinerea b05.10 proteins were clustered based on sequence similarity by Markov clustering using the orthoMCL function in Biolayout 3D [80]. A self BlastP search on the combined S. sclerotiorum 1980 and B. cinerea b05.10 complete proteomes with e-value cutoff 1e-30 was used as input for orthoMCL. Repeats and transposable elements were identified using RepeatMasker on S. sclerotiorum 1980 supercontigs with the cross_match method at slow speed and “Fungi” as a DNA source. Genomic distances and genome architecture heatmaps were generated according to [81].

Plant and fungus cultivation, inoculation procedure
Arabidopsis thaliana accession Shahdara and Rubezhnoe-1 were grown in Jiffy pots for four weeks at 22°C with cycles of 9 hours of light per 24 hours. Tomato (Solanum lycopersicum cv. Heinz) were grown for six weeks in pots containing disinfected soil in a greenhouse at 23°C with cycles of ~14 hours of light per 24 hours. Nicotiana benthamiana plants were grown for four weeks at 21°C with cycles of 16 hours of light per 24 hours. S. sclerotiorum strain S55 was first grown for 4 days on PDA plates at 25°C in the dark. Fifty mL of liquid PDB medium were inoculated with 3 agar plugs of PDA cultures and incubated for 4 days at 25°C in the dark, with 150 rpm shaking. Three independent inoculation experiments were performed in which fully grown plant leaves were cut and placed right side up on a wet paper towel in large petri dishes. Mycelium was washed twice in PDB, filtered on Miracloth (Calbiochem, CA), and spread over whole leaf surfaces. Inoculated leaves were incubated for up to 3 days at 25°C with 14 hours of light per 24 hours.

Effectors candidate gene expression by quantitative RT-PCR
Plant leaves were harvested immediately and 6, 24 and 48 hours after inoculation, and ground in liquid nitrogen. Total RNA was extracted using a Nucleospin RNA II kit (Machery- Nagel) according to manufacturer's instructions. RNAs were analyzed and quantified on an Agilent 2100 Bioanalyzer. The first-strand cDNA was synthesized using TRT reverse transcriptase (Roche) according to

(See figure on previous page.)

**Figure 7** In planta expression analysis for selected S. sclerotiorum effector candidates on four different hosts. (a) Transcriptional profiles of 16 S. sclerotiorum effector candidate genes. Overrepresented (yellow) and underrepresented transcripts (blue) in planta are shown as log2-fold changes relative to expression in vitro, normalized using Actin expression. Hierarchical clustering based on Pearson correlation coefficients delimited five clusters. The SS1G_11173 ubiquitin 16 gene was used as a non-induced control (b) Sequential transcriptional activation of effector gene candidates during the infection of N. benthamiana (left) and A. thaliana Sha. accession (right). (c) in planta expression pattern of candidate effector genes showing host-independent expression (SS1G_06213, left) and host-dependent expression (SS1G_08685, right). (d) Differential expression patterns of two candidate effector genes on susceptible (Sha) and resistant (Rub) A. thaliana accessions. Relative gene expression shown as log2-fold changes relative to expression in vitro, normalized using Actin expression. Error bars show standard deviation calculated from two technical replicates on each of three independent biological experiments. Rub., Rubezhnoe; Sha., Shahdara.
manufacturer’s instructions. Real-time PCR reactions included 3.5 μL of SYBR green mix (Roche), 1 μL of 5 μM primers (Additional file 3: Table S2) and 200 ng of cDNA. Reactions were performed on a Light Cycler 480 II machine (Roche) under the following conditions: 95°C for 5 minutes; 45 cycles of 95°C for 15 seconds, 65°C for 20 seconds and 72°C for 20 seconds; then 95°C for 10 seconds; 65°C for 15 seconds followed by a progressive increase in temperature at 0.11°C/second up to 95°C to obtain melt curve. *S. sclerotiorum* actin (SSIG_O8733) and ubiquitin 16 (SSIG_11173) genes were used as controls. The expression of effector gene candidates relative to Ct values of the control genes was determined and analyzed using the LightCycler 480 SW 1.5 software. Fungal cultures were grown *in vitro* for 3 days and either harvested immediately (Day 0) or inoculated to plants. Values are given as log2 ratio over Day 0 expression. Error bars represent standard deviation calculated from two technical replicates on each of three independent biological experiments.

Additional files

Additional file 1: Table S1. Summary table of annotations for the 486 SPEP genes, including homology, domain and motif searches and expression data. Inclusion into sequence-based clusters, into COG groups, KA/Ks values and closest structural analogs are indicated where relevant.

Additional file 2: Figure S1. Representative symptoms of detached leaves inoculated with *S. sclerotiorum* S55 at 6, 24 and 48 hours post inoculation (hpi).

Additional file 3: Table S2. List of primers used in this work.

Competing interest

The authors declare that they have no competing interests.

Authors’ contributions

KG and CB performed experiments and analyzed transcriptomic data. KG and SR performed secratome analysis. SR designed the research. DR and SR wrote the manuscript, all authors commented on the manuscript before submission. All authors read and approved the final manuscript.

Acknowledgements

SR is supported by a Marie Curie CIG grant ("SEPARATE", contract 334036) and a starting grant of the European Research Council ("VariWhim", contract 336808). This work was supported by the French Laboratory of Excellence project "TUUL" (ANR-10-LABX-41; ANR-11-IDEX-0002-02).

Received: 3 February 2014 Accepted: 27 April 2014

Published: 4 May 2014

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