**Variation in growth rates and aggressiveness of naturally occurring self-fertile and self-sterile isolates of the wilt pathogen *Ceratocystis albifundus***

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*Ceratocystis albifundus* is the most important fungal pathogen of black wattle (*Acacia mearnsii*) grown in plantations in southern and eastern Africa. It is a homothallic fungus but also undergoes unidirectional mating type switching. As a result, the ascospore progeny can be either self-fertile or self-sterile. The only apparent difference between these mating types is the deletion of the *MAT1-1* gene in self-sterile isolates. There is some evidence suggesting that self-sterile isolates grow more slowly than self-fertile isolates, but this has not been tested rigorously. The aim of this study was to determine whether self-sterile isolates are less fit by examining growth rate, relative germination rate and pathogenicity. Five self-sterile isolates were generated from each of five self-fertile isolates of *C. albifundus* and these 30 isolates were compared. The results showed that the self-sterile isolates grew consistently slower and were less pathogenic than the self-fertile isolates. The germination ratio of self-fertile to self-sterile isolates from single ascospores collected from the ascomata of five self-fertile isolates was on average 7:3. This could be a consequence of the self-sterile isolates having a lower germination rate. This observation, and the lower growth and pathogenicity levels, suggests that self-sterile isolates are not likely to compete effectively in nature, raising intriguing questions regarding their role and value to *C. albifundus* and other fungi having a similar mating system.

**Keywords**: *Ceratocystis albifundus*, fitness, self-fertile mating type, self-sterile mating type

**Introduction**

*Ceratocystis albifundus* is an important fungal pathogen, posing a considerable threat to forestry plantations based on Australian *Acacia mearnsii* trees. This fungus is the causative agent of black wattle wilt disease in southern and eastern Africa (Morris et al., 1993; Wingfield et al., 1996; Roux & Wingfield, 1997; Roux et al., 2005). Disease symptoms that develop after infection by *C. albifundus* include the formation of cankers and lesions in the bark, gum exudation and streaked discolouration of the vascular tissue. Infected trees can wilt and die within 6 weeks after infection (Roux et al., 1999).

Mating in ascomycetes is strictly controlled by a single mating locus with two allelic or idiomorphic forms, *MAT1-1* and *MAT1-2*, which represent alternate alleles that differ in length, sequences and structure (Turgeon et al., 1993; Coppin et al., 1997). *Ceratocystis albifundus* is a homothallic fungus that is capable of selfing, enabling it to produce sexual offspring without having a partner of opposite mating type. This is possible because fertile isolates have all three *MAT* genes: *MAT1-1-1*, *MAT1-1-2* and *MAT1-2-1*. Furthermore, unidirectional mating type switching can occur during meiosis, giving rise to self-sterile mating type strains carrying only *MAT1-1-1* and *MAT1-1-2* genes as a result of the deletion of the *MAT1-2-1* gene. The *MAT1-2-1* gene is entirely lost from the genome and these isolates are thus unable to revert back to their self-fertile status (Perkins, 1987; Harrington & McNew, 1997; Witthuhn et al., 2000; Wilken et al., 2014). Wilken et al. (2014) further showed that self-fertile isolates of *C. fimbriata* are genetically identical to the parent strain, while the only apparent difference between the self-sterile isolates and the parent isolate is that they lack the *MAT1-2-1* gene from their entire genome.

Homothallic fungi do not require a compatible mating partner for sexual reproduction due to the ability to self-mate with their own mitotic descendants, giving rise to genetically identical progeny (Lin & Heitman, 2007; Billiard et al., 2011). In this regard, the haploid selfing in homothallic mating systems could result in low levels of genetic diversity and a likelihood of inbreeding depression. Nevertheless, many fungi are homothallic and this form of reproduction is common in nature (Nelson, 1996; McGuire et al., 2001; Gioti et al., 2012). In the case of unidirectional mating type switching, it might be argued that the self-sterile isolates are forced to mate with other
isolates, thus increasing genetic diversity within a population, although fungi having the homothallic mating system do have the potential to outcross. There is, however, some evidence that the self-sterile isolates are less fit than self-fertile isolates (De Beer, 1994; Harrington & McNew, 1995, 1997; Roux, 1996; Wittthuhn et al., 2000), but this has not been tested for Ceratocystis species.

Fitness of microorganisms is characterized by phenotypic and genotypic traits heritable from generation to generation through successful reproduction, and consequently successful survival in a given environment (Pringle & Taylor, 2002). It has been suggested that growth rate is a simple and convenient means to measure the fitness of fungal pathogens (Pringle & Taylor, 2002). In addition, spore production rate and pathogenicity have been applied as quantitative traits to estimate the fitness of fungal pathogens (Pariaud et al., 2009). Traits such as faster growth and higher levels of pathogenicity give a competitive advantage to individual pathogens, which ultimately facilitates more effective host colonization (Lockhart et al., 2005; Meyer et al., 2010).

There are many phenotypic differences between self-fertile and self-sterile mating type strains in filamentous ascomycetes. These, for example, include ascospore size and dimorphism, and formation of fruiting bodies, as well as more rapid linear growth rates (Webster & Butler, 1967; Uhm & Fujii, 1983; Perkins, 1987; Harrington & McNew, 1995, 1997; Wittthuhn et al., 2000). Specifically, Wittthuhn et al. (2000) reported that self-fertile mating strains in some Ceratocystis species have more rapid linear growth than the self-sterile isolates. In this study, the possible differences in fitness between self-fertile and self-sterile strains of C. albifundus were considered by comparing growth rates, levels of aggressiveness in inoculation studies and percentage germination.

Materials and methods

Collection and identification of isolates

All isolates of the wilt pathogen C. albifundus used in this study (CMW39129, CMW39130, CMW39131, CMW39132 and CMW39133) were collected on the Bloemendal Experimental farm, near Pietermaritzburg in the KwaZulu-Natal Province of South Africa, where A. mearnsii trees are propagated commercially. Pieces of bark bearing characteristic ascomata of C. albifundus were removed from bark flaps on tree stumps of recently felled trees. Cultures were made from the spore drops that were produced at the apices of the ascocarps collected from different trees. These were transferred directly onto separate Petri dishes of each ascospore isolate. Ascospore masses transferred from the apices of the ascocarps were streaked over the surfaces of 35 mm Petri dishes with sterilized surgical scalpel blades, transferred into 1.5 mL Eppendorf tubes and ground to a fine powder with a pestle. Genomic DNA was extracted using PrepMan Ultra (Applied Biosystems) following the manufacturer’s instructions. The quantity and quality of DNA extracted was assessed with a spectrophotometer (ND-1000; NanoDrop) to calibrate the concentration and purity of DNA as PCR templates.

The PCR amplification reactions were conducted on a Veriti 96-well thermal cycler (Applied Biosystems). The total volume of each PCR reaction mixture was 15 μL, containing 0.5 μL genomic DNA, 0.3 μL (10 pm) of each primer (forward and reverse), 3 μL MyTaq PCR buffer (Bioline) and 0.09 μL MyTaq DNA polymerase (Bioline). The PCR cycling profile consisted of an initial denaturation at 97°C for 5 min; 35 cycles of denaturation at 95°C for 1 min, annealing at 55°C for 1 min, and extension at 72°C for 1 min; and a final extension at 72°C for 10 min. Sequencing was performed in both directions using primers ITS1F and ITS4 (White et al., 1990) and the Big Dye Terminator v. 3.1 cycle sequencing premix kit (Applied Biosystems) according to the instructions provided by the manufacturer. Sequences were determined using an ABI PRISM 3100 Genetic Analyzer (Applied Biosystems). Sequencing results were compared with published sequences of C. albifundus, obtained from GenBank, using BioEdit v. 7.0.9.0 sequence alignment editor (Hall, 1999).

Mating type proportion from single ascospore progenies

Single ascospore cultures were made directly from ascocarps produced by the five self-fertile isolates confirmed to be C. albifundus. Ascospore masses transferred from the apices of the ascocarps were suspended in 50 μL isoparaffin solvent, Soltrol 130 (Chemfit) to facilitate dispersal (Whitney & Blauel, 1972) of the hydrophobic Ceratocystis ascospores. About 5 μL of spore suspension was streaked over the surfaces of 3.5% water agar (WA; 35 g L⁻¹ agar, Difco) in 90 mm Petri dishes using a platinum wire loop and incubated at 25°C for 72 h until germination of single ascospores was observed. Agar disks containing individual germwells were excised with an aseptic syringe needle and transferred onto 2% malt yeast extract agar (MYEA; 20 g malt extract and 2 g yeast extract, Biolab, 20 g agar, Difco). Fifty single ascospore isolates were made from each of five parental isolates of C. albifundus and these represented self-fertile and self-sterile mating type progeny. The plates were incubated at 25°C for 2 weeks, and the ratio of self-fertile to self-sterile strains of C. albifundus was calculated.

Both morphological characters and molecular characterization were used to ensure the mating type identity of the self-fertile and self-sterile strains obtained from each of the five parental isolates. The presence or absence of ascocarps was used as an initial indication of the mating type of the isolates (Webster & Butler, 1967). Two primer sets, Albi_MAT1-2_F/Albi_MAT1-2_R and Albi_MAT2-1_F/Albi_MAT2-1_R, were used to amplify the MAT1-1-2 and MAT1-2-1 genes in the MAT locus DNA sequence comparisons as described below. Ceratocystis albifundus can be distinguished from all other Ceratocystis species based on its characteristic light/cream-coloured ascomatal bases, bearing black necks and producing hat-shaped ascospores (Wingfield et al., 1996). The five selected isolates were then identified using sequences of the internal transcribed spacer regions (ITS1, ITS2) and 5.8S rDNA gene regions, as follows.

All C. albifundus cultures were incubated for 2 weeks to facilitate mycelial growth. Mycelium of the fungus was scraped from the surfaces of Petri dishes with sterilized surgical scalpels, transferred into 1.5 mL Eppendorf tubes and ground to a fine powder with a pestle. Genomic DNA was extracted using PrepMan Ultra (Applied Biosystems) following the manufacturer’s instructions. The quantity and quality of DNA extracted was assessed with a spectrophotometer (ND-1000; NanoDrop) to calibrate the concentration and purity of DNA as PCR templates.

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in five representative isolates from each of the mating types (Table 1). The amplification reactions were conducted on a Veriti 96-well thermal cycler. The total volume of the PCR reaction mixtures were 15 μL, containing 0.5 μL genomic DNA, 0.3 μL (10 pm) of each primer (forward and reverse), 3 μL MyTaq PCR buffer (Bioline) and 0.09 μL MyTaq DNA polymerase (Bioline). The thermocycling profile for amplification of MAT genes consisted of an initial denaturation at 97°C for 1 min; 35 cycles of denaturation at 95°C for 15 s, annealing at 55°C for 30 s, and extension at 72°C for 30 s; and a final extension at 72°C for 7 min. The amplification products were analysed by electrophoresis using a Mini-Sub gel GT cell (Bio-Rad) and a PS500X DC Power Supply (Hoefer Scientific) on a 1% agarose gel in 1X TAE buffer (40 mM Tris, 20 mM acetic acid, 1 mM EDTA, pH 8), and then visualized using a Bio-Rad Gel Doc EZ imager and IMAGE LAB v. 3.0 software.

Assessment of growth rates

Five confirmed self-sterile isolates from each of the five self-sterile parental strains, as well as the parental strains themselves, were compared for growth rate. Growth rates were assessed by transferring 5 mm plugs from the edges of actively growing cultures onto the centres of 75 mm Petri dishes containing 2% MYEA. The cultures were then incubated at 25°C for 2 weeks. Two measurements for colony diameter at right angles to each other were taken for each of the 25 self-sterile and five self-fertile isolates, daily for 2 weeks. The average growth and standard deviations were then calculated. The entire experiment was repeated once.

Assessment of aggressiveness

In order to detect possible differences in aggressiveness between self-fertile and self-sterile mating type strains of C. albifundus, inoculation trials were conducted on c. 2-year-old A. mearnsii trees located in a jungle stand near Pretoria East, Gauteng, South Africa. Trees of c. 70 mm diameter were inoculated at c. 1.5 m above ground. A total of 50 trees were inoculated with each of the five parental self-fertile strains (250 trees) and 20 trees were inoculated with each of 25 self-sterile isolates, five of which were derived from each of the five parental strains (total 500 trees). Twenty trees were used as controls by inoculating them with sterile MYEA.

Discs of agar bearing mycelium were taken from the margins of actively growing, 2-week-old cultures with a 7 mm cork borer. For the inoculations, a wound was made on the stems of trees, using a 7 mm cork borer, to expose the cambium and the agar discs overgrown with the test strain were inserted into the wounds, with the mycelium facing the cambium. All inoculation points were covered with masking tape to prevent desiccation of wounds, with the mycelium facing the cambium. All inoculated trees were assessed after 6 weeks during December 2012 and January 2013, respectively, and lesion lengths were measured. Reisolations were made from two trees for each treatment and the resulting fungi were identified using a Bio-Rad Gel Doc EZ imager and IMAGE LAB v. 3.0 software.

Results

Collection and identification of isolates

Ceratocystis albifundus isolates were successfully obtained from the stumps of recently (4 weeks previously) harvested A. mearnsii trees. Fruiting bodies produced on MYEA medium had morphological features typical of C. albifundus, including light-coloured ascomatal bases (Wingfield et al., 1996). This identification based on morphology was supported by ITS sequence data where the maximum percentage identity of the isolates was almost identical (99%) when analysed using BLAST against those of C. albifundus strains in NCBI. All sequence data obtained in this study have been deposited in NCBI (accession numbers KF147144 to KF147148).

Mating type proportion from single ascospore progenies

In total, 50 single ascospore progeny were recovered from each of the five parental self-fertile mating type isolates. The segregation ratio of self-fertile to self-sterile isolates was significantly biased towards the self-fertile mating type for four of the parental strains (CMW39129, CMW39130, CMW39131 and CMW39132). Exceptions were found for isolates CMW39131 in the second trial and CMW39133 in both trials, which both had a segregation ratio close to 1:1 for self-fertile to self-sterile isolation mixtures were 15 μL, containing 0.5 μL genomic DNA, 0.3 μL (10 pm) of each primer (forward and reverse), 3 μL MyTaq PCR buffer (Bioline) and 0.09 μL MyTaq DNA polymerase (Bioline). The thermocycling profile for amplification of MAT genes consisted of an initial denaturation at 97°C for 1 min; 35 cycles of denaturation at 95°C for 15 s, annealing at 55°C for 30 s, and extension at 72°C for 30 s; and a final extension at 72°C for 7 min. The amplification products were analysed by electrophoresis using a Mini-Sub gel GT cell (Bio-Rad) and a PS500X DC Power Supply (Hoefer Scientific) on a 1.5% (w/v) agarose gel in 1X TAE buffer (40 mM Tris, 20 mM acetic acid, 1 mM EDTA, pH 8), and then visualized using a Bio-Rad Gel Doc EZ imager and IMAGE LAB v. 3.0 software.

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The inoculation study was conducted in two separate trials established in November 2012 and in December 2012, respectively. The inoculated trees were assessed after 6 weeks during December 2012 and January 2013, respectively, and lesion lengths were measured. Reisolations were made from two trees for each treatment and the resulting fungi were identified to ensure that the inoculated fungi were responsible for the infections observed (Koch’s postulates).

Statistical analysis

A chi-square analysis was used to test for statistical support of possible differences in the proportion of each mating type emerging from the single ascospore isolations; 95% confidence was accepted as an acceptable level of significance. The value for the degrees of freedom was established based on the number of mating types (self-fertile and self-sterile). The null hypothesis was that no significant differences would be observed in the production of self-fertile and self-sterile mating type progeny from single ascospore isolations. A test of independence was also conducted to determine whether there was a significant difference in each single ascospore isolation trial.

Welch’s two-sample t-test and analysis of variance (ANOVA) and Tukey’s honestly significance difference (Tukey’s HSD) test were used to determine whether there were significant differences in the growth rates or aggressiveness of self-fertile and self-sterile mating types based on a P-value computed using R v. 2.5.1 (http://www.r-project.org/; R Core Team, 2012). In addition, general linear models (GLM) using R v. 2.5.1 were applied to test the hypothesis that the ability of the self-fertile mating type isolates to grow faster is closely correlated with a higher level of aggressiveness compared to the self-sterile mating type isolates.

Table 1 Primers used for amplification of MAT genes

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>Product size (bp)</th>
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</thead>
<tbody>
<tr>
<td>Albi,MAT1-2_F</td>
<td>ATAGCAAGGGTATCGGCTC</td>
<td>834</td>
</tr>
<tr>
<td>Albi,MAT1-2_R</td>
<td>GCCGTGCAGAGATCCTA</td>
<td>586</td>
</tr>
<tr>
<td>Albi,MAT2-1_F</td>
<td>CCCCTTCATTGGCCCAT</td>
<td>596</td>
</tr>
<tr>
<td>Albi,MAT2-1_R</td>
<td>CATCAAGTCTGTCATCCA</td>
<td>596</td>
</tr>
<tr>
<td>Albi,MAT2-1_F</td>
<td>CATCAAGTCTGTCATCCA</td>
<td>596</td>
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</tbody>
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Gene sequence data where the maximum percentage identity of the isolates was almost identical (99%) when analysed using BLAST against those of C. albifundus strains in NCBI. All sequence data obtained in this study have been deposited in NCBI (accession numbers KF147144 to KF147148).
mating types (Table 2). The greatest deviation from a 1:1 segregation ratio was found for isolates CMW39130 and CMW39131 in the first trial and CMW39132 isolate in the second trial, where this ratio was c. 3:1. Overall, the segregation ratio of self-fertile to self-sterile was c. 1/5 on average (Table 2). The bias of segregation towards the self-fertile mating type was supported statistically, based on chi-square analysis at a 95% confidence level (Table 2).

Assessment of growth rates
All the self-fertile isolates exhibited more rapid growth in comparison to the five self-sterile isolates derived from each of them, although isolates representing each of the two mating types had similar growth rates and produced similar growth patterns on MYEA at 25°C (Fig. 1). These observations were also strongly supported statistically (Welch’s two sample t-test, P < 0.001).

Assessment of aggressiveness
In both inoculation trials, self-fertile isolates consistently produced longer lesions than the self-sterile isolates derived from them (data shown only for the second trial which were identical to those of the first trial; Figs 2 & 3). Self-fertile strains resulted in distinct black lesions on the stems, gum exudation and streaking in the cambium 6 weeks after inoculation. In contrast, self-sterile strains were not found to be associated with the same severe symptoms. Reisolations from the lesions consistently yielded self-fertile and self-sterile isolates morphologically indistinguishable from the inoculated isolates. The hypothesis that there is a significant difference in aggressiveness between self-fertile and self-sterile isolates was statistically supported by analysis of variance (P = 2E–16) and Tukey’s HSD test (P = 0 at the 95% confidence level).

Correlation of growth and aggressiveness
Four measurements including growth rate and aggressiveness of isolates, representing both mating types, were applied to GLM tests available in R v. 2.5.1. The results showed that self-fertile isolates have superior fitness based on growth rate as well as pathogenicity. This was evident from a significant correlation between growth rate and aggressiveness of self-fertile and self-sterile isolates based on GLM tests (coefficient = 4.534, standard error = 1.031, t-value = 4.397, probability (>|t|) = 0.002).

Discussion
This study showed that there are significant differences in growth and aggressiveness between self-fertile parental and self-sterile offspring strains of the wilt pathogen *C. albifundus*. These parental and offspring strains are essentially identical because this fungus is haploid, although minor differences in the strains could have resulted as the consequence of sexual recombination. Self-sterile isolates of *C. albifundus* lack a single gene at the mating type locus. This deletion appears to have resulted in self-sterile isolates being significantly less fit than their self-fertile counterparts.

Fungal pathogens that reproduce sexually would typically be expected to produce equal frequencies of progeny representing the two different mating types (Fisher, 1930). The same results have been reported for

<table>
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<tr>
<th>Isolate</th>
<th>Mating type of successfully germinated spores</th>
<th>Failed to germinate</th>
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<tr>
<td></td>
<td>1st trial Chi-square</td>
<td>2nd trial Chi-square</td>
</tr>
<tr>
<td>CMW39129</td>
<td>27 Self-fertile</td>
<td>13 Self-sterile</td>
</tr>
<tr>
<td>CMW39130</td>
<td>33 Self-fertile</td>
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<tr>
<td>CMW39132</td>
<td>31 Self-fertile</td>
<td>17 Self-sterile</td>
</tr>
<tr>
<td>CMW39133</td>
<td>26 Self-fertile</td>
<td>18 Self-sterile</td>
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*Significantly different from 1:1 ratio at 0.05 probability level.
Ceratocystis coerulescens (Harrington & McNew, 1997). In contrast, isolates of C. albifundus studied here showed a significant bias towards self-fertile mating. A similar bias is known in other fungal pathogens (Kwon-Chung & Bennett, 1978; Wickes et al., 1996; Nieuwenhuis & Aanen, 2012). The bias towards the self-fertile isolates could possibly arise from ascospores that have a reduced capacity to germinate or, alternatively, fewer ascospores representing the self-sterile form might be produced in each individual ascus. This, however, has not been tested.

Growth rate has been shown to represent a simple and efficient means to estimate the relative fitness of fungal populations (Pringle & Taylor, 2002; Schoustra et al., 2009). In this study, evidence for a lower level of fitness in self-sterile isolates was supported by the growth tests. All self-fertile mating type strains showed more rapid growth than self-sterile mating type strains. Although this has been suggested previously (De Beer, 1994; Harrington & McNew, 1995, 1997; Roux, 1996; Witthuhn et al., 2000), the current study is the first where molecular tools have been used to confirm the mating type of strains beyond doubt. The only apparent difference between the self-fertile and self-sterile isolates might be the deletion in the MAT1-2-1 gene. This seems to be the

![Figure 2: Mean lesion lengths on Acacia mearnsii trees 6 weeks after inoculation with self-fertile (-SF) and self-sterile (-SS) isolates of Ceratocystis albifundus, sterile agar as control. The bar represents standard deviation of the mean.](image-url)
most plausible reason for the reduction in growth, although the mechanism is not understood.

Extensively repeated inoculation tests using a large number of isolates showed that self-fertile isolates of *C. albifundus* are significantly more aggressive than self-sterile isolates. Likewise, growth rate for the test isolates was positively correlated with aggressiveness of isolates. A correlation between growth rate and pathogenicity has been shown for various other ascomycete fungal pathogens (Belisario et al., 2008; Brasier & Kirk, 2010). Furthermore, it has also been suggested that differences in pathogenicity can be attributed to the involvement of mating type genes in fungi (Lockhart et al., 2005; Zhan et al., 2007). However, in both these examples, and unlike *C. albifundus*, the fungi were heterothallic and thus the mechanism underpinning the differences in pathogenicity are likely to be different.

There are very few fungi that undergo unidirectional mating type switching like that found in the tree pathogen *C. albifundus*. Although the evolutionary advantages for this irreversible switching are not understood, the fact that distinct differences in fitness of self-fertile and self-sterile isolates have been shown suggests that such an advantage exists. Unidirectional mating type switching is also a common feature in species of *Ceratocystis* as defined by De Beer et al. (2014) and related to *C. fimbriata* (Harrington & McNew, 1995, 1997; Wilken et al., 2014) and it is likely that similar differences in fitness of self-fertile and self-sterile isolates exist in species in this genus other than *C. albifundus*. Logically, maintaining this system, which would require significant amounts of energy to be channelled into producing progeny with low levels of fitness, would be counterproductive. Nevertheless, there may be some advantages to be gained from this strategy. For example, the self-sterile isolates may increase the genetic diversity in a population by fully exploiting the benefits of sexual recombination. This might be particularly valuable to species that have the potential to self-cross, although homothallic isolates also have the potential to outcross. Future studies should, therefore, consider the level of outcrossing that occurs in *Ceratocystis* species in nature.

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