Regulation of Pathogenesis by Light in *Cercospora zeae-maydis*: An Updated Perspective

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The fungal genus *Cercospora* is one of the most ubiquitous groups of plant pathogenic fungi, and gray leaf spot caused by *C. zeae-maydis* is one of the most widespread and damaging foliar diseases of maize in the world. While light has been implicated as a critical environmental regulator of pathogenesis in *C. zeae-maydis*, the relationship between light and the development of disease is not fully understood. Recent discoveries have provided new insights into how light influences pathogenesis and morphogenesis in *C. zeae-maydis*, particularly at the molecular level. This review is focused on integrating old and new information to provide an updated perspective of how light influences pathogenesis, and provides a working model to explain some of the underlying molecular mechanisms. Ultimately, a thorough molecular-level understanding of how light regulates pathogenesis will augment efforts to manage gray leaf spot by improving host resistance and disease management strategies.

**Keywords**: *Cercospora*, photoreceptor, stomatal tropism

*Cercospora* is one of the world’s most ubiquitous genera of plant pathogenic fungi

The fungal genus *Cercospora* is one of the most diverse and destructive groups of foliar fungal pathogens in the world, comprised of over 3000 named species (Crous and Braun, 2003). *Cercospora* species infect a wide range of important crops and can cause tremendous losses (Mian et al., 2008; Ward et al., 1999; Weiland and Koch, 2004). *Cercospora* species are frequently classified according to host association, e.g., *C. beticola* infects sugar beet (*Beta vulgaris*). Although *Cercospora* species generally share many important morphological and biochemical characteristics, the host range of a given species is quite narrow. The fact that such a large number of *Cercospora* species share numerous physiological characteristics and are closely related at the molecular level (Goodwin et al., 2001), yet exhibit such a high degree of host specificity, seems to indicate that an early progenitor of the taxon evolved a highly effective strategy for foliar pathogenesis that further evolved to overcome defense responses in a wide range of plants.

*Cercospora zeae-maydis* causes gray leaf spot of maize

First identified in Illinois in 1924 (Tehon and Daniels, 1925), gray leaf spot did not become a major concern in the U.S. until the 1980s, when a dramatic increase in both incidence and severity was observed (Latterell and Rossi, 1983). In the 1990s, gray leaf spot caused significant losses in the Midwest and, as an endemic pathogen, is now commonly encountered on an annual basis in most maize-producing regions of the U.S. (Ward et al., 1999). Additionally, during the 1980s and 1990s, the disease also became pandemic throughout Africa and South America, with reported losses of 60% or greater (Ward et al., 1999). In the U.S., two genetically distinct groups of the pathogen were found to cause identical gray leaf spot symptoms, and were designated Groups I and II (Wang et al., 1998). Later, the two groups were determined to be taxonomically distinct species; Group I retained the species name of *C. zeae-maydis*, whereas Group II was designated *C. zeina* (Crous et al., 2006). Interestingly, *C. zeina* appears to be the predominant (if not exclusive) gray leaf spot pathogen in Africa (Dunkle and Levy, 2000; Meisel et al., 2009). Both fungi causes distinctive rectangular lesions delineated by major veins of the leaf, and the disease derives its name from the grayish appearance of lesions when masses of conidia are produced on erumpent conidiophores emerging from stomata. When disease pressure is high immediately before or during grain fill, the loss of photosynthetic capacity can substantially reduce yield and predispose plants to other pathogens, such as stalk-rotting fungi. In general, gray leaf spot is most severe in warm, humid conditions; free moisture is critical for disease development (Rupe et al., 1982).
The disease cycle of gray leaf spot can be divided into four distinct stages (Fig. 1).

1. **Germination** (0–3 days after inoculation [dai]). Asexual spores (conidia) are dispersed onto leaves by wind or rain splash. Under humid conditions, nascent hyphae (germ tubes) emerge and grow across the leaf surface. In the vicinity of a stomate, germ tubes reorient growth to intercept stomata.

2. **Infection** (3–5 dai). Upon reaching stomata, hyphae differentiate into bulbous, multilobed appressoria. Unlike appressoria produced by *Magnaporthe grisea* and other fungi that penetrate the epidermis by generating turgor pressure, those produced by *C. zeae-maydis* appear to function as scaffolds to guide the penetration peg into the stomatal pore. Unlike appressoria produced by *Magnaporthe grisea* and other fungi that penetrate the epidermis by generating turgor pressure, those produced by *C. zeae-maydis* appear to function as scaffolds to guide the penetration peg into the stomatal pore.

3. **Colonization** (5+ dai). After entering the leaf mesophyll via stomata, *C. zeae-maydis* grows intercellularly before switching to a necrotrophic growth habit. The necrotic, expanding lesions induced by *C. zeae-maydis* are delimited by the major veins of the maize leaf, giving them a distinctive rectangular shape.

4. **Sporulation** (7+ dai). Shortly after lesions form, erumpent conidiophores emerge through stomata of colonized tissue and produce conidia. Often, conidia are produced in a zonate pattern, giving lesions a scalariform (ladder-like) appearance. Conidia are dispersed by wind or water to neighboring plants, thus initiating secondary infection cycles.

Gray leaf spot is difficult to manage because many maize hybrids are moderately to highly susceptible to the disease, the fungus overwinters in plant debris, and until recently, chemical control options were not cost-effective in most regions of the world. Given the increasing reliance on conservation tillage in maize production systems throughout the world and the demonstrated ability of *Cercospora* species to evolve resistance to numerous fungicide chemistries (Hanson, 2010), the long-term outlook calls for an increased reliance on genetic resistance to ensure effective and sustainable disease management. However, considerably more information is required about the molecular basis of gray leaf spot in order to optimize efforts to improve genetic resistance through conventional breeding or transgenic approaches.

**Light regulates secondary metabolism and morphogenesis in *C. zeae-maydis***

During pathogenesis, *C. zeae-maydis* and many other species of *Cercospora* produce the non-host-specific phytotoxin cercosporin (Daub and Ehrenshaft, 2000). Cercosporin is a perylenequinone that causes lipid peroxidation and alters membrane permeability by forming reactive oxygen species, thus functioning as a phytotoxin during pathogenesis (Daub and Ehrenshaft, 2000). Of particular importance is the fact that cercosporin biosynthesis is strongly induced by light, although other environmental factors are also involved (You et al., 2008). However, whether cercosporin plays a critical role in gray leaf spot has not been proven conclusively, there is considerable evidence that cercosporin is a broadly conserved, key component of pathogenesis among many *Cercospora* species. Upchurch et al. (1991) reported that a cercosporin non-producing mutant of *C. kikuchii*, generated by UV mutagenesis, was not able to cause typical necrotic lesions on soybean, and Callahan et al. (1999) demonstrated that *C. kikuchii* requires *CFP1*, a gene predicted to encode an ABC transporter, for protection against the toxic effects of cercosporin and virulence on soybean. In *C. zeae-maydis*, Shim and Dunkle (2003) demonstrated that the disruption of *CZK3*, which is predicted to encode a MAP kinase kinase, caused pleiotropic phenotypes, including the abolishment of cercosporin biosynthesis, conidiation, and pigmentation. When inoculated on maize leaves, the *CZK3* disruption mutant failed to elicit typical necrotic lesions, thus implicating *CZK3* in virulence. But due to the pleiotropic phenotype of the mutant, it is not conclusive that the reduction in virulence is caused partially or exclusively by the loss of cercosporin biosynthesis. In addition, a cluster of genes involved in cercosporin biosynthesis were recently identified and characterized (Chen et al., 2007a and b; Choquer et al., 2007; Dekkers et al., 2007). However, the
Is light a global regulator of pathogenesis in *C. zeae-maydis*?

The role of light in regulating conidiation and cercosporin biosynthesis raises questions of whether light influences additional components of pathogenesis. We recently discovered that light is required for stomatal tropism and appressorium formation in *C. zeae-maydis*, and that blue light is specifically required for the induction of cercosporin biosynthesis and the repression of conidiation (Kim et al., unpublished data). At the molecular level, fungal pathogenesis is widely regarded as an intricate and complex genetic regulatory network. In *C. zeae-maydis*, the involvement of light in numerous stages of pathogenesis strongly suggests that a) light regulates fungal development and this ultimately influences pathogen’s ability to infect plants, and b) the mechanisms underlying the perception of light and transduction of the signal are likely to play a central role in the genetic regulatory network underlying pathogenesis.

Known and predicted mechanisms underlying light-regulated pathogenesis in *C. zeae-maydis*

Light is a ubiquitous environmental cue that regulates numerous biologically important processes, and organisms across all kingdoms of life have evolved intricate molecular mechanisms to detect and respond to light (Idnurm and Crosson, 2009). Fungi are no exception, wherein elaborate systems for the detection of light at the molecular level have been characterized (Bahn et al., 2007; Purschwitz et al., 2006). Furthermore, a direct relationship between photoreceptors and pathogenicity has recently been established in fungi (Idnurm and Heitman, 2005; Kim et al., unpublished data; Ruiz Roldán et al., 2008).

How plants and fungi detect light is remarkably different at the molecular level. In plants, blue light has long been known to mediate light-induced responses via two broadly grouped classes of photoreceptors – phototropins and cryptochromes (Purschwitz et al., 2006). Phototropins contain a characteristic flavin-binding domain, the LOV (light oxygen voltage) motif, which imparts the capacity to regulate an array of photoreactive responses (Crosson et al., 2003). In fungi, the best understood category of blue-light photoreceptors is encoded by the well-characterized gene family white collar-1 (wc-1; Dunlap and Loros, 2006; Herrera-Estrella and Horwitz, 2007). wc-1 was initially described in a non-pathogenic ascomycete, Neurospora crassa, wherein it encodes a dual-function photoreceptor/GATA family zinc finger transcription factor WC-1. Additionally, WC-1 interacts with another GATA family zinc finger transcription factor, White Collar-2 (WC-2), via two congruent Per-ARNT-Sim (PAS) motifs to form the heterodimeric white collar regulatory complex (WCC; Ballario et al., 1996; Lee et al., 2003; Purschwitz et al., 2006). The WCC is essential for virtually all light-mediated responses in *N. crassa*, including circadian rhythmicity, morphogenesis, reproduction, secondary metabolism, and phototropism (Ballario et al., 1998; Dunlap and Loros, 2006; Herrera-Estrella and Horwitz, 2007). To date, genes similar to *wc-1* have been identified in a number of diverse fungi, including ascomycetes, basidiomycetes, and zygomycetes (Estrela and Avalos, 2008). The identification and characterization of *wc-1-like* genes has shown the degree of functional conservation, which suggests that the molecular mechanisms underlying responses to light may be at least partially conserved across the fungal kingdom.

Recently, *CRP1* of *C. zeae-maydis*, a homolog to the *N. crassa* *wc-1*, was cloned and functionally characterized (Kim et al., unpublished data). Disruption of *CRP1* via homologous recombination virtually abolished the ability of the fungus to form appressoria, even in the presence of stomata. During infection, *C. zeae-maydis* orients hyphal growth toward host stomata, whereupon the fungus initiates entry into mesophyll tissues by forming appressoria directly over stomatal pores. Hence, the severely reduced ability to form appressoria effectively rendered *CRP1* disruption mutants...
apathogenic, illustrating the importance of photoreception for virulence in \textit{C. zeae-maydis}. Interestingly, disruption of \textit{CPD1} did not abolish the production of cercosporin nor fully depress conidiation under blue light. These results suggest that additional, uncharacterized blue-light photoreceptors may exist in \textit{C. zeae-maydis}.

Cryptochromes, blue-light receptors that function exclusively in signal transduction, have been well-characterized in plants and animals (Lin and Todo, 2005), although much less is known about structurally similar proteins in fungi (Bluhm and Dunkle, 2008; Borkovich et al., 2004; Veluchamy and Rollins, 2008). A recent study identified \textit{PHL1}, a cryptochrome/6–4 photolyase-like gene in \textit{C. zeae-maydis} (Bluhm and Dunkle, 2008); disruption of \textit{PHL1} abolished photoreactivation after exposure to UV light. Additionally, \textit{PHL1} mutants were impaired in the induction of \textit{CPD1}, a cyclobutane pyrimidine dimer (CPD) photolyase, as \textit{RAD2} and \textit{RVB2}, putative orthologs of genes involved in nucleotide excision and chromatin remodeling during DNA damage repair (Bluhm and Dunkle, 2008). Although \textit{PHL1} mutants showed abnormalities in development and secondary metabolism, the disruption of \textit{PHL1} had no discernible effect on the ability of the fungus to infect and colonize maize leaves and does not appear to function as a blue-light photoreceptor involved in regulating gene expression (Bluhm and Dunkle, 2008).

Beyond blue-light receptors, the genome of \textit{C. zeae-maydis} contains a number of genes that encode homologs of red- and green-light photoreceptors in other organisms. A recent study in \textit{C. zeae-maydis} generated a collection of over 27,500 expressed sequence tags (ESTs) and evaluated their expression under a number of conditions, including response to light (Bluhm et al., 2008). A cluster highly similar to phytochromes was identified (Bluhm et al., 2008). Phytochromes are red/far-red light photoreceptors that have been shown to be important in regulating the transition between sexual and asexual reproduction in a select number of fungi (Blumenstein et al., 2005; Purschwitz et al., 2008). Interestingly, a red light-specific morphological or metabolic phenotype has not been observed in \textit{C. zeae-maydis} (Bluhm and Dunkle, unpublished data), and the potential role of a red-light photoreceptor in pathogenesis is not clear. In addition, homologs of photoreceptors in higher eukaryotes were identified, including blue-/green-light sensing opsins (Bieszke et al., 1999; Waschuk et al., 2005) and an additional protein sequence sharing similarity with plant blue-light sensing phototropin. Moreover, the study identified a number of putative light-regulated genes, including circadian clock regulated genes (Bluhm et al., 2008). To fully understand the potential roles, if any, of these genes in pathogenesis, functional genomics approaches are needed to define individual gene function in the context of host/pathogen interactions.

**Potential roles of light in host defense against \textit{C. zeae-maydis}**

One of the early focuses in plant pathology research was on investigating the relationship between light and disease, based on the hypothesis that pathogens utilize light to recognize favorable conditions for attack (Colhoun, 1973). As discussed above, a direct relationship between light perception and pathogen virulence has been established. In fact, without light, plants are unable to mount a proper defense response to pathogens (Guo et al., 1993; Lozano and Sequeira, 1970). Recently, progress has been made in model plants to better understand the role of light in regulating a complex network of signaling pathways in response to pathogen invasion (Roden and Ingle, 2009). Transcription of many plant defense genes is regulated in response to light (Griebel and Zeier, 2008; Karpinski et al., 2003). For instance, light is instrumental in inducing plant innate immunity via the salicylic acid (SA) signaling pathway (Roden and Ingle, 2009). Additionally, the induction of plant defense genes is elicited by disruptions in light harvesting, photosynthesis, and/or carbon metabolism caused by pathogen attack (Bechtold et al., 2005). The same response is observed in plants exposed to an increase in light intensity. Interestingly, disruption in these physiological processes due to host-pathogen interactions induce systemic acquired resistance (SAR; Bechtold et al., 2005). However, the specific contribution of analogous responses in maize to infection by \textit{C. zeae-maydis} remains to be determined.

Oxylipins, a diverse family of oxygenated metabolites derived from polyunsaturated fatty acids, regulate numerous physiological important processes in plants (Prost et al., 2005). In the context of this review, oxylipins are most notable as potent regulators of host defense in response to fungal pathogens (Farmer et al., 2003; Prost et al., 2005; Rosahl and Feussner, 2004). In maize leaves, the oxylipin encoding gene, \textit{ZmLOX10}, is primarily regulated by circadian rhythmicity (Nemchenko et al., 2006). Significantly, \textit{ZmLOX10} expression parallels the diurnal increase in photosynthesis activity (Nemchenko et al., 2006). This suggests that light may be a key factor involved in the induction of host defense regulators, such as oxylipins, in response to circadian rhythmicity.

**Future perspectives: Evidence for the coordinated response of host and pathogen to photoperiod?**

In filamentous fungi, biological rhythms regulate primary metabolism, growth and development (Jerebzojf, 1965). Particularly, light-regulated conidiation has been exten-
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In conclusion, the importance of light in the complex relationship of plant, pathogen, and photoperiod is becoming increasingly evident. However, despite recent advances in elucidating the complexity in such interactions, the underlying molecular mechanisms and signaling pathways remain poorly understood. Future efforts focusing on the molecular dissection of light-dependent components of fungal virulence and plant defense are required to gain a full appreciation of how light influences the development of plant diseases.

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