Nuclear rDNA phylogeny in the fungal genus *Verticillium* and its relationship to insect and plant virulence, extracellular proteases and carbohydrases

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Phylogenetic relationships among 18 isolates in the genus *Verticillium*, representing 13 species of diverse ecomnutional groups (pathogens of insects, plants, mushrooms, nematodes and spiders, and saprobes), were examined by using sequences from the internal transcribed spacer (ITS) and small nuclear (NS) rRNA regions. The isolates were also assessed for their abilities to infect insect larvae (Galleria mellonella) and to cause necrosis in alfalfa (Medicago sativa), and for their proteolytic, chitinolytic and pectinolytic activities. The phylogenetic data suggested that *Verticillium* is polyphyletic in origin and is therefore a form genus. However, the phylogenetic tree supported the plant pathogens (*V. dahliae*, *V. albo-atrum* and *V. nigrescens*) as a clade. The alfalfa isolate of *V. albo-atrum* (isolate 595) was an interesting outlier to the main body of plant pathogens as it clustered with the insect pathogen *V. indicum*. Strains of *V. lecanii* and *V. indicum* were able to infect insects and are present in divergent groups in the consensus tree, suggesting that the ability to infect insects may have evolved independently many times. Similarly, the nematophagous *Verticillium* species appear to have evolved independently along several different routes and one isolate, *V. chlamydosporium*, was able to infect insects. *V. albo-atrum*, *V. nigrescens* and *V. dahliae* all produced high levels of enzymes capable of degrading pectin, a major component of plant cell walls. The ability to excrete pectinase was a broad indicator of the ability to produce lesions on alfalfa. In the plant pathogens, the functions of a broad-spectrum protease were assumed by trypsins which degrade Bz-AA-AA-Arg-NA substrates (Bz, benzoyl; AA, various amino acids; NA, p-nitroanilide). The insect pathogens and mushroom pathogen (*V. fungicola*) were characterized by production of high levels of subtilisin-like proteases active against a chymotrypsin substrate (succinyl-Ala,-Pro-Phe-NA) and the inability to clear pectin. The insect and mushroom pathogens, and several nematode pathogens, were distinguishable from the plant pathogens in their ability to produce chitinases.

Keywords: entomopathogenic fungi, phytopathogenic fungi, nematophagous fungi, saprophytic fungi, rDNA phylogeny

Abbreviations: ITS, internal transcribed spacer; MEPT, multiple equally parsimonious tree; NA, p-nitroanilide; NS, small nuclear; Suc, succinyl; Bz, benzoyl.
The GenBank accession numbers for the sequences reported in this paper are AF10847-AF108500 and AF110529-AF110531.
INTRODUCTION

The genus *Verticillium* Nees contains a heterogeneous group of asexual fungi, many of which are of considerable importance in agriculture as pathogens of plants, insects and nematodes. The taxonomy of the group is still being reviewed and many species have a difficult and controversial characterization (Evans & Samson, 1986; Rowe, 1995). To date, species of *Verticillium* have been characterized primarily on the basis of morphology and by virulence on certain hosts. The two most commercially important of the *Verticillium* plant pathogens, *V. albo-atrum* and *V. dahliae*, cause vascular wilts in many hosts worldwide. The wide host range with little host specificity distinguishes *Verticillium* fungi from many other plant pathogens. As a result, few physiological races of these plant pathogens have been found and characterized (Rowe, 1995).

Several other *Verticillium* species are being studied as potential biocontrol agents for insect and nematode pests. These *Verticillium* spp. have a wide host range with little host specificity. Thus, the insect pathogen *V. lecanii* and the nematode pathogen *V. chlamydosporium* are hyperparasites on other fungi (Leinhos & Buchenauer, 1992). The pathogens of fungi, *V. fungicola* and *V. lamellicola*, are also pathogenic to mites and insects (Balazy et al., 1987). In addition, morphological differences between *Verticillium* species are minor, with virtually no differences in conidial morphology. Nevertheless, in a recent study, strains of *V. albo-atrum* and *V. lecanii* were found to show enzymatic adaptation to the polymers present in the integuments of their plant and insect hosts respectively, implying pathogenic specialization to the requirements of these ecological niches (St Leger et al., 1997). This suggests that *Verticillium* spp. could be a useful model in the study of the evolution of pathogenic mechanisms.

Several current problems in our understanding of *Verticillium* fungi have resulted from the inability to recognize the phylogenetic relationships within this group (Jun et al., 1991). Because of this uncertainty, and because we are interested in understanding the genetic basis for the evolution of pathogenicity, it is important to develop a molecular phylogeny of this group that is independent of morphology. Previous biochemical and molecular studies have focused almost exclusively on the two major plant pathogens *V. albo-atrum* and *V. dahliae*. Methods used for diagnosis of these species include vegetative incompatibility analysis (Joaquim & Rowe, 1990), random amplification of polymorphic DNA (RAPD; Barasubie et al., 1995), and restriction fragment length polymorphisms (RFLP; Okoli et al., 1993, 1994; TYPAS et al., 1992).

Recently, the genetic diversity in *V. dahliae* and *V. albo-atrum* was evaluated using PCR technology and exploiting variations in rRNA or mitochondrial rRNA genes (Griffen et al., 1997). Li et al. (1994) used a pair of conserved PCR primers to amplify a region of the mitochondrial small rRNA gene. Employing this fragment in an RFLP analysis differentiated 10 species of *Verticillium* and allowed development of a *V. dahliae*-specific primer. Robb et al. (1993) utilized the nucleotide variability in the internal transcribed spacer (ITS) regions of nuclear rRNA to develop specific PCR-based assays. The ITS regions between 18S and 28S rDNA were amplified from *V. albo-atrum*, *V. tricolor* and *V. dahliae*, cloned and then sequenced. Primers were then synthesized based on specific nucleotide differences found in the ITS regions that were characteristic of each species (Hu et al., 1993; Moukhamedov et al., 1994; Nazar et al., 1991; Robb et al., 1993). A similar technique has been applied to resolve the genetic variation between isolates of *V. chlamydosporium* using restriction fragment analysis of PCR-amplified ITS and the intergenic spacers (IGS) that separate ribosomal repeat units (Arora et al., 1996). These studies were mostly focused on providing a tool for unambiguous identification and enumeration of individual *Verticillium* species. It seems evident that rDNA is sufficiently variable to provide characters for phylogenetic analysis of this diverse genus. In particular, because it contains interspersions of variable and more highly conserved regions that represent differing rates of evolution, rDNA may allow differentiation and detection at a wide range of specificities.

In the present study we used the ITS1 (internal transcribed spacer) region and a portion of the relatively more conserved small nuclear (NS) region of rDNA to provide phylogenetic information for a fairly broad range of *Verticillium* species. In contrast to the previous studies on *Verticillium* we chose to analyse rDNA sequences as this strategy provides much more phylogenetic information than comparing RFLP banding patterns of the amplified regions of rDNA. We also investigated the relationship of phylogeny with the production of hydrolytic enzymes (putative pathogenicity/virulence determinants) and pathogenicity against a plant (alfalfa) and insect (waxworm larvae) species, with the aim of gaining insight into the evolution of *Verticillium* pathogenicity, and facilitating the design and implementation of more effective screening programmes for biocontrol agents.

METHODS

Fungal strains, cultivation and DNA extraction. The *Verticillium* isolates used in the analysis are listed in Table 1. Cultures were maintained on 15% (w/v) agar containing 0.2% yeast extract, 1% peptone, 2% glucose (YPD) at 22 °C. In liquid cultures, 50 ml YPDY (0.1% (w/v) yeast extract, 1% peptone, 2% glucose, 0.1% yeast nitrogen base) was inoculated with conidia taken from 7–14-d-old agar plates. The 250 ml Erlemeyer flasks were incubated with shaking (100 r.p.m.) at 22 °C for up to 7 d and then aseptically collected on Whatman no. 1 filter paper under vacuum filtration and stored at −70 °C.

Genomic DNA was extracted by grinding the mycelium into a fine powder with a mortar and pestle (approx. 5 min) under liquid nitrogen. A sample of the powder was transferred to a tube containing 4 ml lysis buffer (0.1 M Tris, 0.1 M EDTA, 2% SDS, pH 8.0 with NaOH) and incubated at 65 °C for 1 h. The mixture was centrifuged at 7000 r.p.m. for 10 min and the
Table 1. Species, strains and origins of fungi used in this study

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain no.</th>
<th>Isolation/host</th>
<th>Geographical origin</th>
<th>Source*</th>
</tr>
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<tr>
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<td>10</td>
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</tr>
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<td>Alfalfa</td>
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<td>Spider</td>
<td>Solomon Isl.</td>
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<tr>
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<td>ARSEF</td>
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<tr>
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<td>Lagria vilosa (Coleoptera)</td>
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<td>ARSEF</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>Lymantria dispar (Lepidoptera)</td>
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<td>UAMH</td>
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<td>4785</td>
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<td>1691</td>
<td>Wood saprophy</td>
<td>Canada</td>
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*UAMH, University of Alberta Microfungus Collection and Herbarium; ARSEF, USDA Entomopathogenic Fungus Collection at Cornell University.

DNA amplification and sequencing. A region of the NS rDNA gene and a portion of the ITS1 region were amplified using the primer pairs NS7 and NS8, and ITS1 and ITS2, respectively, and an MJ Research Minicycler with protocols outlined by White et al. (1990). The PCR products were analysed by electrophoresis on 0.8% agarose gels. After determination that the PCR products each contained one band, they were purified (Qiaquick PCR purification kit, Qiagen) using methods outlined in the protocol. Sequencing was performed at the Queen’s University Biochemistry DNA sequencing facility (Kingston, ON) with the same primers as used for PCR.

Sequence analysis and phylogenetics. Two sequences of rDNA each were obtained for 18 taxa of Verticillium. The region of the NS rDNA sequence, a conserved region, was approximately 350 bp in length for all taxa. The ITS1 region, a relatively non-conserved region, was approximately 650 bp in length for all taxa. Constructing phylogenies using molecular data requires five basic steps. Initially the sequences must be aligned, the informative characters tested for phylogenetic signal, the data analysed, a tree selected and, finally, nodal support evaluated (Hillis et al., 1993). The two sets of data were first analysed separately. The sequences were pre-aligned using SeqEd (Applied Biosystems, 1992) and handled independently throughout the following procedures. The sequences were transferred from SeqEd to ESEE3 (Cabor & Beckenbach, 1989), a program that allows visual alignment. The ITS1 region contained many gaps and insertions, making realignment necessary. The NS rDNA region had only one base-pair gap and the alignments with both methods were identical. Any unalignable regions were removed from the analysis because positional homology cannot be assured (Swofford & Olsen, 1990). The sequences were transferred into a MacClade (Maddison & Maddison, 1992) readable format. Any uninformative characters (simple isomorphs) were then deleted, leaving 45 characters for the ITS1 regions and 78 NS rDNA characters. To determine if phylogenetic signal was apparent, the matrices were transferred to PAUP (Swofford, 1991) and the random trees option selected. One thousand random trees were generated and the G1 statistics produced. This value was compared against a G1 table to determine significance (Hillis & Huelsenbeck, 1992). PTP values were determined by transferring the matrices to a Pennsyl (Farris, 1988) format and running PTP analysis in random cladistics (Siddell, 1996). One hundred replicates were assessed.

The MacClade matrices were transferred to PAUP in order to determine the most parsimonious resolutions of the data. Aspergillus niger was designated as the outgroup, and branch and bound searches were performed. All characters were unordered and initially equally weighted. If the outcome revealed multiple equally parsimonious trees (MEPTs) then the characters were reweighted based on the rescaled consistency index and the search repeated. This process was repeated until the number of MEPTs could no longer be reduced. Consistency and retention indices were noted. If there were still MEPTs, strict, semi-strict, 50% majority rule
and Adams consensus trees were analysed. One of the MEPTs was then chosen that matched either the semi-strict or 50% majority rule tree and was used for the analysis.

Nodes on most resulting phylogenies had bootstrap values and the number of characters defining that node. The bootstrap values were obtained by transferring the data to a Hennig86 file (Farris, 1988) and running the Heyjoe program in random cladistics. One hundred replicates were performed and the values placed at the appropriate node on the phylogeny produced above.

The characters were assessed in MacClade using the trace character tool. Each character was assessed for appearance on the phylogeny and whether it was an unreversed or homoplastic character. Total number of characters and number of unreversed characters were placed on each node of the phylogeny.

The above procedure was repeated for the ITS region and NS rDNA sequences with the transversion:transition ratio changed from 1:1 to 5:1. This weighs transversions (i.e. purine to pyrimidine) five times greater than transitions. PTP and bootstrap values were not calculated because the Hennig86 cannot compute ratios different from 1:1. The resulting topologies were compared with the 1:1 ratio phylogenies produced above. The consistency index (CI) and retention index (RI) values were not calculated for these phylogenies. Total number of characters and the number of transversed characters are noted at each node.

The 78 rDNA and 45 ITS characters were combined into one matrix in MacClade and the entire analysis repeated. Thus, two combined phylogenies were produced, one unordered and the other with a transversion:transition ratio of 5:1. Nodal support values were identical to those previously determined.

**Insect virulence.** Wax moth larvae (Galleria mellonella) were allowed to crawl over sporulating cultures of each strain of *Verticillium* (eight larvae per strain) for 15 min. They were then placed individually into 100 mm Petri dishes containing moistened filter paper. After 1 d, larvae were provided with artificial diet (Carolina Biological Supplies) and monitored daily for 14 d. Larvae were also provided with water agar for 3 d to detect potential production of hyphae and conidia. If no external evidence of fungal growth was detected, cadavers were dissected and samples examined microscopically to document hyphal bodies in the haemocoel and fat body.

**Plant virulence.** We adopted a laboratory method based on artificial infection of detached leaves as this has been found to be an efficient method of evaluating resistance of alfalfa plants to *V. albo-atrum* (Huang, 1989). Stem cuttings were excised from 7-week-old alfalfa plants (Medicago sativa, var. Iroquis) and surface disinfected (0.25% sodium hypochlorite, 2 h, and four changes of sterile distilled water). The cuttings were placed on 1% water agarose (containing 0.05% yeast extract and 0.2% (dry weight) colloidal chitin adjusted to pH 5.5. Plates were prepared as described by Harkin & Anagnostakis (1975) with an underlay of 15% agar and a 3 mm overlay of the chitin medium. Pectinase activity was detected using minimal medium supplemented with polymethylgalacturonan (pectin) buffered at pH 8.0 (0.05 M HEPES) and containing calcium-rich agar (Durrands & Cooper, 1988). Pectinase activity was detected on this medium by addition of 1% (w/v) cetyltrimethylammonium bromide, which precipitated undegraded substrate.

**Protease assays.** Conidia (2 x 10⁶) were inoculated into 15 ml cuticle/basal salts medium (St Leger *et al.*, 1997) and incubated with shaking (100 r.p.m.) at 23 °C for 3 d. Protease production was evaluated by assaying culture filtrates (Whatman no. 1 filter paper) against a range of substrates previously used to detect subtilisins and trypsins in insect and plant pathogens (St Leger *et al.*, 1987, 1997).

**RESULTS**

**rDNA phylogenetic analysis of *Verticillium* species**

Fig. 1 is one of the MEPTs and also the 50% majority rule tree. The total tree length was 1273 characters (when all characters were reweighted by 10). All data matrices were shown to contain phylogenetic signal from both the G1 statistic and PTP methods (*P* < 0.001). The combined data set included 123 informative characters. Six MEPTs were produced after successively weighting the characters. The isolates within the *V. lamellicola*, *V. lecanii* and *V. coccosporum* species were each resolved as sister taxa in this phylogeny. The two *V. albo-atrum* isolates could be made monophyletic but only with an addition of 26.5% total tree length. The two *V. fungicola* isolates required an additional 13.5% of the tree length to result in monophyly.

Fig. 2 is one of 12 MEPTs for 20 isolates as well as the strict consensus tree when the combined data set was analysed with a transition:transversion of 5:1. The total tree length was 6070 characters (when all characters were reweighted by 10). There are some differences between this tree and the unordered tree (Fig. 1). *V. fungicola* 895 is resolved as the sister to *V. chlamydo-sporium*. Contrary to most of the unordered MEPTs, the *V. dahliae*, *V. albo-atrum* 5393 and *V. nigrescens* clade now resolved as the sister group to the clade including *V. araneoram* on all 12 MEPTs. When the unordered tree (Fig. 1) is altered to the transversion:transition topology only one additional step is required (0.4% difference), indicating that areas of the unordered phylogeny were unstable. As in Fig. 1, the isolates within the *V. lamellicola*, *V. lecanii* and *V. coccosporum* species were each resolved as sister taxa in this phylogeny. However, to make the *V. albo-atrum* isolates monophyletic required a 19.4% change in tree length, and to produce monophyly in the *V. fungicola* isolates required a 12.5% tree length change.
Verticillium phylogeny

Fig. 1. One of the six MEPTs, the 50% majority rule, produced from all informative characters (ITS plus a region of the NS rDNA) for 19 taxa of fungi. Values given at each node represent the bootstrap value (ratio of total number of characters:number of unreversed characters).

Verticillium phylogeny

Fig. 2. One of the 12 MEPTs, the 50% majority rule, strict and semi-strict consensus of 123 informative characters (ITS plus a region of the NS rDNA) for 19 taxa of fungi. A transversion:transition ratio of 5:1 was used. Values given at each node represent the number of characters (number of transversion characters).

Enzymic diversity of Verticillium spp.

Extracellular protease activity on four substrates was determined in Verticillium isolates (Table 2). Isolates of the insect pathogens, V. lecanii and V. indicum, and the mushroom pathogen, V. fungicola, cleaved the subtilisin substrate Suc-Ala-Ala-Pro-Phe-NA faster than Bz-Phe-Val-Arg-NA, and except for V. indicum, possessed little or no activity against Bz-Val-Gly-Arg-NA or Bz-Pro-Phe-Arg-NA, indicating that the trypsins of these species require a specific substrate sequence (Phe-Val-Arg). The nematode pathogens, V. coccosporum, V. lamelicola and V. chlamydosporium, and the spider pathogen, V. nr. araneaurn, were distinguishable from the insect pathogens by producing lower levels of subtilisin-like proteases as compared to the trypsin activity, although these had the same narrow specificity as the enzymes from the insect pathogens. In contrast the plant pathogens (V. albo-atrum, V. dahliae and V. nigrescens) hydrolysed all three trypsin substrates and cleaved the most susceptible of these (Bz-Val-Gly-Arg-NA) at least twofold faster than the subtilisin substrate, Suc-Ala-Ala-Pro-Phe-NA.

Chitinase and pectinase production by Verticillium strains is shown in Table 3. Chitinase production by strains of Verticillium species determined in plate clearing assays correlated with the ability to produce high levels of subtilisins and narrow-spectrum trypsins in liquid culture. Thus, V. fungicola, a mushroom pathogen, and the insect and nematode pathogens produced clearing zones in chitin medium while the plant pathogens did not, indicating that if chitinases are secreted by this group they were at a level below the sensitivity of the assay. By contrast, V. coccosporum and V. chlamydosporium, and the plant pathogens, were all able to degrade pectin (Table 3). Isolates of the insect pathogens and V. fungicola did not produce detectable levels of pectinase.
**Table 2. Extracellular protease activity and substrate specificity of the *Verticillium* species**

Protease activity was measured spectrophotometrically in culture filtrates after 3 d growth in 1% cuticle/basal salts medium. Results in columns A–E represent mean percentage protease activities for each species. Substrates: A, Suc-(Ala)₃-Pro-Phe-NA; B, Bz-Phe-Val-Arg-NA; C, Bz-Val-Gly-Arg-NA; D, Bz-Pro-Phe-Arg-NA. Results in column E are the absolute values in nmol NA released min⁻¹ ml⁻¹ corresponding to 100% activity (St Leger et al., 1987, 1997).

<table>
<thead>
<tr>
<th><em>Verticillium</em> species*</th>
<th>Percentage of maximum protease activity by substrate:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td><em>V. albo-atrum</em> 595</td>
<td>28</td>
</tr>
<tr>
<td><em>V. albo-atrum</em> 5393</td>
<td>26</td>
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<tr>
<td><em>V. nr. araneatum</em> 1905</td>
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<tr>
<td><em>V. coccosporum</em> 2064</td>
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<tr>
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<td>100</td>
</tr>
<tr>
<td><em>V. nigrescens</em> 6687</td>
<td>53</td>
</tr>
</tbody>
</table>

*V. balanoides* 3173, *V. leptobactrum* 5951 and *V. rexianum* 1691 grew poorly in 1% cuticle/basal salts medium.

**Table 3. Pectinase and chitinase activities of the *Verticillium* species and virulence against wax moth larvae and alfalfa**

Enzyme activities were calculated as an index of the total diameter of the colony plus the clear zone around it divided by the diameter of the colony. A value >1.0 indicates that the isolate is releasing the enzyme into the surrounding medium. Virulence against waxworm larvae is shown as LT₅₀ in d; virulence against alfalfa is shown as the diameter of the zone of necrosis, in mm, after 7 d. All values are means of five measurements. Blank spaces indicate no clearing on pectin or chitin agar plates, LT₅₀ values not lower than the control mortality, or no zone of necrosis.

<table>
<thead>
<tr>
<th><em>Verticillium</em> species*</th>
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<th>Bioassays</th>
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</tr>
<tr>
<td><em>V. nigrescens</em> 6687</td>
<td>2.89</td>
<td>1.01</td>
</tr>
</tbody>
</table>

* No detectable pectinase or chitinase activities, or trace alfalfa necrosis and no insect virulence, was observed from *V. nr. araneatum* 1905, *V. rexianum* 1691, *V. balanoides* 3173, *V. lamellicola* 2055 and 4785.
Pathogenicity profiles

Variations in pathogenicity and host range among the isolates of *Verticillium* species were determined against an insect (the wax moth, *Galleria mellonella*) and a strain of alfalfa susceptible to wilt disease (Table 3). Pathogenicity against insects was evaluated by externally inoculating larvae with conidia. Mortality values were very low for non-inoculated controls (<5%). Consequently mortality among treated larvae was almost always due to fungal infection. Of the 18 *Verticillium* isolates tested, four caused mortality and all of these successfully produced hyphae and spores on the cadaver. *V. indicum* was particularly virulent, killing all the test insects within 3 d of inoculation, and then rapidly covering the cadavers in a thick pelt of hyphae. Isolates of *V. lecanii* were also pathogenic, as was an isolate of *V. chlamydosporium* (2218). Insects exposed to other isolates remained symptomless and showed mortality values less than or equal to the non-inoculated controls.

The alfalfa leaves needed to be wounded to allow infection and necrosis of plant tissues. This was expected, as plant-pathogenic *Verticillium* species normally infect via the roots, where wounds may considerably increase the amount of infection (Evans & Gleeson, 1973). Leaves infected with the known plant-pathogenic species (V. *albo-atrum*, V. *dahliae* and V. *nigrescens*) displayed chlorosis after injection of the fungal inoculum (Table 3). Leaflets infected with most of the nematode pathogens, *V. indicum* and V. *fungicola* remained symptomless.

DISCUSSION

Based primarily on morphological criteria, the genus *Verticillium* is divided into four sections and a residual group (Gams, 1988; Gams & Van Zaayen, 1982). The section *Nigrescensia* comprises the plant pathogens and some saprophytic species with dark resting structures. This feature appears to reflect underlying phylogenetic relatedness, as *V. albo-atrum* (5393), *V. dahliae* and *V. nigrescens* were monophyletic in the phylogenetic trees produced (Figs 1 and 2). The section *Prostrata* contains the soil-dwelling and nematophagous species (V. *chlamydosporium*, V. *balanoides*, V. *lamellicola*, V. *leptobactrum* and V. *cocosporum*) as well as other species such as *V. fungicola* and *V. lecanii* with conidiophores at least partly prostrate. These species are polyphyletic, implying that the section *Prostrata* does not reflect evolutionary history. In fact, V. *balanoides* is the sister taxon of *V. rexianum*, a fuscicolous species classified in the section *Albo-erecta*, with erect conidiophores (Gams & Van Zaayen, 1982). This suggests that the conidiophore-erect criterion, a key morphological characteristic for distinguishing among *Verticillium* species (Gams, 1988), is not a useful indicator of phylogenetic relatedness.

In spite of their phylogenetic divergence, the entomopathogens *V. lecanii* and *V. indicum* were all characterized by the production of high levels of subtilisin-like proteases, suggesting that this may be a broadly predictive indicator of entomopathogenicity. However, subtilisin-like proteases also constituted the major protease component secreted by the mushroom pathogen *V. fungicola*. Broad-spectrum subtilisins are the major proteins produced by the insect pathogens *Metharhizium anisopliae* and *Beauveria bassiana* during infection processes, and have much greater ability than the trypsin-like enzymes to degrade insect cuticle (St Leger & Bidochna, 1996). It would be interesting to determine whether subtilisin-like proteases are also involved in degrading certain mushroom tissues. The question arises as to whether the predominance of subtilisins is a derived condition reflecting adaptation to entomopathogenicity, or is retained from a saprophytic ancestor. The latter may be the case since subtilisin-related proteases are the principal broad-spectrum proteases produced by many saprophytes (Gunkle & Gassen, 1989), so they are unlikely to represent the tools specifically developed by entomopathogenic fungi to implement pathogenicity. Also, the basal position of the subtilisin-producing entomopathogen *V. lecanii* at the convergence point of the consensus tree suggests that it is closest to the lineage that gave rise to the other *Verticillium* spp. Although *V. indicum*, *V. chlamydosporium* and *V. lecanii* do not form a monophyletic group, none of the insect or nematode pathogens produced pectinase, indicating independent adaptation by both lineages to an ecological niche where pectinase is not required.

In contrast to the invertebrate pathogens, the plant pathogens *V. albo-atrum* (isolate 5393), *V. dahliae* and *V. nigrescens* are monophyletic in all MEPTs regardless of weight or sequences used. They also exhibited the most evident correlation (as compared to insect and nematode pathogens) between genetic similarity, enzyme production and pathogenicity. Apparently, during adaptation of these *Verticillium* spp. to the plant cell wall the functions of a broad-spectrum protease were assumed by trypsins. As there are no reports of trypsins being produced by saprophytes, trypsin-related enzymes might have a specific role in pathogenicity. The plant pathogen *Cochliobolus carbonum* also produces a broad-spectrum trypsin, which may be involved in degrading hydroxyproline-rich proteins in plant cell walls (Murphy & Walton, 1996). As befits their status as plant pathogens, *V. albo-atrum*, *V. nigrescens* and *V. dahliae* all produced high levels of enzymes capable of degrading pectin, one of the principal components of plant cell walls.

Two separate studies, one employing RFLP (Carder & Barbara, 1991) and the other RAPD analysis (Barasubiye et al., 1995), demonstrated that isolates of *V. albo-atrum* pathogenic to alfalfa were clearly differentiated from other *V. albo-atrum* isolates and warranted a separate taxonomic status. The alfalfa isolate of *V. albo-atrum* (isolate 595) thus provided an interesting outlier to the main body of plant pathogens as the sister taxon to the insect pathogen *V. indicum*. Isolate 595 was clearly more closely related phylogenetically than physiologically to *V. indicum* as it was very similar to *V. albo-
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atrum 5393 in enzyme production. Similarly, V. fungi-
cola 895 resolved in a monophyletic group that included
V. lamellicola isolates while V. fungicola 2065 resolved
in a monophyletic group that included V. coccosporum
isolates; this suggests that a separate taxonomic status
may also apply to some V. fungicola isolates. However,
the V. fungicola isolates were also physiologically
similar to each other in their protease, pectinase and
chitinase profiles. Whereas neither of the V. fungicola
isolates was pathogenic to G. mellonella larvae, strain
2065 was isolated from gypsy moth larvae (Lymantria
dispar), while strain 895 was isolated from a mushroom.
These results indicated that similar enzyme production
and pathogenicity does not always translate into phylo-
genetic relatedness, and vice versa.

Several investigations have identified genetic variability
in some Verticillium spp. and have attempted to
correlate this to virulence. Okoli et al. (1994) utilized
RFLP to differentiate isolates of V. dahliae with differing
pathogenicities on several plant species. V. fungicola
isolates were differentiated by RAPD analysis, and
limited correlations with fungicide resistance and geo-
graphic origin were found, but not with virulence
toward mushroom or colony morphology (Bonnen &
Hopkins, 1997). Beyond this there is little information
on the relatedness between genotype diversity and
pathogenicity to different host groups, e.g. insects,
plants, nematodes or mushrooms. Determining the
differences between closely related plant and insect
pathogens will indicate probable key virulence charac-
ters for pathogenicity of the two host groups and may
help determine how pathogenicity evolved. The phylo-
genetic analysis suggested that the common ancestor
was an insect pathogen, and that plant pathogenicity
evolved secondarily. This is evident by moving more
basically on the phylogeny and noting the state of the
most recent ancestors.

The nematophagous Verticillium species can be grouped
ecologically into species that become attached to the
surface of free-living nematodes, whence they penetrate
the host (e.g. V. baldanoides and V. coccosporum), and
the species that infect nematode cysts and eggs (e.g. V.
lamellicola, V. leptobactrum and V. chlamydosporum)
(Gams, 1988). Evidently, as with the insect pathogens,
nematophagous Verticillium species have evolved inde-
pendently several times to achieve their success as
pathogens. Nevertheless, a phylogenetic branch is
shared by the nematode pathogens that prefer to
penetrate the host and this is distinctive from a common
branch by the isolates that infect nematode cysts and
eggs. This suggests that the patho-ecological groupings
may correlate with two different lineages. Verticillium
species are known to have a broad host range with little
host specificity (Rowe, 1995), and some of the isolates
of nematophagous species we studied here were originally
obtained from insect hosts. The genetic and biochemical
versatility this implies may (1) reflect maintenance of the
ability to exploit resources that temporarily become
available, and (2) indicate that some of the underlying
mechanisms of fungal pathogenesis may be similar in
insects and nematodes. Both the nematophagous and the
entomophagous Verticillium species produce an ordered
sequence of infection structures including appressoria,
and V. chlamydosporium resembles insect pathogens in
secreting subtilisins to breach the hosts' proteinaceous
cuticles (Segers et al., 1994). However, the major
protease components secreted by the nematode patho-
genens were trypsins with a narrow specificity toward
Phe-Val-Arg. Besides proteins, the outer integuments of
insects, nematodes and mushrooms contain chitin as a
major structural component. Consistent with adap-
tation of enzyme production to the specific needs of
different Verticillium spp. on their particular hosts, the
insect, nematode and mushroom pathogens are dis-
tinguishable from the plant pathogens by their ability to
produce chitinases.

The ability to infect nematodes may pre-adapt Vertici-
llum species to infect insects also. However, the converse
is unlikely to be true as, in spite of its abundance, V.
lecanii very rarely attacks nematodes (Gams, 1988). V.
chlamydosporium is most commonly regarded as a
saprophytic soil fungus and, though a potent pathogen
of nematode eggs, may be the least dependent on
nematodes for its development (Gams, 1988). Unlike
the insect and plant pathogens, the nematode pathogens V.
chlamydosporium and V. coccosporum produced pec-
tinases and chitinases. This broad spectrum of poly-
saccharide-hydrolysing enzymes may be indicative of a
less specialized nutritional status than that shown by the
plant and insect pathogens, consistent with the ability to
grow on a greater variety of living and non-living
organic substrates.

ACKNOWLEDGEMENTS

M. J. B. expresses thanks to the Natural Sciences and En-
gineering Research Council of Canada for financial assistance
in the form of an operating grant. A. E. S. would like to thank
NSERC for financial assistance through a PGSB grant. We
thank Richard Humber and Lynne Sigler for providing us with
cultures from the USDA Entomopathogenic Fungus Collection
at Cornell University and the University of Alberta Micro-
fungus Collection and Herbarium, respectively.

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Received 8 September 1998; revised 11 December 1998; accepted 15 December 1998.