Kingdom-Wide Analysis of Fungal Small Secreted Proteins (SSPs) Reveals their Potential Role in Host Association

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Fungal secretome consists of various functional groups of proteins, many of which participate in nutrient acquisition, self-protection, or manipulation of the environment and neighboring organisms. The least characterized component of the secretome is small secreted proteins (SSPs). Some SSPs have been reported to function as effectors, but most remain to be characterized. The composition of major secretome components, such as carbohydrate-active enzymes, proteases, lipases, and oxidoreductases, appear to reflect the lifestyle and ecological niche of individual species. We hypothesize that many SSPs participate in manipulating plants as effectors. Obligate biotrophs likely encode more and diverse effector-like SSPs to suppress host defense compared to necrotrophs, which generally use cell wall degrading enzymes and phytotoxins to kill hosts. Because different secretome prediction workflows have been used in different studies, available secretome data are difficult to integrate for comprehensive comparative studies to test this hypothesis. In this study, SSPs encoded by 136 fungal species were identified from data archived in Fungal Secretome Database (FSD) via a refined secretome workflow. Subsequently, compositions of SSPs and other secretome components were compared in light of taxa and lifestyles. Those species that are intimately associated with host cells, such as biotrophs and symbionts, usually have higher proportion of species-specific SSPs (SSSPs) than hemibiotrophs and necrotrophs, but the latter groups displayed higher proportions of secreted enzymes. Results from our study established a foundation for functional studies on SSPs and will also help understand genomic changes potentially underpinning different fungal lifestyles.

Keywords: fungi, lifestyle, secretome, small secreted proteins, effectors

Abbreviations: SSPs, small secreted proteins; SSSPs, species-specific SSPs; CSSPs, conserved SSPs.
INTRODUCTION

Diverse groups of pathogenic fungi threaten plant health, whereas certain fungi, such as endophytes and mycorrhizal fungi, allow plants to explore new niches, manage biotic and abiotic stresses better, and/or efficiently acquire key nutrients. In both types of plant-fungus interactions, the outcome of interaction is influenced heavily by fungal secretomes, various proteins secreted or injected to plants (Girard et al., 2013). In the secretome, certain small secreted proteins (SSPs) are known to be responsible for disease development as virulence factors or cause resistance (R)-gene mediated defense as avirulence factors (Rep, 2005; Deller et al., 2011; Hacquard et al., 2012). Such SSPs are termed effector proteins and modulate key defense signaling pathways and downstream responses, to attenuate microbe-associated molecular pattern (MAMP) triggered immunity (MTI; Jones and Dangl, 2006). Plants have evolved to activate effector-triggered immunity (ETI) by sensing specific effectors or molecular changes caused by effectors, mainly using the nucleotide binding, leucine-rich repeat class of R-gene products (Dodds and Rathjen, 2010).

Initially, effectors were considered as virulence factors secreted by pathogens (van Esse et al., 2007; Stergiopoulos and de Wit, 2009; Lo Presti et al., 2015). However, it has become apparent that effector-mediated manipulation of MTI is required even for symbiotic associations, because microbial partners also display MAMPs (Zamioudis and Pieterse, 2012; Gourion et al., 2015). Many SSPs have been identified as putative effectors in beneficial plant-associated bacteria (Soto et al., 2006) and mutualistic fungi, such as Glomus intraradices (Kloppholz et al., 2011) and Laccaria bicolor (Plett et al., 2011), expanding the definition of effectors as secreted microbial products that facilitate the establishment of various plant-microbe associations ranging from beneficial to detrimental. Furthermore, SSPs that resemble effector proteins of pathogenic fungi have been identified in saprotrophic fungi, suggesting additional roles of SSPs (Rovenich et al., 2014; Seidl et al., 2015). Driven by the discovery of diverse putative effector proteins in fungi representing different lifestyles, several studies have analyzed the repertoire of putative secreted proteins encoded by various fungi and the potential relationship between their secretomes and lifestyles (Lowe and Howlett, 2012; Krijger et al., 2014; Meinken et al., 2014; Lo Presti et al., 2015). Analysis of the size of secretome relative to the total proteome in 48 fungal species by Lowe and Howlett (2012) suggested its potential relationship with lifestyles. Another comparative study by Meinken et al. (2014) proposed that the secretome prediction of previous study may be overestimated because only SignalP was used for the prediction, but they drew the same conclusion. However, these studies did not consider individual components of the secretome. The study by Krijger et al. (2014) suggested that phylogenetic position strongly influenced both the secretome size and its composition by analyzing 33 fungal species but did not include major secreted enzyme groups. In addition those displaying different modes of pathogenesis (biotroph, hemibiotroph, and necrotroph) were combined as a single lifestyle in the last two analyses. Lastly, the review on fungal effector proteins by Lo Presti et al. (2015) only considered plant cell wall degrading enzymes in order to mine putative effector proteins. The secretome contains not only effector proteins but also groups of enzymes involved in the breakdown of cell walls, self-protection or nutrient acquisition, such as carbohydrate-active enzymes (CAZymes), oxidoreductases, proteases, and lipases (Girard et al., 2013). Not surprisingly, biotrophs encode fewer CAZymes than hemibiotrophs and necrotrophs (Zhao et al., 2014). To investigate whether the composition and size of putative effectors correlates with different lifestyles, such enzymes should also be analyzed separately.

A wide range of validated and suspected protein effectors encoded by bacteria and oomyctes have been identified, which was facilitated by the conserved delivery machinery to plant cells (Cornelis and Van Gijsegem, 2000) and sequence motifs present in effectors (Whisson et al., 2007), respectively. Although, a conserved IGY motif has been identified in a novel SSP family of Dikarya fungi (Cheng et al., 2014), known fungal effectors do not show conserved features, hampering their identification (Rafiqi et al., 2012; Giraldo and Valent, 2013). One or more of the following features have been used to predict candidate effector proteins in fungal secretomes: (a) presence of the signal peptide, but no transmembrane domain or GPI-anchor sites; (b) small sized proteins (usually fewer than 300 amino acids) that are present only in specific species or isolates; (c) expression in planta or during infection; (d) rich in cysteine residues; and (e) presence of a conserved motif within effector candidates like in oomycete and fungal effectors (Birch et al., 2008; Godfrey et al., 2010; Zuccaro et al., 2011; Cheng et al., 2014). Using these features candidate effector proteins have been identified in three types of plant pathogenic fungi: biotrophs such as rusts (Duplessis et al., 2011), smuts (Schirawski et al., 2010), and Blumeria graminis (Spanu et al., 2010), hemibiotrophs including Verticillium dahliae (Santhanam and Thomma, 2013) and Magnaporthe oryzae (Kim et al., 2010), and necrotrophs including Fusarium graminearum (Brown et al., 2012) and Sclerotinia sclerotiorum (Guyon et al., 2014). However, direct comparisons between these fungi are hampered because different bioinformatics approaches and criteria have been used for the prediction of effector proteins. The lack of robust pipelines that can be applied to mine candidate effector proteins from rapidly increasing genome sequences of phylogenetically diverse fungi and limited in planta expression data for the genes encoding SSPs also have hampered large-scale comparative analyses of putative effectors.

In this study, we refined multiple secretome components with the focus on SSPs for 136 fungal species archived in Fungal Secretome Database (FSD; Choi et al., 2010) via a data extraction pipeline consisting of multiple programs. This refined data set was analyzed in the context of the phylogenetic position and lifestyle of individual species. Secreted enzymes that likely play important roles in colonizing host plants, including CAZymes, oxidoreductases, which are likely secreted for protection against host-produced reactive oxygen species (Chi et al., 2009), and lipases and proteases, which participate in nutrient acquisition and manipulation of host defense, were also compared. Resulting data helped determine which secretome components might function as major lifestyle determinants. In addition, we mined...
candidate effector proteins for functional validation and also showed the pattern of evolutionary changes associated with several known effector proteins.

**MATERIALS AND METHODS**

**Phylogenetic Analysis**

The phylogenetic trees of 136 fungal species shown in Supplementary Figure S2 was constructed using CVTree v4.2.1 with k-tuple 7 (Xu and Hao, 2009). The tree only shows topology and ectopically positioned *Taphrina deformans* was manually curated based on NCBI Taxonomy. The lifestyle of each fungus was annotated based on literature review.

**Secretome Data Collection, Refinement, and Annotation**

The SP, SP$^2$ and SL classes of secretory proteins, which include proteins carrying a classical signal peptide, were downloaded from FSD (Choi et al., 2010). Hence, only the proteins secreted via canonical pathway were considered in this analysis. In order to predict SSPs, a two-step mining pipeline was employed. The first step, adopted from Brown et al. (2012), involves refining the secretome by selecting proteins predicted to be secreted by both ProtComp v9.0 (detected as secreted) and WoLF PSORT v0.2 (extr => 10; Horton et al., 2007), which are protein localization prediction programs trained with fungal data. Proteins that may be secreted but probably membrane bound were filtered out using Phobius v1.01 (TM = 1; SP = N; Käll et al., 2004), a program that detects signal peptides and transmembrane helixes, and UTProt (GPI-anchored = Y), a fungal specific GPI-anchor prediction tool (Cao et al., 2009). These programs were run on local Linux computers and the parameter settings were determined with various fungal effector proteins listed in the review by Stergiopoulos and de Wit (2009). The second step was grouping proteins within individual refined secretomes based on their predicted functions. To identify CAZymes, relevant HMM profiles from dbCAN release 3.0 (Yin et al., 2012) were employed. Oxidoreductases, lipases, and proteases were identified using BLASTP (E-value cutoff of 0.001) with individual refined secretomes as queries against BLAST databases of these enzyme sets. Fungal oxidoreductases were downloaded from Fungal Peroxidase Database (Choi et al., 2014). Sources for the lipase and protease datasets were the Lipase Engineering Database (Fischer and Pleiss, 2003) and MEROPS (Rawlings et al., 2014), respectively.

**Mining and Annotation of SSPs**

Protein length and cysteine content were analyzed using in-house Python scripts, and putative species-specific proteins were identified by running BLASTP against all other species, followed by BLASTP against NR database excluding itself to reduce false positives caused by the limited phylogenetic coverage of certain taxa. Both species-specific SSPs (SSSPs) and conserved SSPs (CSSPs) were annotated using pre-computed InterPro terms, which were retrieved from Comparative Fungal Genomics Platform 2.0 (CFGP 2.0; Choi et al., 2013), and mapped to PHI-base effector proteins using BLASTP (Urban et al., 2015).

**Clustering of CSSPs and Molecular Evolutionary Analysis**

All by all BLASTP analyses were performed with CSSPs and PHI-base effector proteins prior to Markov Cluster Algorithm (MCL) clustering. Resulting data were clustered via MCL with the inflation option 1.4 for high granularity (Enright, 2002), which produced the least number of singletons. The predicted protein families containing PHI-base effectors were analyzed further. Proteome data from CFGP 2.0 were used to construct species trees, and proteins in each of the analyzed families were used to construct gene trees. CVtree v4.2.1 was used for fungal species tree with k-tuple 7 (Xu and Hao, 2009), ClustalW in MEGA 6.06 was used for alignment of proteins, and maximum-likelihood gene trees were constructed using the default setting (Hall, 2013). After reconciling the species and gene trees using Notung 2.6 (Chen et al., 2000), potential gene duplication and loss events were annotated.

**RESULTS**

**Analyzed Species Cover Diverse Taxa and Lifestyles**

Genome sequences of the 136 fungal species used in this study (Supplementary Table S1) are publicly available. The taxa covered include Microsporidia, Zygomycota, Glomeromycota, Ascomycota, and Basidiomycota (Table 1). The lifestyles represented include animal pathogens, biotrophic, hemibiotrophic and necrotrophic plant pathogen, symbionts, and saprotrophs (Table 1). The necrotrophs were further divided into crop-infecting and wood-decaying types. The symbionts are fungi associated with plants and resulting beneficial/mutualistic effects in the interactions. These include both ecto- and arbuscular mycorrhizal fungi, endophytes, and a plant growth promoting fungus with symbiotic activity (Vargas et al., 2009).

| Phylum                  | Subphylum (Division/Class) | Number of species | Lifestyle             | Number of species |
|------------------------|----------------------------|-------------------|-----------------------|-------------------|
| Ascomycota             | Pezizomycotina             | 72                | Animal pathogen       | 27                |
| Saccharomycotina       | 10                         |                   |                       |                   |
| Taphrinomycotina       | 4                          |                   | Biotroph              | 10                |
| Basidiomycota          | Agaricomycotina            | 34                | Hemibiotroph          | 9                 |
| Pucciniomycotina       | 4                          |                   | C-necrotroph          | 23                |
| Ustilaginomycotina     | 5                          |                   | W-necrotroph          | 26                |
| Glomeromycotina        | Glomeromycetes             | 1                 | Symbiont              | 6                 |
| Microspora             | Microsporidia              | 5                 |                       |                   |
| Zygomycota             | Mucoromycotina             | 1                 | Saprotroph            | 35                |
| Total                  | 136                        |                   |                       | 136               |

C-necrotroph, Crop-infecting necrotroph; W-necrotroph, Wood-decaying necrotroph.

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**TABLE 1 | Taxonomic distribution and lifestyle of the species analyzed in this study.**

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Refined Fungal Secretomes Show High Degree of Size Variance

Since, fungal effectors are expected to be secreted into the host apoplastic space or cytoplasm, mining fungal secretomes is the first step for their identification. Secretome data for most sequenced fungi have already been archived in FSD (Choi et al., 2010), an online platform that was built to identify and archive secreted proteins via six different programs detecting extracellular proteins and a trans-membrane helix detection program. The predicted secretomes were then categorized into three different classes. To predict bona fide SSPs, the total secretome data from FSD were further refined via additional filtrations (Figure 1). Putatively membrane-bound proteins were eliminated using additional programs that detect transmembrane-helixes and GPI-anchors (see Materials and Methods).

After this refinement, the size of secretome was reduced by 33.7% on average compared to that predicted using only SignalP 3.0 and by 70.9% compared to the total secretome predicted by the pipeline used for building FSD (Supplementary Figure S1). The number of proteins in refined secretomes ranged from 19 (Pneumocystis jirovecii, an opportunistic human pathogen) to 1940 (Auricularia subglabra, a wood-decaying necrotroph; Supplementary Table S1). The refined secretome accounted for 5.5% of the total proteome on average, with the lowest being 0.5% (P. jirovecii) and the highest being 11.0% (Magnaporthe oryzae, a hemibiotrophic plant pathogen). Both P. jirovecii and M. oryzae belong to the phylum Ascomycota, illustrating high degrees of variance within individual phyla (Supplementary Figure S1).

Patterns Observed Among Refined Secretomes in the Context of Phylogenetic Positions and Lifestyles

Several general patterns associated with the size of refined secretome were observed (Figures 2A,B). On average, fungi belong to Pucciniomycotina encode the largest refined secretomes, whereas Microsporidia code for the smallest ones (Figure 2C). However, the size of refined secretome also varied widely within individual taxa as illustrated in Supplementary Figure S2. For example, in Ascomycota, the species belong to Pezizomycotina have much larger secretomes than those of Saccharomycotina and Taphrinomycotina, and in Basidiomycotia, the species belong to Pucciniomycotina have larger secretomes than those of Ustilaginomycotina.

We analyzed the composition of refined secretomes among groups that represent different lifestyles to investigate their relationship. On average, the size of refined secretome for plant-associated species was larger than those of saprotrophs and animal pathogens (Figure 3A). Among the plant-associated species, pathogens encode larger refined secretomes than symbionts. Among the pathogens, crop-infecting necrotrophs code for the largest refined secretome, followed by hemibiotrophs, wood-decaying necrotrophs, and biotrophs. Proportions of CAZymes, proteases, lipases, and oxidoreductases in 136 species were analyzed in relation to their lifestyles (Figure 3B; Supplementary Table S1). Although these enzymes facilitate nutrient acquisition and defense against reactive oxygen species from host (Rogers et al., 1994; Sreedhar et al., 1999; Chi et al., 2009; Blümke et al., 2014), they were not considered for mining effector-like SSPs as in Rep (2005). On average, 46.6% of the refined secretome corresponded to these enzymes. Biotrophs display the lowest proportion for all four types of enzymes, and the highest proportion for all types, except oxidoreductases, was observed in crop-infecting necrotrophs. Wood-decaying necrotrophs exhibit the highest proportion of oxidoreductases. Animal pathogens have higher proportion of proteases than saprotrophs, symbionts, and plant pathogens except necrotrophs, suggesting the importance of proteases in animal pathogenesis and necrosis of plants.
Patterns Associated with Effector-Like SSPs

Most effector-like SSPs belong to the proteins of other functions. Accordingly, we first removed the four groups of enzymes from the refined secretomes before identifying SSPs (Figure 1). Subsequently, three features, including short length (<300 aa), species-specific distribution pattern, and cysteine enrichment, were used to identify effector-like SSPs in the proteins of other functions.

Proteins shorter than 300 aa are abundant in all species (Figure 4A) with the exception of four saprophytic fungi, including *Schizosaccharomyces pombe*, *Pichia pastoris*, *Spathaspora passalidarum*, and *Ophiostoma piceae* (Supplementary Figure S3). In general, biotrophs have the most abundant SSPs. On other hands, proportions of species-specific proteins in the proteins of other functions varied between individual species (Supplementary Figure S4). In general, species that are intimately associated with living plant tissues, such as biotrophs and symbionts, have greater numbers of species-specific proteins than those with different lifestyles (Figure 4B). This suggests that species-specific presence of effectors probably arose via co-evolution with the hosts (Stergiopoulos et al., 2012). The species-specific proteins accounted for 25–50% of the proteins of other functions in 27 species and for over 50% in six species (Table 2). Many wood-decaying necrotrophs have large proportions (but not exceeding 50%) of species-specific secreted proteins. Proportions among symbionts typically ranged from 25 to 50%, but it is over 50% in *Rhizophagus irregularis*, which is the only Glomeromycota symbiont (Tisserant et al., 2012, 2013). Among the animal pathogens, three species belonging to Microsporidia (*Enterocytozoon bieneusi*, *Nosema ceranae*, and *Antonospora locustae*) and one Ascomycota species (*P. jirovecii*) displayed large proportions (>50%; Table 2).

The proteins of other functions and the four enzymes in the refined secretomes were classified into three groups based on their cysteine content, including 0.0< 3.0%, 3.0< 5.0%, and ≥5.0%, and average percentages of these groups in four lifestyles were compared (Figure 4C). Proteins with 3% or more cysteine were considered cysteine-rich in earlier studies (Stergiopoulos and de Wit, 2009; Saunders et al., 2012). However, a more stringent criterion, over 5%, was also used (Brown et al., 2012; Krijger et al., 2014). The general trend was that both classes of cysteine-rich proteins were more abundant among the proteins of other functions than the enzymes regardless of any lifestyles.
Functional Annotation of SSPs using Interpro Terms and the Known Effector Proteins Rarely Reveals Their Functions

InterPro domain analysis was performed to predict potential functions of both SSPs and CSSPs. Most SSPs, 7960 out of 8275 (96.2%), displayed no defined InterPro terms with only 315 SSPs being annotated with 97 different terms (Supplementary Table S2). Among the annotated SSPs, the most commonly found InterPro term is membrane insertase YidC (IPR019998), which was found in 145 proteins. Other terms that are potentially related to pathogenicity and were found at least twice include IPR008427 (extracellular membrane protein, CFEM domain), IPR003172 (MD-2-related lipid-recognition domain), and IPR016191 (ribonuclease/ribotoxin).

Since the InterPro terms did not suggest any specific functions in association with pathogenicity, 49 known effector proteins from 11 fungi were retrieved from the PHI-base (Urban et al., 2015) and mapped to SSPs (Supplementary Table S3). Nine effector proteins were mapped to four species with a bit score >50 and e-value <1.e-3 (Table 3). Except MGG_10556T0 with a C2H2-type zinc finger domain, which resembles M. oryzae avirulence factor AVR-Pii, the others contained no known domains.

CSSPs were clustered with 49 PHI-base effectors via MCL clustering analysis, and 2786 families containing 19,342 proteins were identified. Among them, 13 families contained at least one PHI-base effector (Table 4). Of these families, only one is associated with InterPro term related to fungal pathogenicity. For example, proteins in the family containing M. oryzae effector MgSM1 carry a well-known domain (cerato-platanin, IPR010829; Chen et al., 2013). In total, 22 out of 49 PHI-base effectors were either matched to a single protein or clustered within a protein family (Supplementary Table S3). The small number of matches may be due to the strain-specific presence of many effectors.

Genomic Contexts of SSSP-Coding Genes

The genomic regions containing the known host-specific virulence genes have been shown to have sparsely distributed genes and AT-rich (Schmidt and Panstruga, 2011). We examined the genomic contexts of SSSP-coding genes in 59 species with number of contigs less than 500 (Supplementary Table S4). The average number of genes and the mean AT-content in each of the 100 kb segments of their genomes were calculated as references. The corresponding data for each the 100 kb windows containing SSSP-coding gene(s) were compared with the reference data.

The gene density around SSSP-coding genes was too variable to establish a clear trend in the context of lifestyles.
However, many SSSP-coding genes in a mycorrhizal symbiont L. bicolor, a plant growth promoting fungus Trichoderma virens and most wood-decaying necrotrophs were often located in regions with low gene density (Supplementary Table S4). Overall, the SSSP-coding genes did not appear to be concentrated within specific genomic region(s). On the other hand, the AT-content around most SSSP-coding genes was clearly lower than the total AT-content of the genomes of these species, except that two animal pathogens (Cryptococcus neoformans and P. jiroveci), one biotroph (Mixia osmundae), and one saprotroph (Wallemia sebi) had higher AT-contents at 52, 72, 44, and 60%, respectively (Supplementary Table S4).

The Number of SSSPs Correlates with the Proteome Size, Lifestyle, and Taxonomic Position

Several studies have reported that the size of fungal secretome correlates with lifestyle (Lowe and Howlett, 2012; Meinken et al., 2014; Lo Presti et al., 2015) and that even stronger correlation exists with phylogenetic position (Krijger et al., 2014). The former studies reported that animal pathogens and saprotrophs have similarly-sized secretomes, but their secretomes are smaller than those in plant pathogens. We assessed the size of SSSPs in individual species to determine whether similar patterns exist. The relationships between the predicted proteomes and refined secretome components are shown in Supplementary Figure S5. The pattern was similar to that observed in a previous study (Lowe and Howlett, 2012). However, the relationship between SSSPs and the total proteome was markedly different (Figure 5A). Biotrophs, symbionts, and some hemibiotrophs usually have larger numbers of SSSPs than animal pathogens, saprotrophs, and necrotrophs. The larger numbers of SSSPs in the former group, which are intimately associated with plants, support the hypothesis that these proteins are important for manipulating plant hosts.

The number of SSSPs within species ranged from 0 to 466. Although not all plant-associated fungi have higher numbers of SSSPs than saprotrophs and animal pathogens (Figure 5B),...
TABLE 4 | Protein families that contain known effector proteins.

| PHI-base accession (Effector name) | Species* | Number of proteins | Number of taxa | InterPro domain |
|------------------------------------|----------|--------------------|----------------|----------------|
| PHI:2118 (MgSM1)                  | M. oryzae| 201                | 91             | IPR009009      |
|                                   |          |                    |                | IPR010829      |
| PHI:2901 (BEC1040)               | B. graminis| 97                | 51             |                |
| PHI:2897 (BEC1019)               | B. graminis| 34                | 33             |                |
| PHI:2906 (BEC1005)               | B. graminis| 12                | 9              | IPR017853      |
| PHI:325 (ACE1)                    | M. oryzae| 9                 | 9              | IPR001509      |
|                                   |          |                    |                | IPR016040      |
| PHI:71 (ECP2)                     | C. fulvum| 7                 | 4              |                |
| PHI:2331 (Ave1)                   | V. dahliae| 5                 | 4              |                |
| PHI:1132(AvrLm4-7)               | L. maculans| 5                | 2              |                |
| PHI:2696 (NIP2)                   | R. commune| 4                | 3              |                |
| PHI:2281 (avrlm1)                 | L. maculans| 3                | 2              |                |
| PHI:1131 (AvrLm6)                 | L. maculans| 2                | 2              |                |
| PHI:379 (Six1)                    | F. oxysporum| 2            | 2              |                |
| PHI:2744 (Pit2)                   | U. maydis| 2                 | 2              |                |

*Species which the PHI-base effectors originated. IPR009009: Barwin-related endoglucanase, IPR010829: Cerato-platanin, IPR017853: Glycoside hydrolase, superfamily, IPR001509: NAD-dependent epimerase/dehydratase and IPR016040: NAD(P)-binding domain.

In general, animal pathogens and saprotrophs had fewer SSSPs compared to plant pathogens. Within plant pathogens, numbers of SSSPs encoded by crop-infecting necrotrophs are similar to those encoded by saprotrophs, and biotrophs generally have larger numbers than these groups. Within biotrophs, four Pucciniomycotina species (Puccinia graminis, P. striiformis, Melampsora laricis-populina, and M. osmundae) and B. graminis (Ascomycota) have the highest numbers. In contrast, two species in Ustilaginomycotina (Ustilago maydis and Sporisorium reilianum) and three Ascomycota species (Cladosporium fulvum, T. deformans, and Ashbya gossypii) encode small numbers of SSSPs, similar to those in saprotrophs. In addition, wood-decaying necrotrophs and hemibiotrophs have similar numbers of SSSPs.

The range of SSSPs numbers was also analyzed with regard to taxonomic positions (Figure 5C). The mean number within Basidiomycota is 100, which is higher than the mean in Ascomycota (43). Within Basidiomycota, Pucciniomycotina encodes the most SSSPs. Within Ustilaginomycotina, two biotrophs (U. maydis and S. reilianum) contained greater numbers of SSSPs compared to other members. In Ascomycota, the range was widest in Pezizomycotina with many outliers being present at both sides of the mean. Three hemibiotrophs, including M. oryzae, Leptosphaeria maculans, and Mycosphaerella graminicola, one necrotroph Stagonospora nodorum, one biotroph B. graminis, and one saprotroph Pyronema confluens encode much larger SSSPs than the mean. Saccharomycotina and Taphrinomycotina encode noticeably smaller SSSPs than Pezizomycotina.

The biotrophs in Pezizomycotina code for relatively large numbers of SSSPs. Similarly, T. deformans, the only biotroph in Taphrinomycotina, also shows a high number of SSSPs. However, A. gossypii, the only biotroph in Saccharomycotina, has a lower

FIGURE 5 | Number of SSSPs in relation to the lifestyles and taxonomic positions of the analyzed species. (A) The number of SSSPs and the size of predicted total proteome for individual species are shown. (B) The number of SSSPs among species representing different lifestyles. (C) The number of SSSPs encoded by the species in different taxa. The subphyla Glomeromycota and Zygomycota, represented by only one species each, are not included. Pez, Pezizomycotina; Sac, Saccharomycotina; Tap, Taphrinomycotina; Aga, Agaricomycotina; Puc, Pucciniomycotina; Ust, Ustilaginomycotina. C-necrotroph and W-necrotroph correspond to Crop-infecting necrotroph and Wood-decaying necrotroph, respectively.
number and proportion of SSSPs compared to other members (Figure 5C).

**Evolution of Known Effector Proteins Belonging to CSSP Families Suggests Their Other Roles**

We predicted 13 families of CSSPs containing PHI-base effectors (Table 4). Among them, families containing *M. oryzae* effector MgSM1 and the *B. graminis* effectors were found in the most taxa, and the evolution of these families of CSSPs was analyzed. Genes encoding proteins with a cerato-platanin domain (IPR009009) were conserved in 91 species, all belonging to Pezizomycotina or Agricomycotina (Figure 6). In total, 23 gene duplications and 49 gene losses were observed in the family, but the gene duplications and losses were skewed toward Agaricomycotina, which consisted mostly of wood-decaying necrotrophs. However, these genes were not found in Ustilaginomycotina and Pucciniomycotina, the other subphyla of Basidiomycota that mostly consisted of biotrophic species. The genes encoding proteins that carry the ancestral cerato-platanin domain have undergone at least five duplication events, but many of the genes also have been lost in multiple lineages. As a consequence, 33 species in Dothideomycetes and Eurotiomycetidae contained only one such gene. However, copies in the necrotrophic Sordariomycetes, *Botrytis cinerea* and *Fusarium* spp. have undergone duplication events. The wide distribution of the genes encoding cerato-platanin domain

**FIGURE 6 | Evolution of the protein family carrying the cerato-platanin domain.** The protein family with this domain is the largest cluster identified by MCL analysis with MgSM1 effector protein. Members of this family are found in 91 species within Agaricomycotina and Pezizomycotina. The cerato-platanin protein tree was reconciled with the species tree using the Notung software. The number of genes present in each species is shown at the end of nodes before species name. Gene gains (23 events) and losses (49 events) are noted by blue and red dots, respectively.
proteins in species with pathogenic lifestyles suggested that their products play an important role in pathogenesis. However, numbers of this group were also found in 4 symbionts, 2 nematophagous fungi, and 19 saprotrophs, suggesting other roles associated with them. The gene family identified using the \textit{B. graminis} effector candidate BEC1040 has gone through 16 duplications and 12 losses, and its numbers were present in 50 species in Ascomycota and \textit{Punctularia strigosozonata} in Basidiomycota (Supplementary Figure S6A). Only one duplication event occurred in the family identified with BEC1019 (Supplementary Figure S6B), but both duplication and loss events occurred in the family identified with BEC1005 (Supplementary Figure S6C). Overall, many CSSPs that resemble known effectors were identified in non-plant pathogens, raising the possibility that they are remnants of degenerated genes or play roles other than facilitating plant infection.

**DISCUSSION**

Rapid progresses in sequencing fungal genomes, in combination with various “omics” tools, have facilitated large-scale comparative genomic analyses to uncover the genetic and evolutionary basis of various traits or functions of fundamental and practical significance. In this study, we developed a pipeline for mining SSPs as effector candidates from fungi with different lifestyles and taxonomic positions in order to conduct their kingdom-wide comparative analysis. It has been commonly hypothesized that biotrophs and symbionts secrete more effectors than necrotrophs, as biotrophic associations require the modulation of the host defense system to keep host cells alive for nutrient acquisition while preventing the launch of strong defense responses. Since necrotrophs utilize CAZymes and toxins to kill host cells to obtain nutrient, such manipulations of host defense likely play less critical roles. Because only a small number of fungal proteins in selected plant pathogens have been identified as effectors, we tested this hypothesis by comparing SSPs as effector candidates.

We refined secretomes to identify SSPs as previously reported in analyzing the secretome of \textit{F. graminearum} (Brown et al., 2012). In addition, we mined and compared other components within the refined secretomes to investigate any lifestyle-associated genomic adaptations. Three previous studies examined potential relationships between the secretome and lifestyle (Lowe and Howlett, 2012; Krijger et al., 2014; Lo Presti et al., 2015). However, Lowe and Howlett (2012) and Lo Presti et al. (2015) used only one signal peptide prediction program for mining secretomes and Krijger et al. (2014) did not eliminate putative membrane-bound proteins, which likely inflated the size of secretomes. Our refined secretome pipeline more rigorously identified secretory proteins via two additional protein localization detection programs and the elimination of transmembrane and GPI-anchor proteins. For comparison, the number of SSPs and the effector candidates in powdery mildew predicted by Spanu et al. (2010) is the same, and the refined secretomes predicted in both corn pathogens \textit{S. reilianum} and \textit{U. maydis} are similar to those reported by Schirawski et al. (2010). The corn pathogen study found that many effector candidate genes from both pathogens were orthologous, consequently the number of SSPs for them were drastically reduced in our study. When the refined secretome of \textit{F. graminearum} was compared to the data by Brown et al. (2012), their 539 out of 574 proteins were included in our 961 secretome. Although we predicted larger secretome than their refined secretome, this is due to the lowered parameter settings for prediction as our parameters were determined based on the effectors listed in Stergiopoulos and de Wit (2009) for kingdom-wide analysis. We also used a greater number of species to cover more diverse taxa and further divided plant pathogens into four groups to perform comprehensive analyses.

The roles of fungal effector proteins regarding lifestyles are previously discussed by Lowe and Howlett (2012) and Lo Presti et al. (2015). These studies suggested that fungi with same lifestyles have similar secretome proportions. The secreted CAZymes also showed similar pattern that biotrophs encode fewer CAZymes than hemibiotrophs and necrotrophs (Zhao et al., 2014; Lo Presti et al., 2015). Although our overall conclusion on refined secretome and CAZymes may be similar with the previous studies, the numbers of SSPs show lifestyle adaptation, different from secretomes and CAZymes analyses. In addition, secretomes contain not only CAZymes for cell wall degradation and utilization of its components as nutrients but also proteases, lipases, and oxidoreductases for breakdown of other macromolecules, and self-protection and/or pathogenesis. For example, AVR-pita of \textit{M. oryzae} is a zinc metalloprotease that acts as an avirulence factor in its host rice (Zhang and Xu, 2014). The lipase effector FGL1 in \textit{F. graminearum} suppresses callose formation in wheat and is required for host infection (Blümke et al., 2014). Oxidoreductases have been investigated in phytopathogenic fungi for their roles in scavenging plant reactive oxygen species and pathogenicity (Chi et al., 2009). Since their importance in pathogenesis has been established or suggested, we examined the proportions of these enzymes relative to the whole proteome in light of the lifestyle and taxonomic position of individual species. Proteases, lipases, and oxidoreductases seem to be more abundant in plant pathogens, especially in hemibiotrophs and necrotrophs, than the species with different lifestyles. Since the majority of known fungal effector proteins do not possess enzymatic activity, we excluded the above enzyme sets prior to mining SSPs, which include short proteins (\textlesss\text{300} aa) with a signal peptide, but no transmembrane domain or GPI-anchor. Species-specific presence was also analyzed to classify SSPs into SSPs, which likely act as host-specific effectors. Overall, we found that the size of refined secretomes and SSPs varies widely between species, but some patterns associated with lifestyles.

In general, phytopathogenic fungi tend to have larger secretomes than non-pathogens. Although, Lowe and Howlett (2012) suggested that animal pathogens generally have a lower proportion of secretome, the sizes of secretomes and proteins of other functions in certain animal pathogens, such as nematophagous and entomopathogenic fungi, were similar to some necrotrophic plant pathogens. This is not too surprising considering that nematophagous and entomopathogenic fungi secrete diverse proteins to facilitate infection and consumption of hosts (Andersson et al., 2013; Staats et al., 2014). The host-specific animal pathogens that co-evolved with hosts for longer
period of time may have large secretome, yet smaller than those of phytopathogens. Although symbionts intimately interact with plant hosts, their secretomes are smaller than those of pathogens, a pattern that was found in previous studies on symbionts such as L. bicolor (Martin et al., 2008) and Tuber melanosporum (Martin et al., 2010). This is due to the reduced number of CAZymes compared to necrotrophs and hemibiotrophs. Biotrophs encode smaller sets of CAZymes, but larger secretomes than non-pathogens mainly due to their abundant SSSPs. This reflects the lifestyle of biotrophs which causes minimal damages to the hosts to maintain a long-term feeding relationship. Not surprisingly, reduced numbers of SSSPs are observed among crop-infecting necrotrophs. However, wood-decaying necrotrophs conspicuously possess a similar level of SSSPs to hemibiotrophs. Their roles of SSSPs in wood-decay remain unclear. Within saprotrophs, fruit-body forming fungi and fungi displaying antifungal activities encode greater numbers of SSSPs than yeasts and extremophiles. However, the number of SSSPs has no correlation with host range within necrotrophs, types of rot for wood-decaying necrotrophs, or types of association for symbionts. As illustrated by Neurospora crassa, a saprotrophic fungus that has been reported to be associated with pine trees in harsh conditions and even pathogenic (Kuo et al., 2014), fungal lifestyles may not be a fixed attribute, but is changeable depending on environmental conditions.

Another concern is that the number of small proteins could be affected by the minimum protein size used to annotate each genome. However, we strictly used the annotated data associated with published genomes, thus we can assume that the numbers of annotated proteins were comparable to each other. In addition, the majority of genome data were also from JGI and Broad Institute, which followed the conventional JGI annotation process and Broad Gene Finding Methods, respectively. If the minimum cutoff length for gene prediction was stated, only short peptides without EST support were eliminated. Since, the majority of fungal genome studies did not show the cutoff size for protein-coding gene prediction, we analyzed and compared the length distribution of the annotated proteomes for validation (the last column of Supplementary Table S1). In result, no parameters were possibly used for 50 species and additional 72 species with the cutoff of 30 aa. The rest 14 species contained proteins with length at least 50 aa. The number of short proteins in the first category was extremely high for a few species such as C. fulvum and P. confluens. However, only the proteins secreted with canonical pathways were considered in this analysis, which means the minimum size of protein is bound to the length of signal peptide that is 15 to 40 aa (Choo et al., 2005). Although many short proteins were annotated using no or very short cutoff, they were eliminated if the signal peptide was absent. Overall, we believe that the numbers of proteins are generally comparable like in other secretome studies.

Many genes coding candidate effectors in plant pathogens and virulence-associated genes of animal pathogens have been reported to reside in gene-sparse, AT-rich, and telomere-proximal genomic regions (Schmidt and Panstruga, 2011). However, genomic distribution patterns of SSSP-coding genes did not display similar trends, with the exception of those in the wood-decaying necrotrophs. Most SSSP-coding genes in these fungi were found in genomic regions with low gene numbers. A previous study reported that 23% of the proteins encoded by these wood-decaying fungal species are unique (Riley et al., 2014), indicating that the SSSP-coding genes recently arose from non-coding sequences as suggested by Carvunis Model (Carvunis et al., 2012). However, there are still many SSSP-coding genes located in AT-rich regions. Therefore, the genomic context could be considered for prioritizing validation of effector functions. Although effector proteins are often thought to be species-specific due to co-evolution with the host, there are cases of conserved effectors within related species. For example, the C. fulvum effector Ecp2 (Stergiopoulos et al., 2012) and the cerato-platanin proteins (Chen et al., 2013) are conserved only within Ascomycota and Basidiomycota. Moreover, the clustering analysis of CSSPs showed that none of them seem to be conserved across the fungal kingdom, indicating that they are typically limited to specific genera. However, some CSSPs of B. graminis, e.g., BEC1040, BEC1019, and BEC1005, were found in fungi having different lifestyles other than biotroph. These observations were further supported by the facts that these proteins are involved in fungal development, and resemble metalloprotease and glucanase, respectively (Pliego et al., 2013). Taken together, these suggest that the numbers of CSSPs may not be correlated with fungal lifestyles.

In conclusion, different secretome components reflect lifestyle-associated genomic adaptations in fungi. Results from this comparative study provide new insights into the genetic basis and molecular evolution of fungal lifestyles and also establish a solid foundation for future discovery and functional validation of effectors.

**AUTHOR CONTRIBUTIONS**

KK and YL designed this project. KK, JJ, HS and GC performed computational analyses. JC and KC provided the secretome and the genome data for analyses. KK, JJ, SK, and YL wrote the manuscript. SK and YL supervised the research. All authors read and approved the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016.00186
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