Mitogen-activated protein kinases (MAPKs) are evolutionarily conserved proteins that function as key signal transduction components in fungi, plants, and mammals. During interaction between phytopathogenic fungi and plants, fungal MAPKs help to promote mechanical and/or enzymatic penetration of host tissues, while plant MAPKs are required for activation of plant immunity. However, new insights suggest that MAPK cascades in both organisms do not operate independently but that they mutually contribute to a highly interconnected molecular dialogue between the plant and the fungus. As a result, some pathogenesis-related processes controlled by fungal MAPKs lead to the activation of plant signaling, including the recruitment of plant MAPK cascades. Conversely, plant MAPKs promote defense mechanisms that threaten the survival of fungal cells, leading to a stress response mediated in part by fungal MAPK cascades. In this review, we make use of the genomic data available following completion of whole-genome sequencing projects to analyze the structure of MAPK protein families in 24 fungal taxa, including both plant pathogens and mycorrhizal symbionts. Based on conserved patterns of sequence diversification, we also propose the adoption of a unified fungal MAPK nomenclature derived from that established for the model species *Saccharomyces cerevisiae*. Finally, we summarize current knowledge of the functions of MAPK cascades in phytopathogenic fungi and highlight the central role played by MAPK signaling during the molecular dialogue between plants and invading fungal pathogens.

INTRODUCTION

The development of all organisms relies on the ability of cells to sense and respond rapidly to changes in the surrounding environment. To coordinate appropriate cellular actions, eukaryotes use diverse receptors that perceive specific cues and relay information to intracellular signaling pathways. The interface created during the interaction between a phytopathogenic fungus and its host is a rich source of signals that feed into this molecular dialogue, resulting in rapid and highly structured responses in both protagonists.

Despite completely different lifestyles, fungi and plants rely on many analogous signaling pathways to coordinate their respective cellular actions. Among evolutionarily conserved pathways, mitogen-activated protein kinase (MAPK) cascades function as key signal transducers that use protein phosphorylation/dephosphorylation cycles to channel information. MAPK cascades in all eukaryotes generally consist of three interlinked protein kinases (PKs) that are sequentially activated (Widmann et al., 1999). Activated MAPK kinase kinases (MAP3Ks) first phosphorylate two Ser and/or Thr residues located within the activation loop of MAPK kinases (MAP2Ks). Activated MAP2Ks in turn trigger MAPK activation through dual phosphorylation of a highly conserved activation loop that possesses the hallmark motif -TXY-. Activated MAPKs can then phosphorylate downstream substrates, affecting their biochemical properties and leading to specific output responses.

*Arabidopsis thaliana* has been widely employed as a model to examine MAPK functions in the plant kingdom. Plant MAPKs have been shown to regulate numerous cellular processes, including biotic stress relief (Pitizhke et al., 2009; Andreasson and Ellis, 2010). In fungi, the study of MAPKs from *Saccharomyces cerevisiae* has provided pivotal insights that have contributed greatly to our understanding of MAPK signaling in all eukaryotes (Chen and Thorner, 2007). This has been particularly important for studies of other fungi, including both human and plant pathogens, where orthologous MAPK signaling modules have been found to be involved in the control of infection-related morphogenesis (IRM), virulence, cell wall biogenesis, and stress responses (Xu, 2000; Zhao et al., 2007; Rispail et al., 2009).

In this review, we aimed at identifying the full complement of MAPK signaling components in a range of taxonomically diverse fungi that yet all interact with plants. Based on the analysis of recovered protein sequences, we could resolve the fungal MAPK family into four separate clades, while three distinct subgroups were defined for each of the fungal MAP2K and MAP3K protein families. As clustering of these signaling components is recapitulated in
the model fungal species \textit{S. cerevisiae}, we propose a unified nomenclature that relies on gene/protein names from yeast to identify homologous candidates from the investigated fungal species. To avoid ambiguous annotation of orthologous candidates, we also make use of three-letter species acronyms that indicate the origin of each gene or protein (Table 1). Finally and as a complement to this, we summarize current knowledge of the functions of MAPK cascades in phytopathogenic fungi and highlight the central role played by MAPK signaling during the molecular dialogue between plants and fungal pathogens.

\section*{MAPKs FROM BUDDING YEAST}

In \textit{S. cerevisiae}, MAPK cascades constitute essential signaling pathways that are involved in key aspects of the yeast life cycle (Chen and Thorner, 2007). As a result, these protein modules as well as both their upstream regulators and downstream substrates are highly conserved even in distantly related fungi, including plant pathogens (Rispal et al., 2009). \textit{S. cerevisiae} thus represents an excellent model for the study of MAPK signaling in fungi. To allow better understanding of MAPK signaling in phytopathogenic fungi, we first provide a brief description of prototypical MAPK cascades from yeast.

In the budding yeast, five MAPK pathways regulate mating, invasive growth, cell wall integrity, hyperosmolarity responses, and ascospore formation (Chen and Thorner, 2007). The yeast mating response depends on G protein–coupled receptors (Ste2 and Ste3) that bind cognate peptidic pheromones (Figure 1). This binding results in dissociation of the inhibitory G\textsubscript{i} subunit Gpa1, from Ste4 and Ste18, which respectively function as stimulatory G\textsubscript{b} and G\textsubscript{y} subunits. Released G\textsubscript{b}/G\textsubscript{y} subunits associate with the scaffolding protein Ste5, and the p21-activated kinase (PAK) Ste20, to create a protein complex that activates the MAP3K Ste11. Ste11 is the entry point for a MAPK cascade that also includes MAP2K Ste7 and two partially redundant MAP3Ks, Kss1 and Fus3. The cyclin-dependent kinase (CDK) inhibitor Far1 and the transcription factor (TF) Ste12, which respectively control cell cycle arrest and expression of mating-responsive genes, are primary targets of the pheromone response pathway (Figure 1). Many components of the pheromone pathway, including Ste20, Ste50, Ste11, Ste7, Kss1, and Ste12, are also involved in regulating filamentous growth (Figure 1). This response is stimulated upon starvation and leads to invasive growth in haploid cells or pseudohyphal development in diploid cells. Under nutrient-rich conditions, inactive Kss1 localizes to the nucleus, where it sequesters TFs such as Ste12 and Teco. Phosphorylation of Kss1 relieves its negative regulation and results in TF release. However, upstream components of the pheromone pathway, such as the pheromone receptors, heterotrimeric G proteins, and the Ste5 scaffold protein, are not required for filamentation.

\begin{table}[h]
\centering
\begin{tabular}{llll}
\hline
Phylum & Organism & Prefix & Plant Disease & Infection Strategy \\
\hline
\textbf{Ascomycota} & \textit{A. brassicicola} & Abr & \textit{Brassica} dark leaf spot disease & Neotrophi \\
\textbf{Ascomycota} & \textit{B. cinerea} & Bci & Polyphage (e.g., grape noble rot) & Neotroph \\
\textbf{Ascomycota} & \textit{C. heterostrophus} & Che & Southern corn leaf blight disease & Neotroph \\
\textbf{Ascomycota} & \textit{C. parasitica} & Cpa & Chestnut blight disease & Neotroph \\
\textbf{Ascomycota} & \textit{F. graminearum} & Fgr & Wheat head-blight disease & Neotroph \\
\textbf{Ascomycota} & \textit{F. oxysporum} & Fox & Polyphage (several \textit{formae speciales}) & Hemibiotroph \\
\textbf{Ascomycota} & \textit{Fusarium verticillioides} & Fve & Rice bakanae disease & Neotroph \\
\textbf{Basidiomycota} & \textit{H. annosum sensu lato} & Han & Conifer root rot disease & Neotroph \\
\textbf{Basidiomycota} & \textit{L. bicolor} & Lbi & None: mutualist & Ectomycorrhizae \\
\textbf{Ascomycota} & \textit{M. oryzae} & Mor & Rice blast disease & Hemibiotroph \\
\textbf{Basidiomycota} & \textit{M. larici-populina} & Mlp & Poplar leaf rust disease & Obligate biotroph \\
\textbf{Ascomycota} & \textit{Mycosphaerella fijiensis} & Mfi & Banana (\textit{Musa} spp) black leaf spot disease & Neotroph \\
\textbf{Ascomycota} & \textit{M. graminicola} & Mgr & \textit{Septoria} leaf blotch of wheat & Neotroph \\
\textbf{Ascomycota} & \textit{Mycosphaerella pini} & Mpi & Pine needle blight disease & Hemibiotroph \\
\textbf{Ascomycota} & \textit{Mycosphaerella populorum} & Mpo & \textit{Septoria} leaf spot and canker of poplar & Hemibiotroph \\
\textbf{Basidiomycota} & \textit{P. graminis-tritici} & Pgr & Wheat and barley stem rust disease & Obligate biotroph \\
\textbf{Basidiomycota} & \textit{T. tritici} & Put & Wheat leaf rust disease & Obligate biotroph \\
\textbf{Ascomycota} & \textit{Pyrenophora tritici-repentis} & Pte & Wheat yellow leaf spot disease & Neotroph \\
\textbf{Ascomycota} & \textit{S. cerevisiae} & Sce & None: does not interact with plants & None \\
\textbf{Ascomycota} & \textit{S. nodorum} & Sno & \textit{S. nodorum} blotch of wheat & Neotroph \\
\textbf{Ascomycota} & \textit{Sclerotinia sclerotionum} & Ssc & White mold disease & Neotroph \\
\textbf{Ascomycota} & \textit{Trichoderma virens} & Tvi & None: biocontrol agent & Mycoperasitism \\
\textbf{Ascomycota} & \textit{T. melanosporum} & Tme & None: mutualist & Ectomycorrhizae \\
\textbf{Basidiomycota} & \textit{U. maydis} & Uma & Maize smut disease & Biotroph \\
\textbf{Ascomycota} & \textit{V. dahliae} & Vda & Wilt disease & Neotroph \\
\hline
\textbf{Not included in phylogenetic analysis} & & & & \\
\textbf{Ascomycota} & \textit{C. purpurea} & Cpu & Ergot of rye and other grains & Biotroph \\
\textbf{Ascomycota} & \textit{C. orbiculare} & Cor & \textit{Cucumber} (\textit{Cucumis sativus}) anthracnose disease & Hemibiotroph \\
\textbf{Basidiomycota} & \textit{P. striiformis} & Pst & Wheat stripe rust & Obligate biotroph \\
\textbf{Ascomycota} & \textit{P. teres} & Pte & Barley net blotch & Neotroph \\
\hline
\end{tabular}
\caption{Plant-Interacting Fungi and Three-Letter Species Prefixes}
\end{table}
In this developmental context, activation of the Ste11-Ste7-Kss1 module instead depends on osmosensors Sho1 and Msb2 (Figure 1).

A third yeast MAPK cascade forms part of the cell integrity sensing pathway and consists of the MAP3K Bck1, two redundant MAP2Ks, MKK1 and MKK2, as well as the Slt2 MAPK (Figure 1). This cascade is required to maintain cell wall integrity and is involved in compensatory responses against environmental stresses like high temperature or exposition to heavy metals. Surface sensors perceive external cues and relay information to Rom2, a guanine exchange factor (GEF) that regulates the activity of the Rho1 GTPase. Active Rho1 then activates PKC1, a homolog of the α, β, and γ isoforms of mammalian PK C. PKC1 then phosphorylates Bck1 to initiate MAPK signaling through the Slt2 cascade. Associated with this module is the Spa2 adaptor protein, which directs appropriate MAPK signaling components to cell growth sites. Activated Slt2 then modifies downstream targets, such as the TFs Rlm1, Sbf1, and Swi4/6, which are involved in regulating the transcriptional programming required for cell wall biogenesis (Figure 1).
In yeast cells grown under hypertonic conditions, the high osmolarity glycerol (HOG) pathway (Figure 1) is required for accumulation of osmoregulator molecules and maintenance of an osmotic gradient across the stressed plasma membrane. This pathway leads to the activation of Hog1, a stress-related MAPK. The Pbs2 MAP2K and three MAP3Ks (Ste11, Ssk2, and Ssk22) complete this MAPK module, which can be activated by two signaling input branches converging at the level of Pbs2. While activation of Ssk2 and Ssk22 depends on a two-component His kinase phosphorelay system (Sin1, Ypd1, and Ssk1), activation of Ste11 depends on a signaling pathway shared with the filamentous growth pathway and comprising proteins Sho1, Msb2, Cdc42, Ste20, and Ste60. Several TFs, including Hot1, Msn2, and Sko1, lie downstream of Hog1 and control expression of genes involved in osmotic as well as oxidative stress responses (Figure 1).

Smk1, the fifth MAPK from S. cerevisiae, is unique because it is only expressed during late stages of meiosis, prior to ascospore enclosure (Krisak et al., 1994). Smk1, which has no homolog in other fungi except in some ascomycetous yeasts, regulates spore morphogenesis by controlling assembly of the ascospore wall. The Smk1 pathway is the least well defined, with neither the upstream MAP2K nor MAP3K so far identified (Figure 1). Since PAK Sps1 is also involved in ascospore wall assembly, it is believed to work upstream of the Smk1 MAPK (Friesen et al., 1994).

**PHYLOGENY OF MAPK FAMILIES IN PLANT-INTERACTING FUNGI**

Considering the importance of MAPK cascades in fungal biology and virulence against plants (see the MAPKs from phytopathogenic fungi section), we took advantage of genomic databases that are available following completion of genome sequencing projects (U.S. Department of Energy Joint Genome Institute, Broad Institute of MIT and Harvard, and Genoscope Sequencing Center, France; see Supplemental Table 1 online) to identify the full complement of associated MAPK signaling components. Overall, the genomes of 24 taxonomically diverse fungi were analyzed in our survey. A total of 260 genes and corresponding proteins were identified, including 112 MAPKs, 74 MAP2Ks, and 74 MAP3Ks (see Supplemental Data Sets 1 to 3, respectively, and Supplemental References 1 online). Importantly, more than 50% of all identified candidates correspond to PKs that have never been described in the literature, suggesting that the assembled information will provide a useful resource for researchers wishing to compare and investigate MAPK modules in plant-interacting fungi. In the case of model pathogen Magnaporthe oryzae, Ustilago maydis, Fusarium spp, and Botrytis cinerea, our screen was accurate enough to retrieve all the previously identified MAPK signaling components (Dean et al., 2005; Garcia-Pedrajas et al., 2008; Rispail et al., 2009; Ma et al., 2010; Amselem et al., 2011; see Supplemental Data Sets 1 to 3 online). We therefore presume that remaining gene models and associated proteins identified here correspond to the full complement of MAPK signaling components in the investigated fungi.

**MAPKs**

Full-length MAPK protein sequences were aligned (see Supplemental Data Set 4 online), and a phylogenetic analysis was conducted (Figure 2). This showed that the fungal MAPK family can be resolved into four separate clades, each of which contained at least one representative member from each of the investigated fungal species (Figure 2). Certain MAPK homologs display a strikingly high level of similarity, even though they belong to different fungal species (e.g., Kss1/Fus3-type homologs from Dothideomycetes; Figure 2; see Supplemental Data Set 1 online). With the exception of Verticillium dahliae, which has two Hog1 homologs, all the ascomycetes examined have four conserved MAPK candidates, including a single homolog for each of the yeast Kss1 (or Fus3), Slt2, and Hog1 prototypical MAPKs (Figure 2; see Supplemental Data Set 1 online). On the other hand, basidiomycete species generally possess a larger number of MAPKs, with up to seven or eight candidates in the Agaricomycetes (Figure 2; see Supplemental Data Set 1 online). The increased number of MAPKs in these species most likely results from duplication events that created extra gene copies retained during diversification of the various basidiomycete species. Gene duplication is observed for some Slt2 and Hog1 homologs, although this feature is particularly prominent for MAPKs homologous to yeast Kss1 and Fus3 (Figure 2; see Supplemental Data Set 1 online). Since most of the duplicated MAPKs are found on scaffolds that have not yet been assigned to a specific chromosome, it is not clear which kind of duplication events might be involved in the expansion of the MAPK family in basidiomycetes. However, since no obvious MAPK gene cluster could be identified, this suggests that family expansion is not associated primarily with tandem gene duplication, but rather with whole-genome or large segmental duplication events. Interestingly, the dichotomy existing between ascomycetes and basidiomycetes is recapitulated in the clustering of MAPKs (Figure 2; this is also true for MAP2Ks and MAP3Ks). For basidiomycetous species, the segregation of MAPK candidates also reflects the established phylogeny associated with fungal classes (Ustilaginomycetes, Puccinimycetes, and Agaricomycetes; Figure 2). All the investigated species also contain a single MAPK-related protein that is homologous to yeast Ime2 (Figure 2; see Supplemental Data Set 1 online). When compared with classical MAPKs, Ime2 homolog sequences have diverged more extensively, yet these proteins have been sufficiently preserved to define a fourth clade of MAPKs (Figure 2).

**MAP2Ks**

The results of phylogenetic analysis of MAP2K sequences indicate that this protein family can be divided into three clades, each of which contains at least one representative from each of the examined fungi (Figure 3; see Supplemental Data Set 5 online). With the exception of the two Agaricomycetes, all surveyed fungal species have three MAP2K candidates, which appear to be the homologs for the yeast Ste7, MKK1 (or MKK2), and Pbs2 prototypical MAP2Ks (Figure 3; see Supplemental Data Set 2 online). In contrast with the pattern observed among the basidiomycetous MAPKs, it seems that gene duplication has not led to expansion of the fungal MAP2K family. The only obvious duplication events, including the presence of two MKK1/2 homologs in Heterobasidion annosum sensu lato (also referred to as H. irregulare; see Olson et al., 2012) and two Ste7 homologs in...
Figure 2. Phylogenetic Relationships of MAPKs from Plant-Interacting Fungi.
**Laccaria bicolor** (Martin et al., 2008), seem to be lineage-specific events since they are not recapitulated in the other Agaricomycetes examined, namely, *L. bicolor* and *H. annosum sensu lato* (Figure 3; see Supplemental Data Set 2 online). The relatively small number of MAP2Ks encoded in these genomes also suggests that each of these proteins likely activates multiple MAPKs, a scenario that may be more prominent in basidiomycetes that possess recently duplicated gene pairs in several clades of MAPKs. Experimental support for this prediction comes from the study of *U. maydis*, where the MAP2K Fuz7 (here renamed Uma-Ste7 according to our unified nomenclature; see below) has been shown to operate upstream of several MAPKs (Garrido et al., 2004; Di Stasio et al., 2009).

**MAP3Ks**

Phylogenetic analysis of fungal MAP3K sequences demonstrates that this family also consists of three well-resolved clades, defined by homologs of the yeast Ste11, Bck1, and Ssk2/Ssk22 prototypical MAP3Ks (Figure 4; see Supplemental Data Set 6 online). Surprisingly, the ascomycetes *Alternaria brassicicola* and *Stagonospora nodorum* possess only two predicted MAP3Ks and lack Ssk2/Ssk22 or Bck1 homologs, respectively (Figure 4; see Supplemental Data Set 3 online). This situation may be due to sequencing artifacts, since MAPKs and MAP2Ks that presumably function downstream of the apparently missing MAP3Ks can be identified in the genome of both fungi. With the exception of *Fusarium oxysporum*, all examined ascomycetes have a single candidate for each of yeast Ste11, Bck1, and Ssk2/Ssk22 MAP3Ks (Figure 4; see Supplemental Data Set 3 online). In *F. oxysporum*, duplication of Bck1 homologs seems lineage specific, as this pattern is not conserved even in closely related *Fusarium* species (Figure 4; see Supplemental Data Set 3 online; Ma et al., 2010). Interestingly, lineage-specific duplication of Bck1 homologs can also be identified in basidiomycetes such as the rust fungus *Melampsora larici-populina* and *Puccinia graminis-tritici* as well as within the ectomycorrhizal fungus *L. bicolor* (Figure 4; see Supplemental Data Set 3 online; Martin et al., 2008, 2010). Although they belong to distinct taxa (*Basidiomycota* for *L. bicolor* and Ascomycota for *T. melanosporum*), both of these fungi establish ectomycorrhizal symbioses with forest trees. Genome sequence analysis shows that MAPKs are relatively well conserved in *T. melanosporum* compared with other ascomycetes, whereas *L. bicolor* presents a few gene family expansions compared with other fungi (for more details, see phylogenetic classification of MAPK families in the preceding sections). Genome-scale transcriptome studies show that MAPK genes are relatively highly expressed in the free-living mycelium, fruiting bodies, and ectomycorrhiza compared with other conserved genes (Martin et al., 2008, 2010), indicating that these signaling pathways likely play a role in the development and functioning of the symbiotic association. Although no genome sequence is yet available from endomycorrhizal fungi in the Glomeromycota, the recent report of nonredundant virtual transcripts from the fungus *Glomus intraradices* confirms the presence and expression of several MAPK signaling components (Tisserant et al., 2012).

At the functional level, only limited information is currently available on MAPK signaling in symbiotic fungi. In the ectomycorrhizal fungus *Tuber borchii*, the Kss1/Fus3-type MAPK TBMK (here renamed Tbo-Kss1 according to our unified nomenclature; see below) becomes phosphorylated during interaction of the fungus with its host tree, *Tilia americana* (Menotta et al., 2006). Interestingly, ectopic expression of this MAPK in *F. oxysporum* strains lacking *Fmk1* (here renamed Fox-Kss1; see below) partially restores their ability to grow invasively (Menotta et al., 2006). This suggests that MAPK signaling could play an important role during the presymbiotic colonization phase, perhaps by modulating expression of target genes necessary to allow host infection and establishment of functional ectomycorrhizae. The induction of fungal MAPKs...
Figure 3. Phylogenetic Relationships of MAP2Ks from Plant-Interacting Fungi.
UNIFIED NOMENCLATURE FOR FUNGAL MAPK FAMILIES

In the past years, MAPK signaling has been studied in a wide array of fungi, including several plant pathogens and symbionts (reviewed in Xu, 2000; Zhao et al., 2007; Rispail et al., 2009; this article). During this time, research reports have generated disparate and confusing gene/protein designations that do not necessarily reflect putative orthologous relationships between homologs from distinct fungal species (see Supplemental Data Sets 1 to 3 online). Trivial naming also marked the plant MAPK homologs from distinct fungal species (see Supplemental Data Set 1 online). For each family. MAP2Ks should now belong to the Ste7, MKK1/MKK2, or Pbs2 type (Figure 3; see Supplemental Data Set 2 online). In relevant cases, previous MAP2K nomenclature is indicated in parenthesis.

MAPKs FROM PHYTOPATHOGENIC FUNGI

Following characterization of the first MAPK from a fungal plant pathogen (Xu and Hamer, 1996), similar strategies have been employed to examine the role of MAPKs in other phytopathogenic fungi. Generally, degenerate primers allow PCR-based amplification of MAPK genes, and associated functions are assessed through loss-of-function studies. Knockout strains can, for instance, be examined for phenotypes involving altered vegetative growth and virulence. More recently, comparative genomics has facilitated study of fungal signaling pathways (Rispail et al., 2009), and complete MAPK cascades have been functionally defined in model plant pathogens.

The Kss1-1/Kss1-2 MAPK Cascade in U. maydis

U. maydis is the causal agent of maize (Zea mays) smut disease, an infection characterized by the development of tumor-like structures called galls (Kahmann and Kämper, 2004; Brefort et al., 2009). This basidiomycete has been widely used as a model species to study dimorphism, a process that refers to the ability of certain fungi to switch between unicellular and multicellular growth forms to maintain resource uptake under changing conditions (Nadal et al., 2008). For some phytopathogenic fungi, this morphology switch is a prerequisite to plant infection, and in most cases, an encounter with the host is responsible for triggering the dimorphic shift.

In the absence of host plants, U. maydis develops as a saprobic, haploid, unicellular organism that is incapable of infecting maize (Kahmann and Kämper, 2004; Brefort et al., 2009). Upon contact with the host, compatible haploid cells mate and initiate formation of a dikaryotic filament. This transition marks the initiation of a parasitic growth stage typified by appressorium formation, penetration, and the formation of filamentous hyphae inside plant tissues. Mating of U. maydis cells depends on a pheromone-receptor system that is coupled to MAPK

predicted evolutionary relationships and that conservation of biological function within most of these relationships remains to be confirmed. Indeed, disruption of a particular MAPK type may well have unequal phenotypic effects in various fungal species, since those effects will be modulated by interactions of the kinase within the larger genetic and cellular context of each species. Nonetheless, the systematic nomenclature proposed here should facilitate species-to-species comparisons and limit the expansion of trivial naming that already marks the MAPK family from phytopathogenic fungi (see Supplemental Data Set 1 online).
Figure 4. Phylogenetic Relationships of MAP3Ks from Plant-Interacting Fungi.

Genome assembly from various fungi was searched using amino acid sequence of yeast MAP3Ks as queries. Retrieved gene models were accepted only if corresponding protein contained consensus sequences for Ser/Thr PKs, including conserved Asp and Lys residues within the active site (D[L/I/V]K motif).
signaling (Figure 5; Kahmann and Kämper, 2004; Brefort et al., 2009). Continued parasitic growth depends on the same MAPK cascade, as well as on another signaling pathway that is activated by modulation of the intracellular level of cAMP (Figure 5).

The *U. maydis* mating apparatus includes a seven-transmembrane receptor (Pra1 or Pra2), which recognizes cognate lipopeptide pheromones (Mfa2 or Mfa1, respectively; Figure 5; Kahmann and Kämper, 2004; Brefort et al., 2009). Downstream of the receptor lies a MAPK cascade consisting of the MAP3K Uma-Ste11 (previously named Kpp4/Ubc4), the MAP2K Uma-Ste7 (Fuz7/Ubc5), and the MAPK Uma-Kss1-1 (Kpp2/Ubc3; Figure 5; Banuett and Herskowitz, 1994; Mayorga and Gold, 1999; Andrews et al., 2000). Ste7 is essential for mating, filamentous growth, and virulence and was the first component of the cascade to be characterized (Banuett and Herskowitz, 1994). It was later found that deletion of Kss1-1 partially compromises mating, formation of filamentous dikaryons, and virulence (Mayorga and Gold, 1999; Müller et al., 2003b). Partial phenotypes are correlated with the existence of Uma-Kss1-2 (also named Kpp6; Figure 5), a closely related MAPK that possesses an unusually long N-terminal extension (Brachmann et al., 2003). Kss1-2 participates in regulation of virulence since deletion mutants are unable to penetrate the plant cuticle (Brachmann et al., 2003). Most importantly, Kss1-1/Kss1-2 double knockouts fail to mate and are nonpathogenic, suggesting that these MAPKs possess at least partially overlapping functions during mating and virulence (Figure 5; Zhao et al., 2007; Brefort et al., 2009; Rispail et al., 2009). Consistent with this, Kss1-1 and Kss1-2 share Ste7 as an upstream activator, as well as Rok1, a member of the dual-specificity protein phosphatase family, as a negative regulator (Figure 5; Di Stasio et al., 2009).

A loss-of-function study also indicates that *Ustilago* MAP3K Ste11 lies upstream of the Ste7-Kss1-1 module, since Δste11 mutants are defective in both mating and virulence (Figure 5; Andrews et al., 2000; Müller et al., 2003b). Ste11 also interacts with Ubc2 (Figure 5), an adaptor protein with homology to Ste50 from *S. cerevisiae* (Figure 1; Klosterman et al., 2008). Ubc2 had previously been associated with regulation of the Kss1-1 cascade, since its deletion impairs pheromone response and virulence (Mayorga and Gold, 2001). Ubc2 harbors sterile α motif (SAM) and Ras association domains that are typical of Ste50-like adaptors from ascomycetous fungi, as well as Src homology 3 domains that are conserved only in Ubc2-like homologs from basidiomycetous fungi (Klosterman et al., 2008). Yeast two-hybrid (Y2H) assays indicate that Ubc2 and Ste11 interact via their respective SAM domains, and deletion experiments have shown that the SAM and Ras association domains are indispensable for mating and filamentous growth, while Src homology 3 domains specifically regulate virulence (Klosterman et al., 2008).

One of the Ras proteins from *U. maydis* has also been positioned upstream of Ubc2 and its cognate MAPK cascade (Figure 5; Lee and Kronstad, 2002). Thus, expression of a dominant active ras2 allele promotes pseudohyphal growth in a manner dependent on the Kss1-1 MAPK cascade. One possible activator of Ras2 is the GEF protein Sql2 (Figure 5; Müller et al., 2003a). However, unlike ras2 mutants, sql2 knockouts are defective in plant infection but not in mating. It has been suggested that Sql2 may activate Ras2 only under certain circumstances, for example, following perception of host-derived signals (Klose et al., 2004). Other signaling components acting between the *Ustilago* pheromone receptors and the Kss1-1 MAPK cascade remain unknown and may differ from those operating in the yeast pheromone pathway (Figure 1). For instance, none of the four Gα subunits from *U. maydis* could be linked with pheromone receptors (Zhao et al., 2007). In addition, the PAK Smu1 is dispensable for mating and plant infection (Figure 5; Smith et al., 2004), even though it shares high homology with PAK Ste20 from yeast (Figure 1).

In *U. maydis*, pheromone perception induces transcription of several target genes, including those located within the loci “a” and “b” (Kahmann and Kämper, 2004; Brefort et al., 2009). The “a” locus contains two genes encoding mating pheromones and receptors, while the “b” locus contains two homeodomain genes named *be* and *bw* (Figure 5). Regulation of these genes requires the Kss1-1/Kss1-2 cascade as well as cAMP signaling. These pathways converge toward Prf1, a high mobility group domain TF that recognizes pheromone response elements in regulatory regions of target genes (Figure 5). The promoter of Prf1 itself contains two pheromone response elements, which most likely allows self-regulation of the TF (Figure 5; Kahmann and Kämper, 2004; Brefort et al., 2009).

Regulation of Prf1 also depends on its phosphorylation status (Figure 5), and this TF has six putative MAPK phosphorylation sites (Müller et al., 1999). Mutation of these sites creates an allele that retains its ability to drive basal expression of “a” genes but loses its capacity to induce “b” gene expression following pheromone stimulation. Refined delineation of Prf1 phosphorylation patterns demonstrated that only three MAPK phosphorylation sites are required for Prf1 function (Figure 5; Kaffarnik et al., 2003). Prf1 also contains two PKA phosphorylation sites that are sufficient and essential for “a” gene induction (Figure 5; Kaffarnik et al., 2003). Kss1-1 and PKA both interact with Prf1 in vivo, and kinase assays confirm their ability to phosphorylate this TF. Taken together, these results imply that regulation of “a” genes depends solely on Prf1 phosphorylation by PKA, while...
Figure 5. Mating, Filamentous Growth, and Virulence in *U. maydis*.

In *Ustilago*, several MAPKs regulate mating by acting downstream of the pheromone receptor (see text for details). Along with cAMP signaling, MAPKs are also involved in filamentous growth and virulence. The cAMP pathway comprises heterotrimeric G proteins that function upstream of the adenylate cyclase Uac1. cAMP-dependent PKA holoenzyme works as a tetramer comprising two catalytic subunits (Adr1) and two regulatory subunits (Ubc1). When cAMP level is low, binding of regulatory subunits prevents catalytic activity. At higher cAMP levels, conformational changes allow release of catalytic subunits that enter in the nucleus. MAPK and cAMP signaling converge toward TF Prf1, which controls expression of pheromone- and virulence-induced genes. Hap2 is another MAPK substrate controlling expression of Prf1. In light of this study, previous MAPK names are depicted in parentheses. This figure has been adapted from Nadal et al. (2008).
expression of “b” genes requires phosphorylation by both PKA and MAPKs (Figure 5). Prf1 therefore constitutes a key crosstalk point at which signals derived from both MAPK and cAMP signaling pathways are integrated, with differential phosphorylation status controlling the ability of Prf1 to discriminate between target gene promoters (Figure 5; Kaffarnik et al., 2003).

Hap2 is another TF that was shown to operate downstream of MAPKs in U. maydis. This protein was identified as a Kss1-2–interacting protein but was later found to associate equally well with Kss1-1 (Mendoza-Mendoza et al., 2009). Hap2 regulates Prf1 gene expression by binding to a cis-element called the CCAAT-box (Figure 5). Deletion of Hap2 abolishes mating and results in defective expression of pheromone-responsive genes. Mutation of predicted Hap2 MAPK phosphorylation sites attenuates the pheromone response (Mendoza-Mendoza et al., 2009), suggesting that phosphorylation of this TF is required to fine-tune the expression of Prf1 (Figure 5). Epistasis studies also suggest that more unidentified TFs likely operate downstream of Kss1-1 and Kss1-2 to regulate conjugation tube formation and penetration of the plant cuticle (Figure 5).

The Kss1 MAPK Cascade in M. oryzae

Upon host sensing, many phytopathogenic fungi differentiate specialized structures allowing hyphal penetration of plant tissues. MAPKs have been associated with this process, also referred to as IRM. Typically, IRM is initiated shortly after germination of asexual spores called conidia. Emerging germ tubes grow and differentiate into a dome-shaped appressorium that adheres tightly to the hydrophobic plant surface. Following formation of a penetration peg, the appressorium generates strong turgor pressure that ultimately breaches underlying plant tissues (Figure 6; Talbot, 2003; Ebbole, 2007; Wilson and Talbot, 2009).

One of the most well-studied appressorium-forming fungi is the ascomycete M. oryzae (formerly named Magnaporthe grisea), a hemibiotroph that causes a devastating rice (Oryza sativa) blast disease (Talbot, 2003; Ebbole, 2007; Wilson and Talbot, 2009). In this pathogen, cAMP regulates recognition of the plant surface, a prerequisite for the initiation of IRM. This process also depends on Mor-Kss1 (previously named Pmk1), a MAPK that stimulates appressorium formation (Figure 6). Kss1 knockouts still recognize hydrophobic surfaces and respond to cAMP but fail to form appressoria or produce lesions on susceptible rice plants (Xu and Hamer, 1996). Δkss1 strains also lack the ability to mobilize glycogen and lipid reserves that allow glycerol accumulation and generation of high hydrostatic turgor (Figure 6; Thines et al., 2000). However, it is apparent that Kss1 expression is induced during appressorium formation, and the corresponding protein predominantly accumulates in the appressorium nucleus (Bruno et al., 2004). Mutant strains are also unable to colonize wounded rice tissue, indicating that this MAPK is not only required for early penetration steps but also for infectious growth once inside the plant (Xu and Hamer, 1996).

Based on sequence comparisons with yeast and epistasis studies, the MAP2K Mor-Ste7 (Mst7) and the MAP3K Mor-Ste11 (Mst11) have been positioned upstream of Mor-Kss1 (Figure 6; Zhao et al., 2005). M. oryzae strains lacking ste7 and ste11 fail to form appressoria and are nonpathogenic, while ectopic expression of an active Ste7 allele restores appressorium formation in ste7 and ste11 genetic backgrounds. Moreover, expression of an active Ste7 allele results in phosphorylation of the downstream Kss1 MAPK. On the other hand, appressoria formed by such complemented lines fail to penetrate rice leaves, indicating that active Ste7 can only partially restore defects of ste7 and ste11 mutants (Zhao et al., 2005). Coimmunoprecipitation and bimolecular fluorescence complementation assays also indicate that Ste7 and Kss1 physically interact during appressorium formation and that this interaction depends on a MAPK-docking site located at the N terminus of the MAP2K (Zhao and Xu, 2007).

Y2H assays also reveal that Ste7 and Ste11 interact with Mst50 (Figure 6; Zhao et al., 2005), a homolog of the yeast adaptor protein Ste50 (Figure 1). This ternary association most likely results in the formation of a complex that stabilizes the otherwise weak interaction between Mor-Ste7 and Mor-Ste11 (Zhao et al., 2005). Strains lacking Mst50 are defective in appressorium formation and fail to infect rice (Park et al., 2006). Expression of active Ste7 complements appressorium formation, but not lesion development, in mst50 knockout lines, and coimmunoprecipitation experiments confirm the physical interaction between Ste7, Ste11, and Mst50 (Figure 6; Park et al., 2006).

Adaptor protein Mst50 also interacts with the GTPase Cdc42 (Figure 6; Park et al., 2006), a homolog of Cdc42 from yeast (Figure 1). However, Mor-Cdc42 is unlikely to participate in the activation of the Mor-Kss1 cascade since it is dispensable for appressorium formation and plant infection (Zhao et al., 2007). Two M. oryzae PAKs, Mst20 and CHM1, have also been characterized in search of other candidate working as upstream regulators (Li et al., 2004). Mst20 is dispensable for appressorium formation and plant infection, whereas Δchm1 mutants are only reduced in appressorium formation and still grow invasively on wounded rice leaves. This indicates that pak mutants are phenotypically dissimilar from Δkss1 strains and suggests that Mst20 and CHM1 individually play no critical role in activating the Kss1 pathway from M. oryzae (Figure 6; Li et al., 2004).

Mst50 and Ste11 also interact with GTPases Ras1 and Ras2 (Figure 6; Park et al., 2006). Ras1 knockouts have no defect in appressorium formation or plant infection, whereas lethality hinders characterization of Ras2 knockouts. Nonetheless, expression of an active Ras2 allele stimulates appressorium formation in a wild-type M. oryzae strain, but not in mst50, ste11, ste7, or kss1 mutants, indicating that at least one function of Ras2 is mediated through the Kss1 pathway (Figure 6; Park et al., 2006).

Examination of the Magnaporthe infection process also uncovered Kss1-dependent responses involving the TF Mst12 (Figure 6; Park et al., 2002). Mst12 belongs to a family of proteins that are homologous to Ste12, a well-known substrate of yeast MAPKs (Figure 1). Mutant strains lacking Mst12 are nonpathogenic in rice (Park et al., 2002) and produce classic dome-shaped appressoria that, however, fail to penetrate plant surface. It is thought that Mst12 controls cytoskeleton reorganization associated with penetration peg formation (Park et al., 2004a) and that mutants lacking this TF fail to orient the physical forces exerted by appressorium turgor. mst12 mutants are also compromised in infectious growth since inoculation through wound sites does not
Figure 6. The Pathogenicity Pathway from *M. oryzae*.

Following sensing of host signals, the pathogenicity pathway is activated. Components mediating early steps of signal transduction converge toward a protein complex comprising Mst50 and MAPK signaling components. Following activation by Ste7, the Kss1 MAPK stimulates differentiation of appressoria and controls formation of turgor pressure. Nuclear localization of Kss1 is consistent with the fact that TFs accomplish MAPK function. Study of Mst12 indicates that IRM and infectious growth are separated processes, both required for full pathogenicity. Mst12 is not involved in appressorium formation but functions downstream of Kss1 to control penetration peg formation and proliferation inside host tissues. Sfl1 is another MAPK substrate regulating expression of stress-related genes. In light of this study, previous MAPK names are depicted in parentheses. This figure has been adapted from Park et al. (2006). AP, appressorium; CO, conidia; CU, cuticle; IH, infection hyphae; N, nucleus; PC, plant cell; PP, penetration peg.
allow fungal spreading (Park et al., 2002). The Kss1 cascade from *M. oryzae* thus regulates appressorium differentiation and infectious growth by relying on output responses that are uncoupled, with yet unidentified TF(s) mediating appressorium formation and Mst12 promoting penetration peg formation and invasive development (Figure 6).

In light of the direct phosphorylation of Ste12 by yeast MAPKs (Figure 1), and because corresponding Magnaporthe mutants display partially overlapping phenotypes, it is tempting to hypothesize that there is a direct functional relationship between Mor-Kss1 and Mst12. Interestingly, however, Y2H assays detected only a weak interaction between the two proteins (Park et al., 2002). Mst12 homologs have also been characterized in other phytopathogenic fungi (Rispail and Di Pietro, 2010), and again no direct evidence supports a physical interaction between these TFs and upstream MAPKs. In line with this, all Ste12-like TFs from filamentous fungi lack the protein motif that mediates interaction between Ste12 and yeast MAPKs (Wong Sak Hoi and Dumas, 2010). These observations argue for a conserved function of Ste12-like TFs downstream of MAPKs but also for an indirect relationship involving yet unidentified players that may function as bridging factors for these key signaling components (Figure 6).

In an attempt to identify further Mor-Kss1 substrates, a protein named Sfl1 was recently identified out of a screen for phosphorylated TFs (Figure 6; Li et al., 2011). Sfl1 belongs to the heat shock factor family and contains a putative MAPK docking site as well as three predicted MAPK phosphorylation sites. Coimmunoprecipitation experiments indicate that Kss1 and Sfl1 interact in vivo, and deletion of Sfl1 results in significant reduction of virulence against rice and barley (*Hordeum vulgare*; Li et al., 2011). Mutant strains differentiate appressoria but appear to be defective for invasive growth. sfl1 mutants also show increased sensitivity to elevated temperatures, a finding most likely related to the fact that this TF controls expression of genes encoding heat shock proteins (Figure 6; Li et al., 2011). Screening of a CDNA subtraction library also identified GAS1 and GAS2, two Kss1-regulated genes encoding small proteins required for full appressorium function and virulence (Xue et al., 2002). Comparative analysis of ESTs obtained from wild-type appressoria and kss1 mutant germings also revealed other potential genetic targets of this key MAPK (Ebbole et al., 2004).

**Kss1/Fus3-Type MAPKs in Other Phytopathogenic Fungi**

Several MAPKs regulate morphogenetic transitions associated with plant infection by *U. maydis* and *M. oryzae*. Importantly, these infection-related MAPKs are evolutionarily related to Kss1 and Fus3 (Figure 2), which regulate mating and filamentous growth in *S. cerevisiae* (Figure 1). Homologous Kss1/Fus3-type MAPKs have also been characterized in other fungal plant pathogens, and functional analyses confirm that this class of PKs plays important functions in establishment of various infection strategies.

In the case of investigated pathogens that form well-defined appressoria, including *Cochliobolus heterosporus* (Lev et al., 1999), *Colletotrichum orbiculare* (formerly known as *C. lagenarium*; Takano et al., 2000), and *Pyrenophora teres* (Ruiz-Roldán et al., 2001), Kss1/Fus3-type MAPKs are required for appressorium formation. More recently, the MAP3K MEKK1 (Cor-Ste11 according to our proposed nomenclature) was found to act upstream of the MAPK Cmk1 (here renamed Cor-Kss1) in *C. orbiculare* (Sakaguchi et al., 2010). *C. orbiculare* strains lacking Ste11 and Kss1 both fail to form an appressorium, while nuclear localization of a Kss1–green fluorescent protein fusion is diminished in the ste11 deletion mutants. In *C. heterostrophus*, the MAP3K Ste11 was proposed to work upstream of the MAPK CHK1 (here renamed Che-Kss1), since loss-of-function study revealed that Ste11 and Kss1 are both required for appressorium formation (Izumitsu et al., 2009). In addition, the Che-Kss1 MAPK was previously shown to be required for timely induction of genes encoding cellulolytic enzymes and controlling host tissue penetration (Lev and Horwitz, 2003).

Other phytopathogenic fungi do not form appressoria and thus rely on distinct strategies to colonize host tissues. For instance, penetration of plant aerial parts can occur through stomata that provide direct access to underlying photosynthetic mesophyll tissues. In the wheat (*Triticum aestivum*) pathogen *Mycosphaerella graminicola*, the MAPK gene Fus3 (here renamed Mgr-Kss1) is essential for colonization of the host, and knockout mutants fail to penetrate stomata (Cousin et al., 2006). Based on homology with putative orthologs from *M. oryzae*, additional components of the Mgr-Kss1 pathway have been identified, and complementation assays have emphasized the evolutionary conservation of this important signaling pathway (Kramer et al., 2009). In the cereal pathogen *S. nodorum*, deletion of the MAPK gene Mak2 (here renamed Sno-Kss1) produces strains that achieve penetration of stomata but are still nonpathogenic because of their inability to produce infection structures within the host mesophyll (Solomon et al., 2005).

The activity of Kss1/Fus3-type MAPKs is also associated with virulence of several soil-borne pathogens that cause wilt disease symptoms in a variety of crops. In *F. oxysporum*, deletion of the MAPK gene Fmk1 (Fox-Kss1 in our analysis) results in strains that fail to differentiate penetration hyphae and that display reduced transcript level of the pectate lyase gene *pl*1 (Di Pietro et al., 2001). In *V. dahliae*, disruption of the MAPK gene VMK1 (here renamed Vda-Kss1) also results in strains that have reduced virulence against a variety of host plants (Rauyaree et al., 2005). In the necrotrophic fungus *A. brassicicola*, disruption of the MAPK gene Amk1 (here renamed Abr-Kss1) results in strains that are nonpathogenic on intact plants but that still colonize damaged host tissues (Cho et al., 2007). Unlike Mor-Kss1, this suggests that Abr-Kss1 specifically controls host penetration, whereas subsequent proliferation within the host does not predominantly rely on this MAPK. Accordingly, *Alternaria* mutants lacking this Kss1/Fus3-type MAPK produce lower amounts of hydrolytic enzymes that are usually required for penetration of host tissues (Cho et al., 2007). In *B. cinerea*, inactivation of the MAPK gene BMP1 (here renamed Bci-Kss1) results in strains that are essentially nonpathogenic, since infection hyphae fail to penetrate and macerate plant tissues (Zheng et al., 2000; Doehlemann et al., 2006). Independent studies also revealed that deletion of the MAPK gene MAP1Gpmk1 (here renamed Fgr-Kss1) hinders pathogenicity of *Fusarium graminearum*, the causal agent of wheat head-blight disease (Jenczmiionka et al., 2003; Urban et al., 2003). Again, pathogenicity of Fgr-kss1
Recently, it was shown that, upon plant infection, the appressorium wall during host infection (Xu et al., 1998). Mor-Slt2 controls overall cell wall strength as well as remodeling osmotic stabilization. Taken together, these results suggest that colonies grown on plates undergo autolysis in the absence of hypersensitive to cell wall pathogenic since, while they differentiate appressoria, these fail (see Supplemental Data Set 2 online). Mor-Slt2 mutants are non-pathogenic in a murine model system (Ortoneda et al., 2004). These are the MADS box TF Mig1 and the APSES TF Swi6, the respective homologs of Slt2 substrates Rlm1 and Swi6 in yeast (Figure 1). Disruption of Mig1 results in the differentiation of normal appressoria, penetration pegs, and primary infectious hyphae. However, mig1 mutants fail to infect rice tissues, since they are blocked in the differentiation of secondary infection structures (Mehrai et al., 2008). In the case of Mor-Swi6, deletion mutants display reduced hyphal growth, abnormal formation of conidia and appressoria, as well as altered appressorium function and pathogenicity (Qi et al., 2012). Although it remains to be established whether Mor-Slt2 can phosphorylate Mig1 and Mor-Swi6, these results suggest that several TFs function downstream of this MAPK to promote pathogenicity and sustain infectious growth in planta.

The cell wall integrity signaling pathway from U. maydis was also recently resolved and shown to involve MAP3K Bck1, MAP2K Mkk1, and MAPK Mpk1 (here renamed Uma-Slt2-1; Carbó and Pérez-Martin, 2010). In contrast with yeast, where activation of the cell wall integrity pathway arrests the cell cycle in the G2 phase, activation of the corresponding pathway in U. maydis forces cells to escape the G2 phase. This effect is correlated with a decrease in CDK inhibitory phosphorylation, which itself depends on the nuclear accumulation of the mitotic protein phosphatase Cdc25 following activation of the Slt2-1 MAPK (Carbó and Pérez-Martin, 2010). The authors proposed that cell cycle adaptation to stress most likely evolved differently in divergent fungi, so that each species can move toward a cell cycle phase that is most appropriate for responding to the environmental signals encountered.

While the virulence function of the Ustilago cell wall integrity signaling pathway still remains to be established, Slt2 homologs from other plant pathogenic fungi have been linked to pathogenicity. In C. orbi
culare, the MAPK MAF1 (here renamed Cor-Slt2) regulates early penetration steps, and deletion mutants produce elongated germ tubes that lack an appressorium (Kojima et al., 2002). In C. purpurea, the Slt2 homolog (previously known as MK2) is necessary for host penetration and corresponding deletion mutants retain only a limited ability to colonize plant tissues (Mey et al., 2002a). In B. cinerea, bmp3 (here renamed Bci-slt2) mutants are defective in surface sensing, plant omics tend to be localized close to the outer layer of the cell wall, farther from the fungal cell membrane (Fujikawa et al., 2009). Such uneven distribution could favor masking of chitin and β-1,3-glucans, which are highly sensitive to plant hydrolases. Interestingly, accumulation of reinforcing α-1,3-glucans does not occur in Mor-slt2 knockout strains, a situation possibly reflecting one way in which this MAPK regulates strengthening of the fungal cell wall during plant colonization (Fujikawa et al., 2009).

Our understanding of the cell wall integrity pathway from M. oryzae was also expanded by the characterization of MCK1, a MAP3K that we suggest renaming Mor-Bck1 as it displays high homology to Bck1 from yeast. Following epistasis study, Mor-Bck1 was suggested to function upstream of Mor-Slt2 to ensure proper appressorium function (Jeon et al., 2008). Two nuclear localized proteins from M. oryzae were also found to interact with Slt2 and are believed to act as key downstream substrates mediating MAPK function (Mehrai et al., 2008; Qi et al., 2012). These are the MADS box TF Mig1 and the APSES TF Swi6, the respective homologs of Slt2 substrates Rlm1 and Swi6 in yeast (Figure 1). Disruption of Mig1 results in the differentiation of normal appressoria, penetration pegs, and primary infectious hyphae. However, mig1 mutants fail to infect rice tissues, since they are blocked in the differentiation of secondary infection structures (Mehrai et al., 2008). In the case of Mor-Swi6, deletion mutants display reduced hyphal growth, abnormal formation of conidia and appressoria, as well as altered appressorium function and pathogenicity (Qi et al., 2012). Although it remains to be established whether Mor-Slt2 can phosphorylate Mig1 and Mor-Swi6, these results suggest that several TFs function downstream of this MAPK to promote pathogenicity and sustain infectious growth in planta.

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penetration, and induction of necrotic lesions (Rui and Hahn, 2007). In *M. graminicola*, *Sit2* mutants show normal penetration of wheat stomata but are highly reduced in virulence because infectious hyphae fail to branch out (Mehrabi et al., 2006a). Finally, *MGVT* (here renamed *Fgr-Sit2*) was shown to control heterokaryon formation and accumulation of trichothecene mycotoxins in *F. graminearum* (Hou et al., 2002). In light of these various findings, the study of *Sit2*-type MAPKs has defined a second fungal MAPK cascade that plays an evolutionarily conserved role in virulence against plants.

While the role of *Sit2*-type MAPKs in fungal pathogenesis seems to be broadly conserved, their function in preserving cell wall integrity seems to vary among fungal pathogens. As seen for the *Mor-sit2* mutants, *Cpu-sit2*, *Fgr-sit2*, *Mgr-sit2*, and *Una-sit2* mutants all have weakened cell walls and show increased sensitivity to cell wall-degrading enzymes and to compounds interfering with cell wall biosynthesis (Hou et al., 2002; Mey et al., 2002a; Mehrabi et al., 2006a; Carbó and Pérez-Martín, 2010). By contrast, deletion of *Sit2* homologs from *C. orbiculare* and *B. cinerea* has no obvious effect on cell wall sensitivity to lytic enzymes and inhibitors (Kojima et al., 2002; Rui and Hahn, 2007). The reasons for this discrepancy may be related to the possibility that not all *Sit2* homologs have retained their function in cell wall integrity or that a functionally redundant pathway controls this response in some, but not all, phytopathogenic fungi.

**Hog1-Type MAPKs in Phytopathogenic Fungi**

In *S. cerevisiae*, Hog1 lies downstream of a branched signaling pathway that responds to high osmolarity conditions (Figure 1). However, other deleterious environmental conditions also activate the HOG pathway (Hayashi and Maeda, 2006; Marques et al., 2006; Panadero et al., 2006). Consistent with this, Hog1 homologs carry a -TGY- phosphorylation motif, which is a hallmark of stress-activated MAPKs in fungal and mammalian systems.

The *M. oryzae* MAPK OSM1 (here renamed Mor-Hog1) was the first Hog1-type MAPK to be characterized from phytopathogenic fungi (Dixon et al., 1999). Deletion mutants are highly sensitive to osmotic stress and show severe morphological defects when grown under hyperosmotic conditions. These phenotypes are correlated with reduced ability of mutant strains to accumulate osmoprotectant molecules in the stressed mycelium. On the other hand, strains lacking Mor-Hog1 generate normal appressorial turgor since they can still accumulate glycerol in their appressoria. More than 20 years ago, *M. oryzae* containing a -TXY- motif, the hallmark of the MAPK protein family, was identified as a gene expressed exclusively during meiosis (Smith and Mitchell, 1989; Yoshida et al., 1990). It was later identified as a regulatory kinase that controls entry and progression of the meiotic cell cycle via the destabilization of its transcriptional activator, Ime1 (Guttmann-Raviv et al., 2002). Following in-depth characterization of its function, Ime2 is now considered as a key node controlling various steps of meiosis, including ascospore formation, pseudohyphal growth, and sexual reproduction in response to light and nutrient deprivation (Irigeri, 2011).

To date, all identified Ime2 homologs share conserved topology, including an N-terminal region that contains a PK domain displaying high homology to the CDK proteins. The kinase domain of Ime2-type proteins also harbors an activation loop containing a -TXY- motif, the hallmark of the MAPK protein family. Until recently, Ime2 homologs had not been defined as required for virulence of some, but not all, plant pathogens. The reasons for this difference remain unclear but could be related to the disparity of fungal invasion strategies and/or ability to cope with host counterdefenses (Zhao et al., 2007). While deletion of Hog1-type MAPKs has variable and species-specific consequences for fungal virulence, the effect on resistance to stress appears more universal. For instance, hypersensitivity to osmotic stress is not only reported for the *Mor-hog1* mutants (Dixon et al., 1999) but also for mutants of *Cryphonectria parasitica*, *C. orbiculare*, *B. oryzae*, *M. graminicola*, and *B. cinerea* lacking their respective Hog1 homologs (Kojima et al., 2004; Park et al., 2004b; Mehrabi et al., 2006b; Moriwaki et al., 2006; Segmüller et al., 2007). In filamentous fungi, reports have also linked the activity of group III His kinases to the phosphorylation status of Hog1-type MAPKs. For instance, the *C. heterostrophus* His kinase Dic1p promotes stress-mediated activation of Che-Hog1 (Yoshimi et al., 2005), while deletion of His kinase gene HS1 compromises Aal-Hog1 activation in response to stress in *Alternaria alternata* (Lin and Chung, 2010). These studies provide the first hints of how upstream regulation of the HOG pathway might be organized in phytopathogenic fungi.

In several fungal species, it has also been reported that mutants blocked in components of the HOG pathway are more resistant to a variety of fungicides (Kojima et al., 2004; Motoyama et al., 2005, 2008; Yoshimi et al., 2005; Mehrabi et al., 2006b; Moriwaki et al., 2006; Vlaud et al., 2006). In at least three cases, treatment with fungicides also results in the activation of Hog1-type MAPKs (Kojima et al., 2004; Yoshimi et al., 2005; Segmüller et al., 2007). This suggests that the fungicidal properties of several compounds may be due to an overstimulation of the HOG pathway, leading to uncontrolled accumulation of osmoprotectant molecules and concomitant swelling of fungal cells.

**Ime2 Homologs: A Novel Subfamily of MAPKs Conserved in All Eukaryotes?**

More than 20 years ago, *S. cerevisiae* Inducer of Meiosis2 (*Ime2*) was identified as a gene expressed exclusively during meiosis (Smith and Mitchell, 1989; Yoshida et al., 1990). It was later shown that Ime2 is required for entry and progression of the meiotic cell cycle via the destabilization of its transcriptional activator, Ime1 (Guttmann-Raviv et al., 2002). Following in-depth characterization of its function, Ime2 is now considered as a key node controlling various steps of meiosis, including ascospore formation, pseudohyphal growth, and sexual reproduction in response to light and nutrient deprivation (Irigeri, 2011).

To date, all identified Ime2 homologs share conserved topology, including an N-terminal region that contains a PK domain displaying high homology to the CDK proteins. The kinase domain of Ime2-type proteins also harbors an activation loop containing a -TXY- motif, the hallmark of the MAPK protein family. Until recently, Ime2 homologs had not been defined as...
true MAPKs, even though dual phosphorylation of the -TAY-motif found in yeast Ime2 is required for its activity (Schindler et al., 2003). In fact, there is still no evidence that MAP2Ks from yeast can activate Ime2, and modification of the Ime2 -TAY-motif is associated with both autophosphorylation and CDK-activating kinase1 activities (Schindler et al., 2003; Schindler and Winter, 2006). On the other hand, characterization of Cdk-related kinase1, an Ime2 homolog from U. maydis that was here renamed Uma-Ime2, now suggests that at least certain Ime2 homologs behave like true MAPKs and that Ime2-like proteins may indeed constitute a novel subfamily of MAPKs (Figure 2; Garrido et al., 2004).

In Ustilago, Ime2 promotes morphogenetic transition through the control of Prf1 expression (Figure 5; Garrido and Pérez-Martín, 2003; Garrido et al., 2004). As a result, strains lacking this PK are impaired in mating and formation of dikaryotic filaments because hyphal fusion between opposite mating types fails to occur (Garrido et al., 2004). Since mating of Ustilago haploid cells is a prerequisite for plant infection, Ime2 is also necessary for pathogenicity, and mutant strains cause drastically reduced numbers of tumors compared with wild-type strains. Interestingly, the MAP2K Ste7, which stimulates mating and virulence by operating upstream of MAPKs Kss1-1 and Kss1-2, also targets the -TXY- motif of Ime2 for phosphorylation (Figure 5; Garrido et al., 2004). In addition, Ime2 physically interacts with Kss1-1, and its full activity is dependent on this MAPK. This suggests that in U. maydis, activation of Ime2 is tightly linked with signaling that goes through the Kss1-1 MAPK module and that this PK exerts its function both in parallel to Kss1-1 and as a substrate for this MAPK (Figure 5; Garrido et al., 2004; Imriger, 2011).

Ime2 homologs are not only conserved in fungi but also in all eukaryotic taxa examined (Krylov et al., 2003). In mammals, for instance, the Ime2 homolog, male germ cell-associated kinase (Mak), has an expression pattern that correlates with sexual development (Jinno et al., 1993). In Arabidopsis, three Mak-homologous kinases show overall sequence similarity to MAPKs and possess a -TEY- phosphorylation motif that is typically associated with extracellular signal-regulated kinase-type MAPKs (Tena et al., 2001; Champion et al., 2004). In the original plant MAPKs survey, Arabidopsis Mak-homologous kinases were not classified as true MAPKs because they show sequence relatedness to both MAPKs and CDC2-like PKs (Ichimura, 2002). In view of the novel data now linking Ste7 and Ime2 from U. maydis, it will be important to investigate whether MAP2Ks can activate Ime2-type proteins in other eukaryotic taxa, including in plant and animal systems. If this were indeed the case, this exciting new feature would certainly argue for inclusion of the Ime2-like kinases as bona fide MAPKs.

**Molecular Dialogue Between Plants and Fungal Pathogens**

Upon host sensing, fungal MAPKs are important for the virulence of phytopathogens because they control mating, IRM, reinforcement of the fungal cell wall, and expression of virulence factors (Zhao et al., 2007; Rispail et al., 2009; this article). Using mechanical pressure and/or enzymatic activity, fungal pathogens breach the plant surface in order to access desired nutrients (Figure 7, top left). However, during fungal infection of plants, MAPK signaling is critical not only for the attacking fungi but also for activation of defense responses in plant cells.

**Activation of MAPK Cascades in the Infected Plants**

Plant MAPK cascades lie downstream of a sensitive surveillance system that monitors threats of infection (Figure 7). The plant surveillance apparatus includes pattern recognition receptors (PRRs) that allow detection of molecular signatures also known as microbe-associated molecular patterns (MAMPs; Bolier and Felix, 2009). For instance, the extracellular portion of LysM receptors possesses chitin binding activity (Figure 7; Iizasa et al., 2010; Petutschlig et al., 2010), and LysM receptors are necessary for chitin-mediated activation of defense in Arabidopsis (Miya et al., 2007; Wan et al., 2008) and rice (Kaku et al., 2006). Receptor-like kinases harboring extracellular Leu-rich repeats have also been linked to the detection of microbes (Zipfel et al., 2004, 2006), and members from this large PK family likely participate in detection of several fungal MAMPs (Figure 7). Endogenous molecules referred to as damage-associated molecular patterns (DAMPs) can also induce plant defense following their release from damaged plant cell walls (Figure 7). Wall-associated kinases have recently been identified as DAMP receptors (Kohorn et al., 2009; Brutus et al., 2010), and these, together with signals from other PRRs, promote activation of several signaling pathways regulating plant basal defense (Figure 7).

In Arabidopsis, two complete MAPK cascades have been shown to function downstream of PRRs (Figure 7; Asai et al., 2002; Gao et al., 2008; Qiu et al., 2008b). Perception of elicitors thus leads to the activation of at least three stress-responsive MAPKs: MPK3, MPK4, and MPK6 (Asai et al., 2002; Wan et al., 2004; Mészáros et al., 2006; Denoux et al., 2008; Galletti et al., 2011). Further study of plant MAPKs has uncovered several output responses that contribute to plant defense activation. For example, At-MPK3 and At-MPK6 were shown to promote stabilization of certain isoforms of 1-aminocyclopropane-1-carboxylic acid synthase, a key step in Botrytis-induced production of ethylene (Figure 7; Liu and Zhang, 2004; Han et al., 2010). Large-scale in vitro analysis also indicates that TFs are well represented among the primary targets of stress-responsive MAPKs (Feilner et al., 2005; Popescu et al., 2009). In planta, MAPKs have been shown to alter the function of several TFs (Figure 7), including WRKYs in Arabidopsis and Nicotiana (Andreasson et al., 2005; Ishihara et al., 2011; Mao et al., 2011), the bZIP At-VP1 (Djamei et al., 2007), ethylene-related proteins At-EIN3 (Yoo et al., 2008) and At-ERF104 (Bethke et al., 2009), as well as Pti-ZFP1, a poplar (Populus trichocarpa) Zn finger protein (Hamel et al., 2011).

Through posttranslational modification of TFs, plant MAPKs regulate expression of target genes, including several that are involved in signal transduction and defense (Figure 7). Notably, MPK3/6 were shown to promote synthesis of camalexin, the main antimicrobial phytoalexin in Arabidopsis (Ren et al., 2008). Epistasis analysis places both MAPKs upstream of Phytoalexin Deficient2/3 (PAD2/3), two camalexin biosynthesis genes. Moreover, ectopic expression of active upstream MAP2K or MAP3K induces camalexin biosynthesis in the absence of pathogen attack. In wild-type plants, Botrytis-induced production of camalexin is
Fungal pathogens perceive plant-derived signals using plasma membrane receptors. This leads to the activation of fungal MAPK cascades that modulate TF activity and promote expression of genetic targets. Among induced genes are those associated with mating, IRM, mycotoxin biosynthesis, and degradation of the plant cell wall. On the other hand, plant receptors perceive molecular signatures associated with fungal structures or activity. This results in the activation of plant signaling pathways, including modified Ca²⁺ homeostasis, oxidative burst, and changes in MAPK phosphorylation status. Plant MAPKs promote biosynthesis of stress hormone and modulate gene expression through the phosphorylation of TFs. Output responses, including cell wall-degrading enzymes and antimicrobial compounds, affect fungal cell integrity and threaten pathogen survival. Fungal MAPK cascades also participate in compensatory responses allowing protection of hyphae against plant defenses. Overall, MAPK cascades can be viewed as conserved signaling modules involved in the molecular dialogue between fungal pathogens and plants. AP, appressorium; CO, conidia; CU, cuticle; CW, cell wall; CY, cytoplasm; GT, germ tube; N, nucleus; NM, nuclear membrane; PM, plasma membrane; PP, penetration peg; ROS, reactive oxygen species.

preceded by MPK3/6 activation, and the production of this secondary metabolite is compromised in mpk3 and mpk6 knockout mutants (Ren et al., 2008).

More recently, it was shown that WRKY33 is required for MPK3/6-mediated induction of camalexin biosynthesis in Arabidopsis (Mao et al., 2011). Following pathogen attack, wrky33 knockout plants are compromised in the induction of camalexin biosynthesis genes as well as in production of the metabolite. In vivo analysis indicates that WRKY33 is a direct substrate of MPK3/6, and complementation assays show that phosphorylation is involved in WRKY33-mediated induction of camalexin biosynthesis (Mao et al., 2011). Interestingly, WRKY33 was also identified as
part of a multiprotein complex that includes MPK4 and one of its substrates, MKS1 (Andreason et al., 2005). In the absence of pathogen, the MPK4-MKS1-WRKY33 ternary complex is stabilized, preventing WRKY33 from activating target gene promoters (Qiu et al., 2008a). Following pathogen perception, MPK4 is activated, resulting in MKS1 phosphorylation and dissolution of the protein complex (Figure 7). Released WRKY33 can then bind its genetic targets, including the promoter of PAD3 (Qiu et al., 2008a).

Fungal MAPK Cascades and Compensatory Response against Host Defenses

Camalexin produced by the challenged plant probably exerts its antifungal activity by causing cell membrane damage in the attacking fungi (Figure 7). In the necrotroph A. brassicicola, the cellular response elicited by such damage requires both the cell wall integrity and HOG pathways, leading to the activation of fungal MAPKs Abr-Stt2 and Abr-Hog1, respectively (Joubert et al., 2011). Alternaria strains lacking these MAPKs are not only hypersensitive to camalexin, but also to brassinin, a structurally related phytoalexin from Brassica species (Joubert et al., 2011). In F. graminearum, two MAPKs, Fgr-Kss1 and Fgr-Stt2, appear to play a major role in fungal sensitivity to certain plant defenses, a small family of pathogenesis-related proteins (Figure 7; Ramamoorthy et al., 2007). Taken as a whole, these findings suggest that fungal MAPKs are not only required for plant infection but also for regulation of compensatory responses that help to preserve fungal cell integrity during exposure to plant defenses. As a result, fungal strains lacking MAPK signaling components may exhibit reduced virulence because of their inability to cope with plant defenses.

Specificity of Mutually Activated MAPK Pathways

Many features associated with fungal pathogenicity and plant defense are dependent on signaling through MAPK cascades (Figure 7). As a result, these modules can be viewed as crucial players that promote the two-way molecular dialogue between pathogenic fungi and plants. On the other hand, it is not clear whether such patterns of mutual activation are necessarily specific to MAPK modules or if other shared eukaryotic signaling pathways exhibit such a bilateral contribution to the highly complex molecular dialogue between both protagonists.

Part of the answer may come from the comparison of conserved eukaryotic signaling pathways that have been characterized in both plants and fungi. In mammalian and fungal systems, for instance, cyclic nucleotides are viewed as central secondary messengers that promote activation of conserved PKA/PKG/PKC (AGC) signaling pathways. In phytopathogenic fungi like U. maydis, perception of host-derived signals leads to an increase in the cellular concentrations of cAMP, resulting in the activation of PKA signaling and associated promotion of virulence (Figure 5; Kahmann and Kämper, 2004; Brefort et al., 2009). Modulation of cAMP levels also results in altered plant cell signaling, although plant genomes do not appear to encode prototypical PKA enzymes (Robert and Offringa, 2008). Of the 37 AGC kinases identified in Arabidopsis (Galván-Ampudia and Offringa, 2007), none has so far been directly involved in MAMP-triggered immunity. In fact, the majority of plant AGC kinases belong to a plant-specific subgroup, called the AGCVIII kinases, members of which are mainly involved in auxin signaling (Galván-Ampudia and Offringa, 2007; Robert and Offringa, 2008) and other developmental processes (Zhang and McCormick, 2009). Therefore, despite their wide distribution in eukaryotes, including plants and fungi, it seems unlikely that AGC kinases contribute directly to the molecular dialogue between plants and pathogenic fungi, since the plant homologs do not actively participate in early-induced plant immunity.

Proteins allowing perception of calcium (Ca2+), another ubiquitous secondary messenger, are also likely candidates as crucial contributors to the two-way molecular dialogue between plants and interacting fungi. In animals and fungi, intracellular Ca2+ sensors include calmodulins (CaMs), which interact with calcineurin B (CNB) upon Ca2+ stimulation. These protein complexes then bind to calcineurin A, a highly conserved protein phosphatase (Luan, 2009). In phytopathogenic fungi, CaMs and CNBs have been involved in virulence (Lee and Lee, 1998; Warwar et al., 2000; Uhm et al., 2003; Harel et al., 2006; Ahn and Suh, 2007; Egan et al., 2009; Cervantes-Chávez et al., 2011), and comparative genomics of the Ca2+/CaM/CNB pathway suggest that it is highly conserved in fungi (Rispail et al., 2009).

In plants, Ca2+ signaling also depends on CaMs, as well as a group of CNB-related proteins, referred to as the CNB-like (CBL) proteins (Batistic and Kudla, 2009; Tena et al., 2011). Like their animals and fungi counterparts, CBLs were anticipated to act upstream of a protein phosphatase, but instead they were found to form a complex regulatory network with members of the CBL-interacting PK (CIPK) family (Luan, 2009). In plants, CBL–CIPK interactions regulate various stress responses and represent a fundamental paradigm shift that distinguishes signaling in animals and fungi from that in plants (Luan, 2009).

The Ca2+ signaling machinery of plants also possesses an exclusive feature related to the evolution and expansion of the calcium-dependent PK family, a group of plant-specific proteins that combine unique Ca2+ sensing and signaling capabilities within the same gene product (Batistic and Kudla, 2009). Several members of the calcium-dependent PK family are involved in stress responses, including defense gene upregulation in response to MAMPs (Tena et al., 2011). As a result, while the Ca2+ signaling pathways from plants and interacting fungi share some components, striking differences exist in pathway architecture, regulation, and complexity.

Although the full range of signaling strategies operating in plants and fungi has not been defined, comparison of other characterized pathways suggests that very few of these strategies display the striking level of architectural conservation (from membrane receptors to nuclear TFs) that typifies the MAPK pathways. Bilateral activation of MAPK modules from interacting plants and pathogenic fungi may therefore possess a certain level of specificity compared with other signaling pathways, possibly as a consequence of the outstanding MAPK versatility with regards to environmental responsiveness.

UPCOMING CHALLENGES AND CONCLUDING REMARKS

Completion of whole-genome sequencing projects marks the entry of fungal MAPK research in the postgenomic era. Many
questions regarding fungal MAPK signaling remain unanswered, and upcoming studies should, for instance, focus on the understanding of upstream mechanisms leading to MAPK activation. Recently, homologs of the yeast osmosensors Sho1 and Msb2 (Figure 1) have been shown to act upstream of pathogenesis-related MAPK cascades in U. maydis (Figure 5; Lanver et al., 2010), M. oryzae (Figure 6; Liu et al., 2011), and F. oxysporum (Pérez-Nadales and Di Pietro, 2011). Along with the previously identified G protein–coupled receptor Pth11 from M. oryzae (Figure 6; DeZwaan et al., 1999; Kulkarni et al., 2003), these candidate receptors likely ensure MAPK cascade activation following recognition of specific ligands. However, identity of the full array of signaling components lying between surface receptors and MAPK modules is still missing. Comparative and functional genomics should also facilitate characterization of less-well-defined MAPK pathways, including those that engage the Slt2- and Hog1-type MAPKs (Rispail et al., 2009). While upstream regulation of Slt2-type MAPK cascades remains largely unexplored in phytopathogenic fungi, group III His kinases have been shown to operate upstream of Hog1-type MAPKs (Yoshimi et al., 2005; Lin and Chung, 2010). With regards to fungal pathogenicity against plants, complete characterization of all the MAPK signaling pathways is crucial, since genetic evidence points to extensive crosstalk between parallel MAPK cascades operating in C. orbiculare (Takano et al., 2000; Kojima et al., 2002), M. oryzae (Xu and Hamer, 1996; Dixon et al., 1999), and C. heterostrophus (Elaihui et al., 2007; Igbaria et al., 2008).

Comparison of genomic sequences also highlights new challenges regarding the study of MAPK signaling in basidiomycetes. For instance, duplication of several MAPK genes raises the question of the extent of functional redundancy between paralogous MAPKs. In U. maydis, Kss1-1 and Kss1-2 exhibit partially redundant functions during mating and virulence (Zhao et al., 2007; Brefort et al., 2009; Rispail et al., 2009), but careful study of single deletion mutants also shows that these MAPKs still display some degree of specialization. So far, Ustilago is the only species in which functional redundancy between paralogous MAPKs has been addressed experimentally, and the situation may be even more complex in other fungi, where up to three paralogs can be identified in a single MAPK clade. Fine-grained approaches, including promoter-reporter fusions and functional complementation of double and triple knockout mutants, will be needed to decipher discrete functions of such closely related MAPKs. Downstream of the cascade, paralogous MAPKs may also possess unique and shared substrates, and identification of these targets will be a prerequisite to fully understand the output responses controlled by each MAPK. Currently, only a few TFs have been linked to MAPK signaling, and characterization of deletion mutants suggests that more MAPK substrates await discovery. In addition to TFs, MAPKs from other eukaryotic taxa are well known for their propensity to phosphorylate cytosolic as well as microtubule–associated proteins. This pattern has yet to be confirmed in phytopathogenic fungi, even though new potential interacting partners have recently been identified (Zhang et al., 2011).

It is paradoxical that, while tripartite MAPK modules have been conserved throughout evolution of all eukaryotes, these control an extraordinary array of responses that are specific to each species. How such an ancient and evolutionarily conserved mechanism can regulate so many output responses is a fascinating, yet unanswered, question. Study of the interactions between plants and fungal pathogens exemplifies the plasticity of MAPK cascades operating simultaneously in both protagonists. MAPK phosphorylations are not only conserved in phytopathogenic fungi but also in other eukaryotes that interact with plants, including mycorrhizal fungi, oomycetes, insects, and nematodes. Relying on different strategies, these varied organisms positively or negatively affect plant fitness, employing mechanisms that may often rely on well-conserved MAPK signaling pathways. In line with this, it was recently shown that an oomycete MAPK is required for virulence of Phytophthora sojae against soybean (Glycine max; Li et al., 2010). Activation of plant MAPKs also occurs following interactions with other eukaryotes, including following perception of oomycete elicitors (Kro et al., 2003) and feeding by herbivorous insects (Kandoth et al., 2007; Wu et al., 2007). Taken together, these reports strongly suggest that MAPK cascades are not only required for the molecular dialogue between plants and pathogenic fungi but also for the exchanges occurring between plants and eukaryotes from other taxa.

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Table 1.** List of the Genomic Databases Used in This Study.

**Supplemental Data Set 1.** Origin and Gene Model Number of MAPKs from Plant-Interacting Fungi.

**Supplemental Data Set 2.** Origin and Gene Model Number of MAP2Ks from Plant-Interacting Fungi.

**Supplemental Data Set 3.** Origin and Gene Model Number of MAP3Ks from Plant-Interacting Fungi.

**Supplemental Data Set 4.** Amino Acid Alignment Used to Produce Phylogeny of Fungal MAPKs Shown in Figure 2.

**Supplemental Data Set 5.** Amino Acid Alignment Used to Produce Phylogeny of Fungal MAP2Ks shown in Figure 3.

**Supplemental Data Set 6.** Amino Acid Alignment Used to Produce Phylogeny of Fungal MAP3Ks shown in Figure 4.

**Supplemental References 1.** List of the Literature Cited in the Supplemental Data Sets.

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**AUTHOR CONTRIBUTIONS**

L.-P.H. conducted in silico search of gene models and conceived the section describing MAPK signaling components from symbiotic fungi. S.D. and B.E.E. critically revised the article and provided important intellectual content. All authors read, helped to edit, and approved the final version of the article.


MAPK Cascades in Plant-Interacting Fungi

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Mitogen-Activated Protein Kinase Signaling in Plant-Interacting Fungi: Distinct Messages from Conserved Messengers
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