Fitness of Races 0 and 1 of *Phytophthora parasitica* var. *nicotianae*

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**ABSTRACT**


Deployment of tobacco (*Nicotiana tabacum*) varieties with complete resistance to race 0 of *Phytophthora parasitica* var. *nicotianae* has led to a rapid increase in the field populations of race 1 in North Carolina. In a field study, population levels of race 1 decreased relative to race 0 when cultivars with partial resistance to both races were planted, suggesting that race 1 isolates were less fit than race 0 isolates. Experiments were conducted to quantify differences in aggressiveness and survivability of the two races. Tobacco varieties with low, moderate, or high levels of partial resistance were inoculated with 60 pathogen isolates, and symptom development was monitored for 3 weeks. Race 0 isolates were more aggressive than race 1 isolates on cultivars with moderate or high levels of partial resistance; incubation periods were shorter and root rot severity was greater with race 0 isolates. Isolates of race 1, however, caused greater stunting of plants with moderate and high levels of partial resistance than race 0 isolates. Field microplots were infested with either a single race or an equal mixture of each race. Soil samples were collected at the end of two growing seasons and again the following spring. Pathogen populations declined from 40 to 80% during winter months, but population declines for race 0 were lower than for race 1 in each treatment over each winter. Race shifts from race 1 to race 0 that were observed in the presence of cultivars with partial resistance appear to be primarily the result of differences in aggressiveness of the races, with a possible minor effect of enhanced overwintering survival of race 0 compared with race 1.

Additional keywords: black shank, oomycete
Sumatra, but concluded that the low level of disease produced was most likely due to the age of the culture. Powers and Lucas (28) reported a weakly pathogenic strain of *P. parasitica* var. *nicotianae* from North Carolina. Apple noted that over 200 isolates of the tobacco black shank pathogen obtained from 44 North Carolina counties, 8 states, and 5 foreign countries varied from weakly pathogenic on a susceptible cultivar to highly pathogenic on a moderately resistant cultivar (1,2). Van Jaarsveld (42) also observed considerable variation in aggressiveness of South African isolates of *P. parasitica* var. *nicotianae* based on lesion length on ‘Hicks’ (susceptible) and ‘TL33’ (moderately resistant). On cultivars with low (‘NC 2326’), moderate (‘Burley 37’), or high (‘Beinhart 1000-1’) levels of partial resistance, however, there was little to no significant difference between lesion lengths of all isolates examined, including isolates of races 0 and 1 (42).

Development of black shank is related to initial inoculum density and level of partial resistance in the host (16,18,34). Ferrin and Mitchell (16) noted that plant mortality was directly related to initial inoculum density for the moderately resistant ‘Speight G-28’, but not for the susceptible ‘Hicks’. Since the differences observed in isolate fitness in a cultivar-rotation study occurred on cultivars with partial resistance (37), initial inoculum densities may have played an important role in disease development and the resulting race shifts observed in those tests. Thus, survivability may be an important component of fitness for races of this pathogen.

Races 0 has long been anecdotally termed “more fit” than race 1 (2,22). However, studies have not specifically examined fitness of races 0 and 1 of the black shank pathogen. The difference in fitness of races 0 and 1 of *P. parasitica* var. *nicotianae* observed in previous field studies could have been attributed to differences in pathogenic fitness (aggressiveness) or ecological fitness (survival), or both. The objectives of this study were to quantify differences in the aggressiveness and survivability of races 0 and 1 of *P. parasitica* var. *nicotianae* and to determine their relative role in the fitness of the two races of the pathogen.

**MATERIALS AND METHODS**

**Aggressiveness study.** Sixty isolates of *P. parasitica* var. *nicotianae* were chosen at random from plots planted in continuous culture to the flue-cured tobacco cultivars ‘K 326’ (low level of partial resistance), ‘K 346’ (high level of partial resistance), or ‘NC 71’ (complete resistance to race 0 and a low level of partial resistance) (37). The isolates included 20 race 0 isolates from ‘K 326’ plots, 20 race 0 isolates from ‘K 346’ plots, and 20 race 1 isolates from ‘NC 71’ plots. All isolates were obtained from soil samples taken from a single field site during one sampling period (37). The purpose of choosing isolates from the same season and same year was to minimize variability and to eliminate the possibility of reduced pathogenicity that may occur due to the age of the culture (26). The isolates were collected in 2001 from a cultivar-rotation study in Edgecombe Co., NC, at the Upper Coastal Plain Research Station using a soil assay method and a semi-selective agar medium. Soil samples from each plot were thoroughly mixed prior to assay. Multiple 1-g subsamples of soil were taken from each plot to obtain the desired 20 isolates. Each subsample was suspended in 25 ml of deionized water and then dispensed, 5 ml per plate, onto five plates of modified PARPH semi-selective medium, containing V8 juice as the basal medium and amended with pentachloronitrobenzene, hymexazol, ampicillin, rifampicin, and pimaricin (19,33). The race of each isolate was confirmed using a set of host differentials (37).

Four cultivars were used to assess variation in aggressiveness of *P. parasitica* var. *nicotianae*: ‘Hicks’ (susceptible), ‘K 326’ (low level of partial resistance), ‘K 149’ (moderate level of partial resistance), and ‘K 346’ (high level of partial resistance). Plants of each cultivar were seeded into a flat containing potting mix (Metro mix 220, The Scotts Company, Marysville, OH) and allowed to grow 3 weeks. After 3 weeks, the plants were transplanted into cell packs (72 cells, 4 × 4 × 5 cm) containing a 1:1:1 (vol/vol/vol) mixture of steam-pasteurized soil, Metro mix (W. R. Grace), and coarse builder’s sand. All plants were watered daily, fertilized as necessary with 20-20-20 water-soluble fertilizer (Peter’s), and allowed to grow an additional 7 to 10 days before a second transplant into a 10.16-cm-diameter pot (11 cm depth, 400 cm³ volume) containing the same 1:1:1 soil mixture used above.

Inoculum was prepared by placing oat grains sterilized by autoclaving at 121°C for 60 min on three consecutive days onto a 3-day-old carrot agar culture of each isolate of *P. parasitica* var. *nicotianae* collected from the soil assay. The oat grains were left on the cultures for 9 days at room temperature (20 to 23°C). Sixty oat grains were then removed from each plate using sterile technique and used as inoculum for the aggressiveness test. Five 5- to 6-week-old plants of each genotype were root-inoculated with each isolate in the greenhouse by placing three colonized oat grains into each 10.16-cm-diameter pot in a triangular pattern around the growing plant. Three approximately 5-cm-deep holes were made in a triangular pattern approximately 3 cm from the stem, and colonized oat grains were placed in each hole with sterilized forceps. Roots were not damaged prior to inoculation. Symptoms development, including chlorosis, wilting, and lesion development, was recorded at 3, 5, 7, 14, and 21 days after inoculation.

Data on incubation period, severity index, stunting, and root rot were collected. Incubation period was determined as the time required for initial symptom development. Incubation period data also were converted into a severity index that corresponded to the number of days required for the expression of symptoms after inoculation (2). The classes and the corresponding severity values were as follows: 3 days = 10, 5 days = 8, 7 days = 6, 14 days = 4, 21 days = 2, and no symptoms at day 21 = 0. In the calculation of the severity index for an isolate, the number of plants placed in each class was multiplied by the corresponding numerical value; these products were totaled and divided by the number of plants in each treatment. Thus, the severity indices reflect both the incidence of disease and how rapidly the disease developed on a given cultivar (2). Initial and final plant heights were measured, and the change in height relative to noninoculated control plants was used as an indication of stunting. After 21 days, all plant root systems were washed and assigned a root rot severity index from 1 to 7, with: 1 = none, 2 = trace, 3 = 1 to 5%, 4 = 6 to 25%, 5 = 26 to 50%, 6 = 51 to 75%, and 7 = 76 to 100%.

The experimental design was a randomized complete block with five replications for each isolate × variety treatment (1,200 plants per trial); plants in each rep were blocked by initial plant height. The experiment was repeated once. The first trial was conducted from June through July 2002, and the second trial was conducted from September through October 2002. Analysis of variance was performed on the severity index, root rot, and stunting data using the PROC GLM procedure of SAS (version 8e, SAS Institute, Cary, NC), and means separation was conducted using the Waller-Duncan k-ratio test (*k* = 100). Unless otherwise indicated, only significant (*P* ≤ 0.05) differences between treatment means are presented.

**Survival study.** A 3-year microplot experiment was established in May 2000 at the Central Crops Research Station in Clayton, NC, and continued through the spring of 2003. Plots (75 cm diameter, 45 cm depth) were infested with either race 0 or race 1 (single race) or with an equal mixture of both races (mixed race). Plots were infested by uniformly distributing pathogen-colonized oat grains (~ 50 cm² per plot) into each plot in late May 2000. Two isolates of *P. parasitica* var. *nicotianae* were used to infest all microplots: a race 0 isolate from Edgecombe Co., NC, and a race 1 isolate from Mitchell Co., NC. Each isolate was inoculated onto test plants prior to infestation to ensure that each isolate was highly aggressive on cultivars with various levels of partial resis-
tance. Each of the single and mixed race plots received one of three treatments: ‘K326’ (low level of partial resistance), ‘K346’ (high level of partial resistance), or ‘NC71’ (complete resistance to race 0 and a low level of partial resistance) that were continuously cropped over the three growing seasons. Each race × cultivar treatment was replicated 16 times (single race) or 18 times (mixed race), and treatments were arranged in a completely randomized experimental design. Each year plants of each cultivar were transplanted into their respective plots in late May; soil samples were collected in September prior to the removal of plants from each plot, and soil samples were collected again from each plot the following spring (April). Recommended practices for tobacco production were followed, except that the plants were not topped (flower heads removed), and no chemicals were applied to prevent growth of axillary buds at leaf nodes.

Inoculum density of *P. parasitica* var. *nicotianae* was determined for all plots in the fall and spring. In addition, isolates obtained from the mixed race plots were screened for race to determine the percentage of each race in the population. Soil samples were collected from each microplot with a 3-cm-diameter soil probe in late fall and in the following spring and compared using different soil probes for each race treatment to minimize cross-contamination. On average, soil samples were taken to a depth of 15 to 18 cm. Four soil cores were collected from each plot and bulked prior to transport to the lab in coolers. Coolers were stored at room temperature (21 to 23°C) for no more than 4 days before the soil was assayed. Soil samples were thoroughly mixed prior to assay, and inoculum density was determined by placing the soil suspension on PARPH medium as described above for isolate collection. Three subsamples of each bulked soil sample were assayed per sample. After 48 h, the soil suspension was washed from each plate. The plates were then allowed to incubate at room temperature for 24 additional hours, after which *P. parasitica* var. *nicotianae* colonies were enumerated and transferred. The number of propagules of the pathogen per gram of soil was determined by averaging the number of colonies in each of the three 1-g samples after correcting for soil moisture. For race determination in the mixed race plots, a minimum of 10 colonies from each plot, for a total of 180 colonies per treatment, were transferred to fresh PARPH, and additional transfers were made to obtain pure cultures of *P. parasitica* var. *nicotianae*. If 10 propagules per gram were not recovered from the initial soil assay, the soil was assayed additional times until 10 propagules were recovered from each plot, but population data were taken only from the initial assay. All isolates also were plated on nutrient agar to test for the presence of bacteria prior to transfer to 5% clarified carrot agar. Carrot juice (Hollywood or The Hain Celestial Group Inc., Melville, NY) was clarified by filtering the juice through diatomaceous earth (Cellite 545, Fisher Scientific, Fair Lawn, NJ). Fifty milliliters of the filtrate were added to 950 ml of deionized water and 20 g of Bacto Agar (Difco, Detroit, MI) and then sterilized in an autoclave prior to pouring.

To determine the race of each isolate, a host differential was used that included four cultivars or breeding lines: ‘Hicks’ (susceptible), ‘K326’ (low level of partial resistance), ‘NC71’ (complete resistance to race 0 and race 1), ‘K346’ (high level of partial resistance), and ‘Ky 14 × L8’ (*N. longiflora* gene) (37). Plants of each differential were seeded into a flat containing potting mix (Metro mix 220) and allowed to grow 3 weeks. After 3 weeks, the plants were transplanted into cell packs (72 cells, 4 × 4 × 5 cm) containing a 1:1:1 (vol/vol/vol) mixture of steam-pasteurized soil, Metro mix (W. R. Grace), and coarse builder’s sand. All plants were watered daily, fertilized as necessary with 20-20-20 water soluble fertilizer (Peter’s), and allowed to grow an additional 7 to 10 days before inoculation.

Inoculum was prepared as described above and 24 oat grains were removed from each plate using sterile technique and used as inoculum for race determination. Three 4-week-old plants of each genotype were root inoculated with each isolate in the greenhouse by placing two colonized oat grains into each 4 × 4 × 5 cm cell. Roots were not damaged prior to inoculation.

### Table 1. Incubation period for aboveground symptoms of black shank on four tobacco cultivars that vary in level of partial resistance when inoculated with groups of isolates of either race 0 or 1 of *Phytophthora parasitica* var. *nicotianae*

<table>
<thead>
<tr>
<th>Isolate group</th>
<th>Cultivar*</th>
<th>Hicks</th>
<th>K326</th>
<th>K149</th>
<th>K346</th>
</tr>
</thead>
<tbody>
<tr>
<td>Race 0 (K346)</td>
<td>7 b&lt;sup&gt;a&lt;/sup&gt;</td>
<td>14 c</td>
<td>16 b</td>
<td>18 b</td>
<td></td>
</tr>
<tr>
<td>Race 0 (K326)</td>
<td>12 a</td>
<td>16 b</td>
<td>18 a</td>
<td>21 a</td>
<td></td>
</tr>
<tr>
<td>Race 1 (NC71)</td>
<td>13 a</td>
<td>17 a</td>
<td>19 a</td>
<td>&gt;21</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Five-week-old plants were inoculated with colonized oat grains and observed over a 21-day period following inoculation, and the time required for expression of first visible symptoms was recorded. Data represent the mean of 20 isolates per race category.

<sup>b</sup> Hicks, susceptible; K326, low level of partial resistance; K149, moderate level of partial resistance; and K346, high level of partial resistance.

<sup>c</sup> Isolates were collected using a soil assay from field plots planted to cultivars K346 (high level of partial resistance), K326 (low level of partial resistance), or NC71 (completely resistant to race 0 and race 1). Isolates were further characterized as race 0 or race 1 using a host differential.

### Table 2. Severity index on four tobacco cultivars that range from susceptible to highly resistant when inoculated with isolates of race 0 or 1 of *Phytophthora parasitica* var. *nicotianae*

<table>
<thead>
<tr>
<th>Isolate group</th>
<th>Cultivar*</th>
<th>Trial 1</th>
<th>Trial 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hicks&lt;sup&gt;a&lt;/sup&gt;</td>
<td>K326</td>
<td>K149</td>
</tr>
<tr>
<td>Race 0 (K346)</td>
<td>6.68 a</td>
<td>3.92 a</td>
<td>3.50 a</td>
</tr>
<tr>
<td>Race 0 (K326)</td>
<td>4.78 b</td>
<td>3.48 b</td>
<td>2.88 b</td>
</tr>
<tr>
<td>Race 1 (NC71)</td>
<td>4.42 b</td>
<td>3.14 c</td>
<td>2.66 b</td>
</tr>
</tbody>
</table>

<sup>a</sup> Hicks, susceptible; K326, low level of partial resistance; K149, moderate level of partial resistance; and K346, high level of partial resistance.

<sup>b</sup> Isolates were collected using a soil assay from field plots planted to cultivars K346 (high level of partial resistance), K326 (low level of partial resistance), or NC71 (completely resistant to race 0 and a low level of partial resistance). Isolates were further characterized as race 0 or race 1 using a host differential.

<sup>c</sup> Five-to-six-week-old plants of each cultivar were inoculated with colonized oat grains of a given isolate and observed over a 21-day period. Time required for the expression of first visible symptoms was recorded. Time to symptom expression was then converted to a disease index on a 0 to 10 scale, which represented both the time to symptoms and severity of symptoms. The scale is as follows: 3 days = 10, 5 days = 8, 7 days = 6, 14 days = 4, 21 days = 2, no symptoms at day 21 = 0. Data represent the mean of 20 isolates per race category and are averages over two trials.

<sup>d</sup> Severity values in each column followed by the same letter do not differ significantly (Waller-Duncan k = 100).
After 14 days, plants were scored for the presence or absence of symptoms, and the isolate was determined to be either race 1 or race 0. Race 1 was defined by the ability to cause disease on 'NC 1071' and 'Ky 14 × L8'. Race 0 did not cause disease on either of these hosts, but caused disease on the other two, 'Hicks' and 'K 326'. Race 1 also caused disease on 'Hicks' and 'K 326'.

Percent survival of the pathogen was calculated by dividing the inoculum density (ID) in the spring sample by the ID in the fall sample and multiplying by 100. For the mixed race plots, race determinations were used to determine what percentage of the total pathogen population recovered could be attributed to each race. Analysis of variance was performed on the percent survival data using the PROC GLM procedure of SAS (version 8e), and means separation was conducted using the Waller-Duncan k-ratio test ($k = 100$). Unless otherwise indicated, only significant ($P \leq 0.05$) differences between treatment means are presented.

**RESULTS**

Aggressiveness study. Significant variations were observed in aggressiveness between the groups of isolates representing races 0 and 1. These differences occurred in all fitness parameters measured, including incubation period, disease severity index, root rot, and stunting. Significant variation also was observed between the two groups of race 0 isolates for these same measures of fitness.

As the level of partial resistance in the four tobacco cultivars increased, the incubation period increased for all isolate groups (Table 1). Differences were observed between groups in incubation period, but there was no cultivar by isolate-group interaction ($P > 0.4$); race 0 isolates from 'K 346' caused symptoms faster than race 0 isolates from 'K 326' and race 1 isolates on each cultivar (Table 1). For example, on the susceptible 'Hicks', race 0 isolates from 'K 346' had a severity index of 6.68, race 0 isolates from 'K 326' had an index of 4.78, and race 1 isolates had index value of 4.42. Again, ranking of isolate groups remained constant across the four levels of partial resistance. Differences were also observed within race 0 isolates, where those collected from 'K 346' had higher severity indices than race 0 isolates from 'K 326'. Race 1 isolates did not cause symptoms on 'K 346' and thus had severity indices of 0.00 in both trials.

Distribution of severity indices also varied within each isolate group (Fig. 1A to C). All isolates, except for race 1 isolates on 'K 346', had a severity index of 2 or greater on all tobacco cultivars (Fig. 1A).

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**Fig. 1.** Percentage of isolates in each isolate group with severity indices (0 to 10 scale) of A, 2, B, 4, or C, 6 on tobacco cultivars Hicks (susceptible), K326 (low level of partial resistance), K149 (moderate level of partial resistance), and K346 (high level of partial resistance). A severity index was based on number of days to symptom development: 2 = 15 to 21 days, 4 = 8 to 14 days, and 6 = 1 to 7 days.
However, only race 0 isolates from ‘K 346’ had severity index values of 4 or greater on all four tobacco cultivars. Less than 10% of race 0 isolates from ‘K 326’ and race 1 isolates had an index of 4 on the moderately resistant ‘K 149’ (Fig. 1B). On the susceptible ‘Hicks’, an index of 6 was observed for 65, 10, and 5% of race 0 isolates from ‘K 346’, race 0 isolates from ‘K 326’ and race 1 isolates, respectively, (Fig. 1C).

There was not a significant effect of trial on root rot severity, so data were combined from the two runs of the experiment. There was a significant cultivar × race interaction \( (P < 0.001) \), so data are shown for each isolate group by cultivar. There was little or no difference among isolate groups in root rot severity on the susceptible cultivar or the cultivar with the low level of partial resistance (Fig. 2A and B). However, race 0 isolates had significantly more root rot than race 1 isolates on cultivars with moderate and high levels of partial resistance (Fig. 2C and D).

There was a significant effect of trial on stunting; plants grew less overall and had slightly less stunting in trial two. However, trends were similar across trials, so only data from the first trial are presented (Fig. 3). There was no significant difference in stunting on the susceptible ‘Hicks’, or on ‘K 326’ with low partial resistance (Fig. 3A and B). In sharp contrast to observations on root rot, race 1 isolates caused more stunting than either group of race 0 isolates on the moderately resistant ‘K 149’ and highly resistant ‘K 346’ (Fig. 3C and D). Race 0 isolates from ‘K 346’ caused more stunting than did race 0 isolates from ‘K 326’. Although most cultivars were stunted 60 to 80% relative to control plants, stunting of ‘K326’ was only about 30% with all isolate groups (Fig. 3B).

**Survival study.** Race 0 populations exceeded race 1 populations at every time point in the mixed-race plots from the microplot experiment. This was true for the plots planted with the cultivars with low or high partial resistance (Fig. 4A). In sharp contrast, on the completely resistant ‘NC 71’, race 1 populations exceeded race 0 populations at every time point (Fig. 4B). Race 0 populations declined steadily in plots planted with ‘NC 71’ and in the spring of 2003 was not detected (Fig. 4B).

There was not a significant effect of trial on survival; plants grew less overall and had slightly less survival in trial two. However, trends were similar across trials, so only data from the first trial are presented (Fig. 3). There was no significant difference in survival on the susceptible ‘Hicks’, or on ‘K 326’ with low partial resistance (Fig. 3A and B). In sharp contrast to observations on root rot, race 1 isolates caused more survival than either group of race 0 isolates on the moderately resistant ‘K 149’ and highly resistant ‘K 346’ (Fig. 3C and D). Race 0 isolates from ‘K 346’ caused more survival than did race 0 isolates from ‘K 326’. Although most cultivars were stunted 60 to 80% relative to control plants, stunting of ‘K326’ was only about 30% with all isolate groups (Fig. 3B).

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Populations of both races declined sharply over each winter (Fig. 5A and B). Differences in survival were observed for the races. For example, in the mixed race plots, where survival data could be combined over the 2 years, race 0 isolates survived significantly better than race 1 isolates (Fig. 5B). On average, 60% of the race 0 population survived the winter, while only 42% of the race 1 population survived. In the single race plots, where there was a significant year and year × race interaction, data could not be combined over years, and differences in survival followed the same trend but were not significantly different (Fig. 5A).

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**Fig. 2.** Root rot caused by *Phytophthora parasitica* var. *nicotianae* on four tobacco cultivars with different levels of partial resistance: A, susceptible, B, low level, C, moderate level, and D, high level of resistance to black shank of tobacco. Bars are the average of five replications per treatment. Bars with the same letter do not differ significantly (Waller-Duncan \( k = 100 \)).
DISCUSSION

Isolates of races 0 and 1 of *P. parasitica* var. *nicotianae* exhibited significant differences in aggressiveness on *N. tabacum* cultivars with different levels of partial resistance. Differences in incubation period, disease severity index, and root rot severity were observed. Variations in aggressiveness among isolates of race 0 of *P. parasitica* var. *nicotianae* have been reported (2,22,42), but this is the first report where variations in aggressiveness between races of the tobacco black shank pathogen have been quantified. Similar observations were reported for races 1 and 2 of *Verticillium dahliae*, where isolates of race 1 caused greater losses than race 2 on cultivars susceptible to both races. For both pathogens, the ability to overcome resistance genes in the host was associated with a measurable loss in aggressiveness on susceptible cultivars (6,41).

The time from inoculation to initial aboveground symptom development increased as the level of resistance in the tobacco cultivars increased for all isolate groups examined. This result indicated that the cultivars used in this study behaved in the greenhouse as they have previously reacted in fields infested with *P. parasitica* var. *nicotianae*. The resistance level (low to high) is generally assigned by extension specialists based on disease development in field trials over many locations and years. The ranking of isolate groups was similar across all cultivars, indicating that the isolates varied only in aggressiveness and not in virulence to the tobacco cultivars (41).

Differences between groups of race 0 isolates collected from ‘K 326’ or ‘K 346’ plots were observed in this study. Dukes and Apple (13) demonstrated that aggressiveness of isolates increased after passage through a resistant cultivar compared to passage through a susceptible cultivar. Their study suggested that under field conditions the pathogen strains would become more aggressive as a result of continuous planting of resistant cultivars. ‘K 346’ has a high level of partial resistance compared with ‘K 326’, which has a low level of partial resistance. The 20 isolates that comprised an isolate group in this study were collected from plots during the fourth year of a cultivar-rotation study, where either ‘K 346’ or ‘K 326’ was continuously cropped. Since ‘K 346’ has a higher level of resistance than ‘K 326’, the higher level of aggressiveness observed in this isolate group supports the results of Dukes and Apple (13). The components of aggressiveness described by Dukes and Apple (14) were not quantified for the isolates in this study.

Race 1 caused significantly more stunting than either group of race 0 isolates, even though the race 0 isolates were more
aggressive in all other measures of disease development. Similar results were obtained by Carlson et al. (8), where a race 0 isolate of *P. parasitica* var. *nicotianae* caused symptoms faster, and disease severity was greater, than with a race 1 isolate. The race 1 isolate, however, caused significant plant stunting, reduced quality, and weakened root systems (8). The mechanisms of pathogenicity have not been investigated for race 0 and race 1 isolates, and deserve further study.

The cultivar with the low level of partial resistance, ‘K 326’, was only stunted 30% on average, while most other cultivars were stunted 50 to 80%, with the exception of ‘K 346’ inoculated with race 0 from ‘K 326’. ‘K 149’ and ‘K 346’ have moderate and high levels of partial resistance, respectively. Jones and Shew (18) showed that ‘Hicks’ and ‘K 326’ had significantly larger root systems (two to three times larger) than moderately resistant or highly resistant cultivars. Production of fewer roots by black shank resistant cultivars may be important in their survival in soil infected with *P. parasitica* var. *nicotianae* and a possible avoidance mechanism (35). Plants that produce few roots are less likely to come in contact with pathogen propagules than those that have a greater rooting intensity (35). Conversely, cultivars with a greater rooting intensity may be more able to tolerate drought stress or pathogen infection by being able to uptake water and nutrients even with infected roots. Perhaps ‘K326’ was more able to overcome the loss of part of its root system compared to the effects on the more resistant cultivars.

Survival studies examined the population dynamics of races 0 and 1 of *P. parasitica* var. *nicotianae* in field soil where either partial or complete resistance was deployed in host cultivars. Even though partial resistance was effective against both races 0 and 1 of *P. parasitica* var. *nicotianae*, race 0 had a higher inoculum density than did race 1. This result also indicated that race 1 was not as fit as race 0 on cultivars with partial resistance. By simply examining the population dynamic curves, however, it was not possible to ascertain if the difference was due to differences in initial inoculum density (survival) as Ferrin and Mitchell (16) suggested as being important or if it was due to differences in aggressiveness.

On the cultivar with complete resistance to race 0, race 1, by far, had the highest inoculum density at all sampling dates. Race 1 is able to reproduce on ‘NC 71’, and thus the curve takes into account survival, reproduction, and pathogenic potential. Race 0, in sharp contrast, cannot reproduce on ‘NC 71’, and thus this curve represents a survival curve in the absence of new reproduction. The increase in race 1 relative to race 0 on the completely resistant cultivar can be explained by examination of the mechanism of resistance. Complete resistance does not prevent infection, but it prevents growth and reproduction of the pathogen, which results in a sharp decline in the pathogen population (35). A similar phenomenon has been observed with *Thielaviopsis basicola* on tobacco cultivars with the *N. debneyi* gene for complete resistance (36). Reproduction of *T. basicola* was very low or nonexistent on cultivars with resistance from *N. debneyi*. *P. parasitica* var. *nicotianae* can survive in soil for 5 years or longer in the absence of tobacco (4,22). Our results indicate that propagules persisted for at least 3 years, because the plots were established in May 2000 and the final samples were collected in the spring of 2003.

The increase in predominance of race 1 of *P. parasitica* var. *nicotianae* in the flue-cured region of North Carolina has corresponded with the widespread deployment of cultivars with the *Ph* gene, which confers complete resistance to race 0 of the pathogen. Race 0 is not able to reproduce on cultivars with the *Ph* gene, and thus race 1 has a selective advantage that allows it to become the predominant member of the population during continuous use of complete resistance. Our results demonstrate that isolates of race 1 are less fit than isolates of race 0.

Simply defined, fitness is the combined ability of an organism to survive and reproduce. Vanderplank (41) considered pathogen fitness to be the driving force in the evolution and stability of a pathosystem in agriculture. He argued that in a freely mutating system, unnecessary virulence genes impose a fitness penalty to the pathogen. He suggested that a mutation from avirulence to virulence would only occur if it was necessary to overcome an R gene, i.e., if it was the only way the patho-

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**Fig. 4.** Population dynamics of *Phytophthora parasitica* var. *nicotianae* in field microplots planted with tobacco cultivars A, K 326 (low partial resistance) or K 346 (high partial resistance), or B, NC71 (complete resistance) in mixed-race plots. Plots were originally infested with equal levels of races 0 and 1.
Although shifts in relative population of resistance to the developing race 0 population for the following year against the developing race 0 population.

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