A Chitin Synthase with a Myosin-Like Motor Domain Is Essential for Hyphal Growth, Appressorium Differentiation, and Pathogenicity of the Maize Anthracnose Fungus

Colletotrichum graminicola

Stefan Werner,* Janeyce A. Sugui,* Gero Steinberg, and Holger B. Deising†

1Martin-Luther-Universität Halle-Wittenberg, Institute of Agricultural and Nutritional Sciences, Phytopathology and Plant Protection, Ludwig-Wucherer-Str. 2, D-06099 Halle (Saale), Germany; 2Max-Planck-Institute of Terrestrial Microbiology, D-35043 Marburg, Germany

Submitted 17 June 2007. Accepted 11 August 2007.

Chitin synthesis contributes to cell wall biogenesis and is essential for invasion of solid substrata and pathogenicity of filamentous fungi. In contrast to yeasts, filamentous fungi contain up to 10 chitin synthases (CHS), which might reflect overlapping functions and indicate their complex lifestyle. Previous studies have shown that a class VI CHS of the maize anthracnose fungus Colletotrichum graminicola is essential for cell wall synthesis of conidia and vegetative hyphae. Here, we report on cloning and characterization of three additional CHS genes, CgChsI, CgChsIII, and CgChsV, encoding class I, III, and V CHS, respectively. All CHS genes are expressed during vegetative and pathogenic development. ΔCgChsI and ΔCgChsIII mutants did not differ significantly from the wild-type isolate with respect to hyphal growth and pathogenicity. In contrast, null mutants in the CgChsV gene, which encodes a CHS with an N-terminal myosin-like motor domain, are strongly impaired in vegetative growth and pathogenicity. Even in osmotically stabilized media, vegetative hyphae of ΔCgChsV mutants exhibited large balloon-like swellings, appressorial walls appeared to disintegrate during maturation, and infection cells were nonfunctional. Surprisingly, ΔCgChsV mutants were able to form dome-shaped hyphopodia that exerted force and showed host cell wall penetration rates comparable with the wild type. However, infection hyphae that formed within the plant cells exhibited severe swellings and were not able to proceed with plant colonization efficiently. Consequently, ΔCgChsV mutants did not develop macroscopically visible anthracnose disease symptoms and, thus, were nonpathogenic.

Additional keywords: force measurement, fungal cell wall.

Chitin, the β-1,4-linked polymer of N-acetylglucosamine, represents a major component of the cell walls of higher fungi and is essential for hyphal growth and rigidity (Bartnicki-Garcia 1968; Cabib et al. 1996). After trans-membrane polymerization of chitin by membrane-localized glycosyl transferases called chitin synthases (CHS), chitin strands are covalently linked with β-glucan to contribute to a network of four structural cell wall components (i.e., chitin, β-1,3- and β-1,6-glucan, and mannanprotein) (Kollar et al. 1997; Sietsma and Wessels 1994; Wessels 1993).

The yeast Saccharomyces cerevisiae contains three CHS with distinct cell cycle functions in cell wall expansion, septum formation, and budding (Cabib et al. 1996, 2001). In comparison, the genome of filamentous fungi may contain as many as 10 CHS genes (Miyazaki and Ootaki 1997), grouped in six classes, with CHS of classes III, V, and VI being typical for filamentous fungi (Munro and Gow 2001). Because certain CHS are active during defined developmental stages of Aspergillus nidulans (Lee et al. 2004), one may argue that individual enzymes are required during different stages of complex life cycles and to adapt to different ecological niches (Ruei-Herrera et al. 2002). Alternatively, different CHS may concertedly contribute to certain processes in vegetative or pathogenic development, with overlapping roles (Roncero 2002). This view is supported by the fact that mutations in single CHS genes often do not cause obvious phenotypes (Motoyama et al. 1994, 1997; Specht et al. 1996), but double and triple mutations result in drastic cell wall defects (Fujiiwara et al. 2000; Ichinomiya et al. 2002; Motoyama et al. 1997; Shaw et al. 1991; Wang et al. 2001). With the exceptions of the maize smut fungus Ustilago maydis, the vascular wilt fungus Fusarium oxysporum, and the gray mold fungus Botrytis cinerea, the role of individual CHS in the infection process of plant-pathogenic fungi has not been studied (Madrid et al. 2003; Soulìé et al. 2006; Weber et al. 2006). These detailed investigations revealed essential roles of class V CHS (Madrid et al. 2003; Weber et al. 2006), suggesting that these myosin-like CHS are of particular importance for virulence of fungal pathogens.

Colletotrichum graminicola, the causal agent of maize leaf anthracnose and stalk rot, is a severe pathogen that caused considerable yield losses throughout the north-central and eastern United States (Bergstrom and Nicholson 1999; Warren et al. 1973). Differentiation of an appressorium is essential for in-
fectation of intact host tissue (Deising et al. 2000; Mendgen and Deising 1993; Mendgen et al. 1996). During maturation of appressoria of *Colletotrichum* spp., melanin is incorporated into the cell wall and osmotically active compounds are synthesized to high concentrations. Measurements with optical waveguides indicated that appressoria of *C. graminicola* generate considerable turgor pressure, which is translated into force that is exerted onto an epidermal cell in order to penetrate the host (Bastmeyer et al. 2002; Bechinger et al. 1999). Conceivably, rigid fungal cell walls are thought to be essential to support the high osmotic pressure and direct force during the infection process. Moreover, invasive growth of fungal hyphae in planta requires cell wall synthesis, which further suggests that wall-synthesizing enzymes are of high importance for fungal virulence. Random mutagenesis experiments allowed identifying a mutant of *C. graminicola* with conidia that burst and hyphal apices that swell in media with low osmotic potential (Epstein et al. 1998). According to recent classification (Munro and Gow 2001), the tagged gene encodes a class VI CHS, which is responsible for synthesis of approximately 30% of the chitin in conidial cell walls (Amnuaykanjanasin and Epstein 2003).

In this article, we report on genetical dissection of chitin synthesis in *C. graminicola* and show that CgChsV, a chitin synthase with a myosin-like motor domain, is essential for hyphal growth, infection structure differentiation, and pathogenicity.

**RESULTS**

Chitin synthesis is required for hyphal morphology, appressorium formation, and pathogenicity of *C. graminicola*.

Chitin represents a major cell wall polymer in filamentous fungi and is synthesized at the apex of a growing hypha. To localize chitin, vegetative hyphae of *C. graminicola* were stained with the tetramethylrhodamine isothiocyanate (TRITC)-conjugated chitin-specific lectin wheat germ agglutinin (WGA) (Fig. 1). Hyphae growing with few branches (Fig. 1A) as well as highly branched hyphae (Fig. 1B) showed strongly fluorescing apices (arrows), indicating chitin biosynthesis and deposition of newly formed chitin at hyphal tips. However, some apices were not decorated by WGA (compare hyphae marked by arrowheads in fluorescence images of Figure 1A and B), suggesting either that these hyphae were not actively growing or that the chitin was not accessible to the agglutinin, due to modification or masking of the polymer.

To investigate the importance of chitin for rigidity and function of cell walls of the maize pathogen *C. graminicola*, saprophytic and pathogenic development of the fungus was assessed in the presence of the competitive CHS inhibitor nikonkycin Z (Fig. 2). In liquid media, branched vegetative hyphae developed (Fig. 2A, left image). When grown in medium containing 50 μM nikonkycin Z, large hyphal swellings filled with vacuoles occurred along hyphae at irregular intervals (Fig. 2A, middle image), suggesting that the inhibition of CHS resulted in cell wall defects that were unevenly distributed along the hypha. Occasionally, swellings were closely associated and formed chains of protrusions (Fig. 2A, right image). These observations clearly indicate that chitin represents an essential structural polymer of vegetative hyphae. Next, we assessed the importance of chitin synthesis in infection structures differentiated in vitro and for infection competence on maize leaves (Fig. 2B through D). On hydrophobic hard surfaces such as polyester, conidia germinate, form a short germ tube, and differentiate a melanized appressorium (Deising et al. 2000). Already at the lowest nikonkycin Z concentration tested (10 μM), approximately 60% of the appressoria were distorted and showed drastically increased diameters (Fig. 2B and C). Although some of the enlarged appressoria were still melanized (Fig. 2C, arrow), others failed to melanize and had collapsed (Fig. 2C, white arrowhead). Occasionally, swollen germ tubes also were observed (Fig. 2C, black arrowhead). Increasing the nikonkycin Z concentration to 100 μM only led to minor increases of the portion of abnormal structures (Fig. 2B). Importantly, conidial germination rates were not significantly affected by the inhibitor at the concentrations tested (Fig. 2B).

Because differentiation of functional appressoria is essential for invasion of host leaves, we anticipated that inhibition of chitin synthesis would affect not only infection-related morphogenesis but also infection rates and disease symptom de-

---

**Fig. 1.** Localization of chitin by labeling with wheat germ agglutinin (WGA). **A,** Newly synthesized chitin localized at the apices of rapidly growing hyphae was stained with tetramethylrhodamine isothiocyanate-labeled WGA (right image, arrow). Some hyphae did not show apical fluorescence (compare arrowheads in left differential interference contrast (DIC) and right fluorescence images). **B,** Localization of chitin in older, heavily branched parts of the mycelium. Arrows indicate WGA-labeled hyphal tips; arrowheads in upper DIC and lower fluorescence images indicate nonfluorescing apices. Bars are 10 μm.
development. Indeed, increasing inhibitor concentrations resulted in reduction of anthracnose symptom severity, and no symptoms were visible when the infection droplets contained 100 μM nikkomycin Z (Fig. 2D). As expected, clear anthracnose symptoms occurred in the absence of the chitin synthase inhibitor, and control leaves mock inoculated with distilled water did not develop disease symptoms (Fig. 2D). Taken together, these data suggest that chitin synthesis is indispensable for vegetative growth and pathogenicity of C. graminicola.

Identification of CHS from C. graminicola.

To isolate CHS genes of C. graminicola, we performed polymerase chain reaction (PCR) studies with degenerate primers designed to amplify class I to III and class IV to VI genes. Four PCR fragments of 619, 638, 660, and 710 bp were obtained. Sequence analyses of the 710-bp PCR fragment amplified with primers CHS3 and CHS4 showed that it corresponds to a part of the class VI CgChsVI gene originally designated chsA (Amnuaykanjanasin and Epstein 2003). This gene has been studied in detail previously; therefore, it was not further analyzed here. The 619- and 638-bp PCR fragments amplified with primers CHS1 and CHS2 were used to screen a cosmid library. Purified cosmid clones contained the complete coding regions of a class I (CgChsI) and a class III (CgChsIII) CHS gene (classification was done according to Munro and Gow [2001]; see also Mellado and associates [2003] and Takeshita and associates [2006]). The complete sequences of the CHS gene of C. graminicola studied here are available under GenBank accession numbers AY052545 (CgChsI), AY052546 (CgChsIII), and AY052547 (CgChsV). Phylogenetic analyses showed that the class I and class III CHS are most closely related with the corresponding enzymes NcChs3 and NcChs1 of Neurospora crassa, with 74 and 71% amino acid identity, respectively (Fig. 3A). The proteins consist of 899 and 912 amino acids, respectively, and show a typical CHS core region (Nagahashi et al. 1995). Hydropathy plots suggest that both enzymes have eight transmembrane domains (Fig. 3B). In addition to the class I and III CHS genes, we identified a class V CHS gene of C. graminicola, designated CgChsV. CgChsV has a 5,709-bp open reading frame, which is interrupted by two introns (nucleotides 311 to
This derived protein, like CgChsVI, has an N-terminal myosin-like motor domain and a CHS core region at the C-terminus (Fig. 3B). The myosin motor domain of CgChsV shows strong similarity (2.2 e-115) with the myosin motor consensus sequence and has a P-loop ATP binding site (GESGSG) as well as the regulatory IQ motif. In comparison, the myosin-like domain of CgChsVI is much shorter and shows only limited similarity with the myosin motor consensus sequence (2.9 e-7). For both CHS of C. graminicola with myosin-like motor domains, six transmembrane domains have been predicted (Fig. 3B). In total, the proteins contain 1,866 (CgChsV) and 1,783 (CgChsVI) amino acids. The expression profile of the four CHS genes identified in C. graminicola was studied by reverse-transcription (RT)-PCRs with RNA isolated from vegetative hyphae and pre- and post-penetration stages of pathogenic development (i.e., from in vitro infection structures that had formed appressoria and from infected leaves). All genes were expressed at all stages of fungal development investigated (Fig. 3C).

**Generation of ΔCgChsI, ΔCgChsIII, and ΔCgChsV mutants.**

To functionally characterize CgChsI, CgChsIII, and CgChsV, mutants were generated in which a part of the CHS coding region was replaced by the hygromycin phosphotransferase (hph) gene of Escherichia coli. As an example, targeted inactivation of CgChsV is shown (Fig. 4). The knock-out (KO) vector consisted of 974 bp of the CgChsV gene 5′ and 1,621 bp 3′ of the hph gene (Fig. 4A) and was transformed into the C. graminicola wild-type strain CgM2. In order to discriminate between transformants with homologous and ectopic integration of the KO vector, Southern blot experiments were performed with HindIII-digested DNA from the wild-type CgM2 and different independent transformants. While the wild-type strain showed a single 5.6-kb band hybridizing with the CgChsV probe, KO mutants exhibited a 7.7-kb band. In addition to the 5.6-kb CgChsV wild-type band, a high molecular weight band was detected in transformants with ectopically integrated KO vector (Fig. 4B).

To demonstrate that partial replacement of the CgChsV gene resulted in failure to synthesize the CgChsV transcript, RT-PCR experiments were performed with total RNA from vegetative hyphae of the wild-type strain CgM2 and the transformants used in Southern blot experiments. As expected, CgChsV transcripts were present in the wild-type strain and transformants with ectopically integrated KO vector, but absent from four transformants with homologous integration of the KO vector (Fig. 4B).
ments, transformants carrying homologous and ectopic integrations of the KO vector for CgChsI and CgChsIII also were obtained (data not shown). However, C. graminicola mutants defective in CgChsI and CgChsIII did not display any obvious phenotype (growth rate, hyphal morphology, and conidiation rates) differing from the wild type and their virulence on maize leaves was not altered (data not shown).

CgChsV confers rigidity to vegetative hyphae of C. graminicola.

Inactivation of CgChsV drastically affected vegetative growth of C. graminicola (Fig. 5). Although the wild-type strain CgM2 and transformants with ectopically integrated KO vector grew comparably well on oatmeal agar (OMA), the ΔCgChsV mutant failed to grow on this substratum. However, on OMA osmotically stabilized with 150 mM KCl, growth of the ΔCgChsV mutant was partially rescued (Fig. 5A). Under these conditions, wild-type hyphae grew straight with some branching (Fig. 5B). Even in the presence of high osmolyte concentrations, significant cell wall defects were obvious in the ΔCgChsV mutant. In some parts of the mycelium, hyphae grew with swellings occurring occasionally (Fig. 5C) whereas, in others, chains of protrusions almost completely lacking filamentous stretches were visible (Fig. 5D). Cell wall defects were reminiscent of those caused by the chitin synthase inhibitor nikkomycin Z, although the latter primarily occurred at the hyphal tips (compare Figs. 2A and 5B through D). Determination of the N-acetyl glucosamine contents suggested that chitin content is reduced by approximately 30% in ΔCgChsV compared with wild-type hyphae (S. Werner, H. B. Deising, and B. Morschbacher, data not shown). Complementation of a ΔCgChsV strain with the entire CgCHSV gene driven by the glyceraldehyde-3-phosphate promoter of A. nidulans complemented growth and hyphal morphology defects of ΔCgChsV, but did not fully restore its ability to produce conidia (data not shown). These data demonstrate that CgChsV has essential roles in wall formation in C. graminicola.

CgChsV is essential for appressorium formation but dispensable for hyphopodium development and function.

Because appressoria need very rigid cell walls to control the high osmotic pressure, it was of interest to see whether or not the ΔCgChsV mutant would be able to differentiate morphologically and functionally normal infection cells (Fig. 6). ΔCgChsV mutants failed to form acervuli with falcate conidia in regular complete medium (CM); however, in liquid CM supplemented with 0.5 M sucrose, both the wild type and the ΔCgChsV mutant produced conidia. Although wild-type conidia were falcate (Fig. 6A), the ΔCgChsV mutant formed both falcate and shorter cylindrical conidia (Fig. 6B and C). Even in the presence of 0.5 M sucrose, the wild-type conidia formed a short germ tube (Fig. 6A, arrowheads) and differentiated a melanized appressorium on the surface of polyester sheets (Fig. 6A, arrows). Conidia of the ΔCgChsV mutant also germinated in CM medium with 0.5 M sucrose and formed appressoria. However, ΔCgChsV appressoria were severely distorted and the appressorial cell wall appeared to be disintegrated (Fig. 6B, arrow). The fact that germ tubes formed (Fig. 6B, arrowhead) suggests that ΔCgChsV conidia do not rupture and release cytoplasm upon germination. Moreover, the size of the distorted cells is comparable with that of wild-type appressoria (compare cells in Figure 6A to C, arrows). The distorted appressoria showed some melanization.

Fig. 4. Targeted inactivation of the CgChsV gene of Colletotrichum graminicola. A, Strategy of inactivation of the CgChsV gene by homologous recombination. In the CgChsV knock-out (KO) vector, a 3,021-bp SalI fragment was replaced by the hygromycin resistance cassette, and this construct was transformed into the C. graminicola wild-type strain CgM2. Bars indicate probe used in Southern blot experiments. B, Southern blot of DNA isolated from C. graminicola wild type and transformants with homologous (KO) and ectopic integration (ectopic) of the KO vector. In reverse-transcription polymerase chain reaction (RT-PCR) analyses, products only occurred in the wild-type strain and in C. graminicola strains with ectopic integration of the KO vector (compare Southern blot and RT-PCR). Fragment sizes are given in kilobases.
(Fig. 6C) and some of these cells were attached to the polyester substratum (not shown). Very occasionally, the formation of an appressorial initial was observed (Fig. 6D, arrowhead); however, during enlargement and maturation of these cells, the appressorial wall was unable to maintain the appressorial shape. These data suggest that CgChsV has essential roles in appressorium differentiation.

*C. graminicola* can also invade the host plant by forming infection structures called hyphopodia. These are lobed melanized infection cells formed by vegetative hyphae (Fig. 6E, arrow), which are thought to functionally resemble appressoria. Surprisingly, the ΔCgChsV mutant differentiated melanized dome-shaped hyphopodia of a size comparable with that of wild-type hyphopodia (Fig. 6F, arrow). However, occasionally large and swollen melanized hyphopodia were found (Fig. 6G, arrow). In hyphopodia, no cell wall distortions comparable with those in appressorial walls were found (Fig. 6G, compare hyphopodium indicated by arrow and disintegrated appressorium indicated by arrowhead). Thus, CgChsV participates in the formation of hyphopodia but, in contrast to the role of this enzyme in appressoria, it is not essential for hyphopodium differentiation. Next, we tested whether hyphopodia of ΔCgChsV mutants were able to exert force in order to invade the host epidermis. Surprisingly, quantitative measurements by optical waveguides revealed that force exerted by wild-type and ΔCgChsV hyphopodia did not differ significantly (Fig. 6D). Moreover, force generated by hyphopodia was comparable with that exerted by wild-type appressoria (Bechinger et al. 1999).

**CgChsV is essential for hyphal growth in planta.**

ΔCgChsV null mutants were able to differentiate hyphopodia, suggesting that they are able to infect plants. To investigate the role of *CgChsV* in pathogenic development, infection assays were performed with the ΔCgChsV mutant and wild-type strains (Fig. 7). Microscopical analyses of infection by wild-type cells showed that hyphopodia (Fig. 7A, asterisk) invaded the host leaf to form thick, primary hyphae (Fig. 7A, large arrowhead) from which thin, destructive secondary hyphae (Fig. 7A, small arrowhead) grew out to colonize the host tissue. Thus, infection hyphae initiating from hyphopodia closely resemble those developing from appressoria. Consistent with the in vitro studies described above, both wild-type and ΔCgChsV mutants formed hyphopodia on the plant surface when osmotically stabilized (Fig. 7A; black bars), and post-penetration infection structures formed at comparable rates (Fig. 7B: gray bars, infection vesicles; hatched bars, primary hyphae; white bars, secondary hyphae). However, the infection hyphae formed by ΔCgChsV mutants within the first host cell showed large swellings (Fig. 7A, arrows). These hyphae were able to penetrate host cell walls to grow into neighboring cells, but the ability to colonize the host tissue was severely reduced.

Consequently, when the inoculum was applied to host leaves in the presence of osmolytes, the ΔCgChsV mutant did not cause any disease symptoms, whereas wild-type and transformants with ectopically integrated KO vector were able to cause anthracnose lesions on wounded and unwounded leaves (Fig. 7C).

These results show that CgChsV is required for post-penetration pathogenic development of *C. graminicola*. In the absence of osmotic stabilization, however, the ΔCgChsV mutant was unable to differentiate hyphopodia on the maize leaf surface (Fig. 7B) and, therefore, infections did not occur under these conditions.

In order to test whether the pathogenicity defect of ΔCgChsV mutants also was due to increased sensitivity to plant defense...
formants were 83.6 ± 15.0% and 87.3 ± 2.3%, respectively (data not shown). Differences in inhibition were not statistically different (P = 0.05). Although these data indicate that pathogenicity defects are primarily due to cell wall defects rather than increased sensitivity to H2O2 (Madrid et al. 2003), it should be noted that vegetative hyphae analyzed in growth assays cannot directly be compared with infection hyphae.

**DISCUSSION**

Chemically, the fungal cell wall consists of 80 to 90% polysaccharides (Bartnicki-Garcia 1968), and the gross monomeric composition of an average ascomycete cell wall shows a predominance of glucose and amino sugars, predominantly N-acetylglucosamine (Sietsma and Wessels 1994). Chitin may contribute more than 50% of the dry weight of the cell wall (Aronson and Machlis 1959) and is thought to be essential for fungal cell wall rigidity. In several filamentous fungi, newly synthesized chitin localizes to the hyphal tip (Weber et al. 2006; Wessels 1993) and, by WGA labeling, we demonstrated that this also holds true for *C. graminicola* (Fig. 1). Strongly stained hyphal tips suggest that chitin is a prominent polymer in this fungus. The fact that only hyphal tips but not older, subapical regions expose chitin at the surface may be due to enzymatic modification of chitin that could be mediated by secreted chitin deacetylases (El Gueddari et al. 2002). Alternatively, lack of hyphal decoration by WGA could be due to apposition of other cell wall layers such as hydrophobin coatings (Wösten et al. 1994). Some hyphal tips of *C. graminicola* were not labeled by WGA, suggesting either that these hyphae were not actively growing or that the entire hyphal surface may have been modified (Sietsma and Wessels 1994), rendering them inaccessible for the lectin.

Studies involving CHS inhibitors, namely nikkomycins and polyoxins, demonstrated the importance of these enzymes and the polymers they synthesize for cell wall rigidity and function and the genomes of filamentous fungi (Munro and Gow 2001) and, although inhibitor studies allow the determination of the importance of chitin synthesis (Zhang et al. 2000), they do not lead to an understanding of the role of individual CHS genes or enzymes. Targeted gene inactivation experiments made it possible to genetically dissect chitin synthesis and to investigate the one or more roles of individual genes in plant (Madrid et al. 2003; Weber et al. 2006) and human-pathogenic fungi (Munro and Gow 2001).

We have cloned and functionally characterized a class I, a class III, and a class V CHS gene of the filamentous ascomycete and maize pathogen *C. graminicola*. This is the first report that a class V CHS with a myosin-like motor domain is indispensable for vegetative growth, differentiation of infection structures, and pathogenic development of this plant-pathogenic fungus.

**Structure of the C. graminicola CHS.**

Due to their high degree of conservation, class I to III CHS genes have been isolated with degenerate PCR primers (Bowen

---

**Fig. 6.** Infection structure differentiation and hyphopodial force exertion by *Colletotrichum graminicola* wild type and the ΔCgChsV mutant strain. Appressorium differentiation by **A**, the wild-type strain CgM2 and **B** through **D**, the ΔCgChsV mutant strain. On a hard, hydrophobic surface, wild-type conidia germinate, form a short germ tube (**A**, arrowheads), and differentiate a melanized appressorium (**A**, arrows). Likewise, conidia of the ΔCgChsV mutant form a germ tube (**B**, arrowhead) and an appressorium (**B**, arrow), but the infection cell is drastically distorted. All fully enlarged appressoria differentiated by ΔCgChsV conidia show lyzed cell walls (**C**, arrows). Appressorial initials (**D**, arrow) develop from germ tubes (**D**, white arrowhead) but exhibit cell wall defects after maturation, as indicated by a second appressorium (**D**, black arrowhead) formed by the same conidium. Differentiation of hyphopodia by **E**, the wild-type strain CgM2 and **F** and **G**, the ΔCgChsV mutant strain. Although the wild-type strain forms lobed, melanized hyphopodia (**E**, arrow), hyphopodia of the mutant also are melanized, but dome-shaped (**F** and **G**, arrows). Occasionally, larger, melanized swollen hyphopodia can be observed (**G**, arrow). Hyphopodia (**G**, arrow) do not exhibit cell wall defects comparable with those found in appressoria (**G**, arrowhead) developing in close vicinity. Infection cells were observed 48 h after inoculation of polyester sheets. Bars indicate 10 μm. **H**, Force measurement using optical waveguides. Force exerted by hyphopodia of the wild-type (WT) and ΔCgChsV mutant strain (KO) were measured by optical waveguides.
et al. 1992). The homologous region is restricted to approximately two-thirds of the carboxyl terminus and, in most cases, hydrophobic membrane-spanning domains are found near this end (Cabib et al. 1996). Although these features have been established with the yeast \textit{S. cerevisiae}, they also hold true for other fungi (Weber et al. 2006). In \textit{C. graminicola}, class I and III CHS exhibit eight hydrophobic membrane-spanning domains at their carboxyl ends and, thus, are typical examples of these enzyme classes. Whether or not they are zymogenic like other class I to III CHS (Mellado et al. 2003) has not been determined.

Class IV to VI CHS genes have been isolated either using PCR primers based on the \textit{ScCHS3} sequence, with primers specifically designed to amplify fragments of CHS with a myosin-like motor domain, or by random insertional mutagenesis (Annuaykanjanasin and Epstein 2003; Madrid et al. 2003; Munro and Gow 2001). In contrast to class IV CHS, enzymes classified as class V CHS have a myosin-like motor domain with significant similarity to the consensus sequence of single-headed myosins, including a classical ATP-binding motifs (P-loop and switch I and II motifs) and the IQ motif. The myosin motor of some CHS (e.g., Chs4 of \textit{Paracoccidioides brasiliensis}, AoChsZ of \textit{A. oryzae}, and UmChs6 of \textit{U. maydis}) are somewhat degraded and, thus, distinct from class V CHS. Nino-Vega and associates (2004) proposed that these CHS should be classified as a new class of enzymes, and enzymes with a shorter myosin-like binding domain lacking classical ATP-binding motifs or the IQ motif have now been designated as class VI CHS (Mellado et al. 2003; Munro and Gow 2001; Takeshita et al. 2006). In \textit{C. graminicola}, typical class V (this study) and VI (Annuaykanjanasin and Epstein 2003) CHS exist, each consisting of approximately 1,800 amino acids and exhibiting six transmembrane domains and either a complete or a shorter myosin-like motor domain (Fig. 3).

**Filamentous growth requires a class V CHS with a myosin-like motor domain.**

Class III, V, and VI CHS have been identified only in filamentous and some dimorphic fungi that are able to differentiate true hyphae. Individual replacement mutagenesis of the class I and III genes \textit{CgChsI} and \textit{CgChsIII} of \textit{C. graminicola} did not cause alterations in hyphal development or conidiation. This may not necessarily be due to lack of function in filamentous growth but, rather, could be due to overlaps in gene and enzyme function. Also, in \textit{A. fumigatus}, deletion of the class I, II, and IV CHS genes \textit{AfchsA}, \textit{AfchsB}, and \textit{AfchsF} had little effect on morphology and growth (Munro and Gow 2001). A class III CHS was not important in the dimorphic basidiomy-

![Fig. 7](image-url) Penetration competence and pathogenicity of \textit{Colletotrichum graminicola} wild-type strain, \textit{ΔCgChsV} mutant strain, and a strain with ectopic integration of the knock-out (KO) vector. **A**, Microscopical analysis of the penetration process. Wild-type and \textit{ΔCgChsV} hyphopodia (asterisks) formed in the presence of 0.5 M sucrose and invaded the host epidermal cell. The wild type formed thick primary hyphae (large arrowhead) and thin secondary hyphae (small arrowhead). Intracellular hyphae of the \textit{ΔCgChsV} mutant showed balloon-like swellings (arrows) and grew more slowly than corresponding wild-type hyphae. **B**, Osmolyte effect on infection structure differentiation by wild-type and \textit{ΔCgChsV} mutant on maize leaves. In the absence of the osmolyte, the \textit{ΔCgChsV} mutant is unable either to grow and form hyphopodia on the leaf surface or to differentiate infection structures in planta. In the presence of 150 mM KCl, rates of infection structure differentiation are comparable. Black bars, hyphopodia; gray bars, infection vesicles; hatched bars, primary hyphae; white bars, secondary hyphae. Error bars indicate standard deviation. Infection structure formation was scored 48 h postinoculation. **C**, Leaf infection assays. Mycelia of the wild type (WT), a \textit{ΔCgChsV} mutant strain, and a strain with ectopic integration of the KO vector were placed onto the surface of intact or wounded maize leaves. Infection droplets contained 0.5 M sucrose. Regardless of wounding, anthracnose symptoms developed after inoculation with the WT strain and a strain with ectopic integration of the KO vector. The \textit{ΔCgChsV} mutant strain (KO) did not cause disease on either intact or wounded leaves. Non-inoculated control leaves (C) also are shown. Photographs were taken 6 days postinoculation.
cete U. maydis (Gold and Kronstad 1994; Weber et al. 2006) and class III CHS do not even exist in Phlycomyces blakesleeanaus (Miyazaki and Ootaki 1997). In contrast, mutants of the gray mold fungus B. cinerea deficient in the class III CHS BcChs3a were severely impaired in growth on solid substratum (Soulié et al. 2006). In U. maydis, two class IV CHS, Chs5 and Chs7, have been shown to be essential for fungal morphogenesis and filamentous growth (Weber et al. 2006), and class IV CHS also have important roles in hyphal growth of the dimorphic fungus C. albicans (Mio et al. 1996) and in the yeast S. cerevisiae (Shaw et al. 1991). However, in the filamentous fungus N. crassa, deletion of class IV CHS was without significant effect (Beth Din et al. 1996). These findings raised doubts about specific cellular functions of individual classes of CHS, and the situation is even more complicated by the fact that CHS of different classes cooperate to mediate hyphal growth and conidiation (Fujiwara et al. 2000; Motoyama et al. 1997).

A growing body of literature shows that mutations in individual class V and VI genes drastically affect filamentous growth. C. graminicola ΔCgChsV replacement mutants showed swollen hyphal tips and balloon-like structures along the hyphae, similar to those seen in A. nidulans mutants deficient in the class V CHS gene csmA (Horiuchi et al. 1999). In F. oxysporum class V CHS mutants, occurrence of balloon-like structures was suppressed in solid synthetic media complemented with osmotic stabilizers, indicating that swellings result from cell wall weakening and, in turn, that class V CHS play a major role in maintenance of hyphal wall integrity and polarized cell wall synthesis (Madrid et al. 2003). As indicated by the inability of C. graminicola ΔCgChsV mutants to form conidia on the surface of solid substrata, the class V CHS also plays a role in asexual sporation. This also has been observed in ΔcsmA mutants of A. nidulans. In addition to cell wall weakening, inactivation of the csmA gene led to formation of hyphae within previously formed older hyphae (intrahyphal growth) (Horiuchi et al. 1999).

Mutating the A. nidulans class VI CHS gene csmB caused formation of balloons, intrahyphal hyphae, lysis in subapical hyphal regions, and drastically reduced conidiation; thus, csmA and csmB mutants show a similar phenotype. Staining experiments indicated that CsmA and CsmB co-localized with actin, but immuno-precipitation experiments did not suggest a physical interaction between these enzymes (Takeshita et al. 2006). In C. graminicola, inactivation of the class VI CHS gene caused conidial budding and swelling of hyphal tips in media with low osmotic pressure, indicating the role of this enzyme in cell wall strength of conidial and vegetative hyphae (Ammuyakanjanasir and Epstein 2003). These data suggest that filamentous growth in several fungi, including C. graminicola, requires class V or class VI CHS with a myosin-like motor domain.

Role of chitin synthases in pathogenesis.

Apical growth is a hallmark of filamentous fungi, allowing invasion and colonization of solid substrata, including living tissues of either plant or animal origin (Wessels 1993). Fungal pathogenesis requires invasive growth (Deising et al. 2000; Stoldt et al. 1997); therefore, one may expect that CHS exclusively found in filamentous fungi are required for pathogenicity or virulence. Functional characterization of enzymes such as chitin and β-1,3-glucan synthases is essential not only to understand fungal pathogenicity but also to identify novel fungicide targets to be used in agriculture and medicine (Debono and Gordee 1994; Georgopapadakou 2001).

In comparison with information on the role of chitin synthases in vegetative development, little is known about the role of chitin synthesis and CHS during pathogenic growth. CHS with myosin motor domains have been found in filamentous fungi, including plant pathogens such as F. oxysporum (Madrid et al. 2003), B. cinerea (Choquer et al. 2004), Magnaporthe grisea (Vidal-Cros and Boccara 1998), and U. maydis (Weber et al. 2006), and also in human pathogens such as A. fumigatus (Auffavre-Brown et al. 1997), Paracoccidioides brasiliensis (Nino-Vega et al. 2004), and Wangiella dermatitides (Liu et al. 2004). The mcsI gene of U. maydis is specifically required for invasive growth in planta. However, the class VI CHS of the corn smut fungus U. maydis, Chs6, also contributes to fungal pathogenicity (Garcera-Teruel et al. 2004; Weber et al. 2006). Likewise, targeted chsV replacement mutants of F. oxysporum failed to colonize the vascular system of tomato plants and to invade wounded tomato fruit (Madrid et al. 2003).

In contrast to U. maydis and F. oxysporum, the corn anhacrose fungus C. graminicola differentiates distinct appressoria. These infection cells are melanized and previous studies have shown that they exert significant force to invade an epidermal host cell (Bechinger et al. 1999). Because nonmelanized mutants are unable to infect corn (Rasmussen and Hanau 1989), generation of turgor and its translation into force are thought to be essential for pathogenesis. On polyester sheets, ΔCgChsV mutants formed germ tubes and appressorial initials, but failed to differentiate functional (i.e., plant cell wall-penetrating) appressoria (Fig. 6). These data clearly indicate that CgChsV is indispensable for synthesis of rigid appressorial cell walls and for appressorium-mediated plant infection. Surprisingly, ΔCgChsV mutants were able to synthesize hyphopodia, which exhibited a distinct morphology, but exerted force and penetrated the host epidermis at rates comparable with wild-type hyphopodia (Figs. 6 and 7), allowing the study of infection structure formation in planta. After entering the epidermal host cell, C. graminicola ΔCgChsV primary hyphae swell markedly, reminiscent of the phenotype of the infection hyphae of U. maydis Δmcs1 mutants (Weber et al. 2006), and anhacrose disease symptoms did not develop. Thus, the class V chitin synthase CgChsV of C. graminicola is essential for both pre- and post-invasive stages of plant infection. Class V mutants of neither C. graminicola nor U. maydis exhibit increased sensitivity toward H2O2, suggesting that reduced virulence is due to cell wall defects per se, rather than to increased sensitivity to plant defense compounds (Weber et al. 2006; this study). Thus, increased sensitivity to reactive oxygen species may be specific for the class V chitin synthase mutant of F. oxysporum (Madrid et al. 2003).

Class V CHS of the plant pathogens C. graminicola, U. maydis, and F. oxysporum, as well as of the black fungal pathogen of humans, W. dermatitidis contribute to virulence (Liu et al. 2004; Madrid et al. 2003; Weber et al. 2006; this study); therefore, it would be tempting to investigate the role of these enzymes in a broader spectrum of pathogenic fungi. It is important to note, however, that pathogenicity assays in plants and animal models are difficult to compare, as indicated by studies with the tomato pathogen F. oxysporum. chsV mutants of this fungus are nonpathogenic on plants but show increased virulence in immunodepressed mice (Ortoneda et al. 2004).

Our study has shown that a class V CHS is a viability and pathogenicity factor in the maize anthracnose fungus C. graminicola and suggests that this class of fungal CHS may be important for infection of plants. Although general conclusions cannot be drawn so far, increasing evidence suggests that class V CHS have important roles in fungal pathogenicity (Madrid et al. 2003; Weber et al. 2006), and one may speculate that these enzymes could represent excellent novel fungicide targets. Entire fungicide classes have been lost due to development of fungicide resistance in fungal pathogen populations; therefore, appropriate fungicide mixtures may not durably be available in
agriculture, and the search for new fungicides will become increasingly important (Deising et al. 2002).

**MATERIALS AND METHODS**

**Fungal cultures and in vitro differentiation of infection structures.**

*C. graminicola* (Ces.) G. W. Wilson (teleomorph *Glomerella graminicola* D. J. Politis) strain M2 was kindly provided by R. L. Nicholson, Purdue University, IN, and used as wild-type strain in this study. The wild-type strain and ΔCgChsI and ΔCgChsIII mutants were grown on OMA (50 g of bled oatmeal [Haferflocken Großblatt aus ökologischem Anbau, Bohlsener Mühle, Boihlen, Germany] and 12 g of agar-agar [Difco Laboratories, Augsburg, Germany] per liter) at 23°C under continuous fluorescent light (Climas Control CIR, UniEqquip, Martinsried, Germany). Alternatively, strains were grown in CM (Leach et al. 1982) in an incubation shaker (UniEquip, Martiensried, Germany). To amplify DNA fragments of class IV to VI chitin synthases, primers CHS3 (5′-AAC CCI GGI AAY MGR GGI AAR MG-3′) and CHS4 (5′-GTT RTG IAC IGT IGA RTT DAT CC-3′) were designed. Reaction mixtures (10 to 20 μl) contained genomic DNA (10 ng), primers (0.5 μM each), 0.02 U of Taq polymerase/μl (Qiagen, Hilden, Germany), 0.2 mM dNTP each, and 1.5 mM MgCl2 in reaction buffer. The PCR program included initial denaturation (5 min at 95°C); 30 cycles of 20 s of denaturation at 94°C, 30 s of annealing at 60°C, and 60 s of chain elongation at 72°C; and a final elongation for 5 min at 72°C. Fragments of 619, 638, 660, and 710 bp were cloned into SmaI-digested pTZ19R.

The PCR fragments were digoxigenin (DIG) labeled by PCR with a dNTP mix containing DIG-dUTP and used to screen a genomic DNA library, as recommended by the manufacturer (PCR DIG Probe Synthesis Kit; Roche Diagnostics, Mannheim, Germany). The cosmid library of *C. graminicola* was constructed in the cosmid vector pSV50 (Vollmer and Yanofsky 1986) and kindly provided by J. A. Rollins and R. M. Hanau, Purdue University, West Lafayette, IN, U.S.A. Cosmid clones containing the entire gene corresponding to the 660-bp fragment have not been found, and inverse PCR (Pang and Knecht 1997) with *Mph*I103I- and *Hind*III-digested genomic DNA were performed to cover the complete coding region.

DNA sequencing was done on an ABI Prism 310 automatic sequencer using BigDye Chemistry (Applied Biosystems; Weiterstadt, Germany), following the manufacturer’s instructions. Sequence analyses were performed using the software supplied by ABI and the DNAStar programs (DNASTAR Inc., Madison, WI, U.S.A.). BLAST search was carried out on the National Center for Biotechnology Information server.

**Targeted gene disruption.**

To generate the CgChsI KO vector, the *CgChsI* gene was digested with *Cfr*I421 and *Bam*HI, removing a 1,037-bp fragment of the coding region. After treatment with the Klenow fragment, the 2,395-bp fragment of this construct was amplified using the Expand Long-Template PCR system (Roche Diagnostics) and the primers CHS11 (5′-GTC ACA AAC ATC AGA CAC CCG T-3′) and CHS14 (5′-CAC CAC GGT AGT AGT CCA TCT G-3′). The amplified fragment was transformed into CgM2 to generate ΔCgchsl mutants.

The 2,691-bp Acc65I-NdeI fragment of pUCATPH carrying the hygromycin resistance cassette was ligated into the CgChsIII gene digested with the same enzymes, so that 1,093 bp of the coding region of this CHS gene was replaced. A 4,91-kbp fragment of this construct was amplified using the Expand Long-Template PCR system (Roche Diagnostics) and the primers CHSH11 (5′-ACC CAA CCT AGG TCT CCT TCC-3′) and CHSH17 (5′-TCT AGA CAG CAT GGT TGG AAT GG-3′) and the Expand Long-Template PCR system (Roche Diagnostics). To generate ΔCgchshIII mutants, the amplified fragment was transformed into CgM2.

The CgChsV gene was digested with *Sal*I, removing a 3,021-bp fragment, and pUCATPH was digested with *Hind*III. T4-DNA polymerase treatment was performed such that the CgChsV ends carried a T- and pUCATPH carried an A- overhang. Thus, after ligation, the KO vector contained 974 bp of the CgChsV gene 5′ and 1,621 bp 3′ of the hygromycin resistance cassette. This construct was transformed into CgM2.
Transformation of C. graminicola basically followed the protocol described by Epstein and associates (1998), with some modifications. SYE medium (0.5 M sucrose and 0.1% [wt/vol] yeast extract) was inoculated to contain 10^6 falcate conidia/ml and incubated in an incubation shaker (Unitron, Infors AG, Bottmingen, Switzerland) at 23°C and 110 rpm for 3 days. Cultures were filtered through two layers of cheesecloth and the filtrate containing oval conidia was centrifuged at 4,260 x g at 4°C for 10 min. The conidia were resuspended in 10 ml of a protoplasting solution containing lysing enzyme from Trichoderma harzianum at 20 mg/ml (Sigma, Deisenhofen, Germany) and 0.1% (vol/vol) β-mercaptoethanol in 0.7 M NaCl, and incubated at 30°C for 3 h with slight agitation. Formation of protoplasts was controlled microscopically. The protoplasts were sedimented by centrifugation (800 x g at 4°C for 10 min), washed in 10 ml of SCT (1 M sorbitol; 50 mM CaCl_2; and 50 mM Tris-HCl, pH 8.0), centrifuged again, and resuspended in 1 ml of SCT. The KO vector (0.5 to 10 μg) was added to 100 μl of the protoplast suspension and incubated on ice for 30 min. After adding 1 ml of PEG solution (40% polyethylene glycol 4000; 0.6 M KCl; 50 mM CaCl_2; and 50 mM Tris-HCl, pH 8.0) and incubation at room temperature for 20 min, 3 to 4 ml of liquid regeneration medium (1 M sucrose, 0.1% [wt/vol] yeast extract, 0.1% [wt/vol] casein, and 0.6% [wt/vol] agar; 45°C) was added and poured onto selection plates containing hygromycin B at 400 μg/ml and 1.5% (wt/vol) agar. After 7 to 10 days, growing colonies were transferred to new selection plates containing hygromycin B at 100 μg/ml. To obtain homokaryotic transformants, colonies that had developed on selection plates were allowed to conidiate on OMA. Conidia again were plated out on selection plates containing hygromycin B at 100 μg/ml.

To identify transformants with homologous and ectopic integration of the KO vector, DNA was extracted and digested with BamHI (CgChsI), AccI (CgChsIII), or HindIII (CgChsV). After electrophoresis, DNA was blotted onto positively charged nylon membranes (Roche Diagnostics) by an alkaline downward procedure (Brown 1999), and hybridized with a DIG-labeled probe, as suggested by the manufacturer (DIG user’s guide; Roche Diagnostics). Probes of 696 bp (CgChsI), 780 bp (CgChsIII), and 560 bp (CgChsV) were amplified with primer combinations CHSII1 and CHSII3 (5′-CGC GTT GGG CTG TGT AAA GG-3′; CHSII6 (5′-GGT TCA GAC AAG ACT GAG CCG C-3′) and CHSII7; and CHSV2.1 (5′-CCG ACA TGG CAC CGG AC-3′) and CHSIV1 (5′-GCC GAG CTT CCT TGT TC-3′)). Chemiluminescence was visualized on Hyperfilm ECL (Amersham, Pharmacia, Freiburg, Germany) after adding CSPD (1% [vol/vol] in detection buffer) (Roche Diagnostics).

**RNA isolation and gene expression analyses.**

Total RNA was isolated following the protocol of Chomczynski and Sacchi (1987).

RT-PCR was performed with the OneStep RT-PCR kit (Qiagen). The total volume of 10 μl contained 115 to 230 ng of RNA, 0.6 μM each primer, 0.4 mM dNTPs, and 0.4 μl of the enzyme mix (OmniScrip, SensiScrip reverse transcriptase, and HotStar Taq-polymerase) (Qiagen).

To exclude DNA contaminations, RNA was incubated with RNase-free DNase (Promega GmbH, Mannheim, Germany). As an additional control, primers flanking an intron were used in RT-PCR analyses.

For CgChsI, the primer combination CHS14 (5′-CGG TAT CCG TAA CTT CTT CCT C-3′) and RT21.2 (5′-TAT AGT TTG GTG TAT GTG GGG-3′) made possible the amplification of a 458-bp fragment with mitochondrial (m)RNA and a 506-bp fragment with DNA as template. For CgChsIII, the primers CHSIII1 and CHSIII3 (5′-TGA TTC TGT GAG ACT GTA TGC G-3′) yielded a 308-bp fragment with mRNA and a 446-bp fragment with DNA as template. For CgChsV, the primer pair CHSVko (5′-TCT CTC ACA CAT ACA CTT C-3′) and CHSV13 (5′-TTG TGC CCG CCA TCA GGT AGT A-3′) made possible the amplification of a 465-bp fragment with mRNA and a 520-bp fragment with DNA as template. Alternatively, the primer pair CHSV3 (5′-TTA CCG GTG TCG ACG TCG CC-3′) and CHSVRT2 (5′-CAA GCA GCT GAA GAA TTA CGC C-3′) yielded a 585-bp fragment with an mRNA template. For CgChsVI, the primer combination ChsvIR1 (5′-CCG AGC TGT ACG AAT ATT TGA TG-3′) and ChsvRT2 (5′-GTG CCG CCG TTG GAC AGT ACG-3′) made possible the amplification of a 461-bp fragment with mRNA and a 516-bp fragment with DNA as template.

Reverse transcription was done at 50°C for 30 min, followed by a denaturation step at 95°C for 15 min. Then, 25 to 35 cycles, consisting of denaturation at 94°C for 20 s, annealing at 57°C for 30 s, and elongation at 72°C for 45 s, were followed by final elongation at 72°C for 10 min.

In order to confirm intron positions, cDNAs of the complete CHS genes were sequenced.

All primers used in this study were from MWG Biotech AG (Ebersberg, Germany).

**Inhibitor studies with nikkomycin Z.**

Nikkomycin Z (Sigma) was added to conidial suspensions at different concentrations (0 to 500 μM) prior to inoculation of polyester transparency sheets or maize leaf segments as described above. In vitro differentiated infection structures were evaluated 24 h after inoculation of transparency sheets; symptoms developed on leaf segments were photographed 3 to 4 days postinoculation.

**Measurement of hyphopodial force exertion by optical waveguides.**

Force measurement was performed with optical waveguides as described previously (Bechinger et al. 1999). Because ΔCgChsV mutants produced neither falcate conidia nor intact hyphae in the absence of osmolytes, pieces of mycelia (approximately 1 mm²) were placed onto optical waveguides in 0.5 M sucrose. Under these conditions, both wild-type and ΔCgChsV mutant differentiated hyphopodia, and force was measured 24 h after inoculation of the waveguides.

**Microscopy and chitin staining.**

Microscopy was done with either a Nikon Eclipse E600 microscope (Nikon, Düsseldorf, Germany) equipped with bright-field and differential interference contrast optics or a Zeiss Axioplan Imaging II microscope (Oberkochen, Germany) and standard rhodamine or FITC filters. Digital images were taken with a DS-5M (Nikon) or a CoolSNAP-HQ CCD camera (Photometrics, Tucson, AZ, U.S.A.) controlled by the imaging software Metamorph (Universal Imaging, Downingtown, PA, U.S.A.). Image processing was done with the software package Lucia 4.61 (Nikon) and the imaging software MetaMorph. For staining of chitin, cells were grown overnight in CM liquid medium supplemented with 0.5 M sucrose, fixed with 3% formaldehyde for 10 min, and incubated with TRITC-labeled WGA following described protocols (Wedlich-Söldner et al. 2000).

**ACKNOWLEDGMENTS**

The financial support of the fungal cell wall biogenesis projects in H. B. Deising’s lab by the Deutsche Forschungsgemeinschaft (De 403/7-1) and by CAPES Brazil to J. A. Sugui is gratefully acknowledged. We thank C.
Bechinger, University of Stuttgart, Germany, and M. Bastmeyer, University of Karlsruhe, Germany, for hyphopodial force measurement, and D. Jany and A. Beutel for excellent technical assistance.

LITERATURE CITED


AUTHOR-RECOMMENDED INTERNET RESOURCE
Cambridge Medical Research Council Laboratory of Molecular Biology
Myosin Home Page: www.proweb.org/myosin