The vertical distribution of Phytophthora sojae was investigated in soil samples collected in the spring of 1994 from soybean fields at 62 locations in Illinois, Indiana, Iowa, and Minnesota. In the fall of 1995, soil samples were collected from 18 additional locations in Illinois and Iowa. Each location consisted of a pair of no-till and conventional-till fields, and soil samples were collected from arbitrarily selected locations in each field at 0- to 7.5-cm and 7.5- to 15-cm depths. Separate intensive sampling was made in the spring of 1995 from two pairs of adjacent no-till and conventional-till fields at the Iowa State University Northeast Research Farm, in which samples were collected from 0- to 30-cm depth in increments of 5 cm. Samples were assayed for P. sojae with the use of a leaf-disk bioassay. In the 1994 regional samples, there was greater recovery of P. sojae (P ≤ 0.05) at 0- to 7.5-cm depth in the no-till samples than in the conventional-till samples for all states except Minnesota. The fall 1995 samples from Illinois followed a similar trend (P = 0.05); whereas samples from Iowa showed no significant difference between tillage systems. At depths greater than 7.5 cm, there was generally no difference in detection frequency of P. sojae between tillage systems. Samples from the Northeast Research Farm followed patterns of vertical distribution similar to those of the regional samples. In no-till fields, the detection frequency of P. sojae was greatest near the soil surface; two to three times greater than that of the conventional-till fields at this depth. In the conventional-till fields, however, the frequency of recovery peaked at 20 cm and was comparable at these depths to those of no-till fields. There was a positive correlation between the percentage of leaf disks colonized and residue dry weights in the no-till fields (r = 0.84, P = 0.04; and r = 0.86, P = 0.03) but not in the conventional-till fields (r = -0.06, P = 0.90; and r = -0.60, P = 0.17). The recovery of P. sojae in greater frequency near the soil surface in no-till fields than in conventional-till fields suggests that the potential for damping-off may be greater in no-till fields than in conventional-till fields.

Additional keywords: conservation tillage, minimum-till, reduced-till, surface residue, Phytophthora root and stem rot

Phytophthora root and stem rot of soybean, caused by Phytophthora sojae, is a major disease of soybean worldwide (23). In the United States, the disease is widely distributed in most soybean-producing regions and is estimated to cause an annual yield loss of up to half a million metric tons (34). Reduction in yield may result from stand losses that arise from pre- and postemergence damping-off or from wilting or stunting of infected older plants. Even though infection can occur at any stage during plant development (24), most of the damage is believed to result from infection at the seedling stage.

As with other diseases caused by Phytophthora species (8), environmental conditions, such as soil compaction, that prolong the saturation of the soil favor development of the disease (10). There is little information on the effect of tillage practices on Phytophthora root and stem rot of soybean. Tachibana (30) observed more severe root rot in conservation tillage (ridge-till) treatments than in conventional tillage treatments. Schmitthenner and Van Doran (25) and Dick and Van Doran (7) reported greater yield reduction of a susceptible cultivar in Phytophthora-infested no-till plots than in conventional-till plots, and they assumed this effect was caused by the disease.

Tillage has been a major component of agricultural practices, with the primary objectives of weed control and seed bed preparation. Generally, conventional tillage practices encompass burial of surface residues and loosening of crust surfaces and dense soil layers. However, in the last few years, there has been increased emphasis on maintenance of surface crop residues, primarily to prevent wind and water erosion (2). Consequently, there is a shift from the traditional clean-till seed bed preparation to adoption of conservation tillage practices that maintain greater than 30% residue cover after planting. Such tillage practices involve little (reduced-till) or no (no-till) soil disturbances between harvest and planting. Within the last decade, no-till acreage in the north central United States has increased from 5.4 to 29.6% of the total soybean production area (2).

The role of crop residues in survival of plant pathogens has long been understood (5,6,16,28). For many plant pathogens, crop residues provide shelter and a food base for reproduction. Crop residues left on the soil surface decompose more slowly than buried residues (9), which may lead to concentration of inhabiting pathogens near the soil surface (20). P. sojae is a homothallic fungus and, therefore, produces abundant oospores in infected root and stem tissues (24,26). Even though oospores may survive in soil (32), infested soybean root and stem residues are believed to be the primary sources of inoculum (24). Therefore, the vertical distribution of the pathogen may be directly related to the vertical distribution of soybean residues, and this in turn may have significant impact on seedling disease caused by the pathogen. The impact may be difficult to demonstrate in the field because of the uncertainty of favorable environmental conditions and the existence of many races of the fungus (1,35). However, one would expect that high population densities of the pathogen near the soil surface may lead to greater risk of damping-off than similar densities in the soil profile. Schmitthenner and Van Doran (25) discussed the possible occurrence of such scenarios and emphasized the need for further investigation. Currently, information is lacking on how the vertical distribution of the pathogen is affected by tillage practices. Such information would be useful in assessing risks associated with particular tillage practices, especially pertaining to Phytophthora root and stem rot of soybean. In addition, this information may be beneficial for making decisions on soil sampling schemes to determine population densities of the pathogen in different management systems. The primary objective of this study was to determine the vertical distribution of propagules of P. sojae in no-till and conventional-till fields.

MATERIALS AND METHODS

In 1994 and 1995, paired soil samples were collected from four states of the north central United States. In addition to these regional samples, separate soil samples were collected in the spring of 1995 from two pairs of no-till and conventional-till
fields at the Iowa State University Northeast Research and Demonstration Farm at Nashua. The regional samples were collected from two soil depths, and the Nashua samples were collected from six depths.

**Regional samples.** In June 1994, soil samples were collected from 62 locations (Fig. 1), each with adjacent conventional-till and no-till soybean fields. Fields were located in Illinois, Indiana, Iowa, and Minnesota and represented 12, 10, 19, and 10 counties, respectively. Samples were collected again in October 1995 from 10 locations in Illinois and 8 locations in Iowa. The 1995 locations in both states were different from those of 1994.

The 1994 locations were selected in cooperation with either the cooperative extension county agents or the assistance of the farmers. In areas where contacts with either county extension agents or farmers were not possible, fields were identified by the presence or absence of the typically erect remains of cornstalks that characterize no-till fields, coupled with the presence or the absence of signs of tillage operations. All 1995 locations were selected with the help of farmers or county extension agents. At each location, a no-till field was first identified and paired with the nearest conventional-till field of similar size. The no-till and the conventional-till fields at each location were either contiguous or, at most, within 1.6 km of each other. All soybean fields had been cropped with corn the previous year except one field in Minnesota that was cropped with alfalfa. From each field, 50 arbitrarily located soil cores at depths of 0 to 15 cm were collected with a 2-cm-diameter soil probe. The soil probe was marked 7.5 and 15 cm from the tip, and soil cores were separated into shallow (0 to 7.5 cm) and deep (7.5 to 15 cm) components. The samples were transported to the laboratory in sealed boxes on ice and stored at 4°C until use. The samples were taken out of cold storage and maintained at 22 to 24°C for 15 days prior to bioassay.

**Nashua Research Farm samples.** Samples were collected from two sites (designated hereafter as sites 1 and 2), each comprising a pair of contiguous no-till and conventional-till fields that had 17 years of the same tillage history. All fields had a prior history of Phytophthora root and stem rot and received similar treatments in all aspects of management except tillage operations. The conventional-till fields were plowed with a moldboard plow every fall after harvest and cultivated every spring before planting. The no-till fields did not receive any tillage operations except the opening of seed furrows with the planter once every spring. The fields at each site had been rotated to corn and soybean for as long as their tillage history. The year prior to sample collection, the fields at site 1 were cropped with soybean, while the fields at site 2 were cropped with corn. The fields at site 1 were nearly flat; whereas the fields at site 2 were located on a gentle slope of 0 to 2%. Each field was slightly less than 0.5 ha.

**Bioassay of regional samples.** Soil samples were forced through a 6-mm screen to break large-sized soil clods and to establish a uniform aggregate size across all samples. Soybean residues that were too large to pass through the screen were returned to the screened soil. For each regional sample, population densities of *P. sojae* were determined from two subsamples using a leaf-disk bioassay (18, 33) in the greenhouse at 23 to 27°C. Sixty cm³ of each subsample were spread uniformly on top of a 9-cm-deep column of pasteurized sandy clay loam (field soil with 52.5, 25,

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#### Table 1. Prevalence of *Phytophthora sojae* in selected locations of adjacent no-till (NT) and conventional-till (CT) fields sampled in spring 1994 and fall 1995 in four states of the north central region

<table>
<thead>
<tr>
<th>Year</th>
<th>State</th>
<th>Locations (no.)</th>
<th>NT</th>
<th>CT</th>
<th>State total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1994</td>
<td>Illinois</td>
<td>15</td>
<td>40.0</td>
<td>13.3</td>
<td>26.7</td>
</tr>
<tr>
<td></td>
<td>Indiana</td>
<td>16</td>
<td>62.5</td>
<td>43.8</td>
<td>53.1</td>
</tr>
<tr>
<td></td>
<td>Iowa</td>
<td>20</td>
<td>85.0</td>
<td>65.0</td>
<td>75.0</td>
</tr>
<tr>
<td></td>
<td>Minnesota</td>
<td>11</td>
<td>72.7</td>
<td>81.8</td>
<td>77.3</td>
</tr>
<tr>
<td>1995</td>
<td>Illinois</td>
<td>10</td>
<td>60.0</td>
<td>40.0</td>
<td>50.0</td>
</tr>
<tr>
<td></td>
<td>Iowa</td>
<td>8</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
</tr>
</tbody>
</table>

* Each location consisted of a no-till (NT) and a conventional-till (CT) field.

The values are means of the corresponding locations. A field was considered positive if *P. sojae* was detected in samples from either of the two depths (0 to 7.5 cm and 7.5 to 15 cm) or both.

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![Fig. 1. Locations in four states of the north central region from which soils were sampled from paired no-till and conventional-till fields during spring 1994 and fall 1995.](image-url)
and 22.5% sand, silt, and clay, respectively) contained in 475-ml perforated plastic cups. The soil columns were watered to saturation and left to drain for 96 h. The samples then were flooded with distilled-deionized water, and ten 0.8-cm-diameter soybean leaf disks were floated on the surface of the water pooled above the soil surface. The leaf disks were obtained from young but fully unrolled leaves of the cultivar Sloan (susceptible to all known races). After 24 h, the leaf disks were removed and surface-sterilized with 0.05% NaOCl for 30 s, then plated onto a selective medium (5 disks per 9-cm-diameter plate) containing hymexazol at 40 mg/liter (14). The plates were incubated in dark at 22 to 24°C. The percentage of leaf disks colonized by *P. sojae* was determined after 4 days. Typical cultures of the isolates that were obtained in this process were shown to be pathogenic to susceptible soybean cultivars (35).

**Bioassay of Nashua samples.** Population densities of *P. sojae* in the Nashua samples were assessed as described above except that four subsamples and 20 leaf disks per subsample were used in the bioassay. Additionally, part of each soil sample from the Nashua fields was used for measurement of residues. Residues were extracted from 500 cm$^3$ (water displacement method) of each sample with a semi-automatic soil elutriator (4) on 0.6-mm-diameter screens. Residues left on the screens were collected, air-dried on greenhouse benches, and weighed.

**Data analyses.** The incidence of *P. sojae* in each of the two fields at each location was determined from the presence or absence of colonization of the leaf disks by the fungus at any of the two sample depths. The significance of the difference in mean incidence between the tillage systems for individual states then was determined with use of McNemar's test (27), a nonparametric equivalent of the paired *t* test.

At some of the locations, *P. sojae* was not detected in either of the two fields. Only data from locations in which *P. sojae* was detected in at least one of the two fields were included in the analyses of the recovery of the pathogen. For each location, data from 0 to 7.5 cm of the no-till field were compared with data from 0 to 7.5 cm of the conventional-till field. Similarly, data from 7.5 to 15 cm of the respective tillage systems at each location were compared. To avoid division by zero, 0.01 was added to each value, and the differences in the percentages of leaf disks colonized between the no-till (NT) and the conventional-till (CT) fields for each location at each depth were standardized to the conventional-till value as: % difference = (NT – CT)/CT. The differences for all locations in each state then were analyzed to determine whether the mean difference was significantly different from zero with the use of *t* test.

For the Nashua samples, the recovery *P. sojae* at each depth of the no-till fields was paired with that of the conventional-till fields at respective depth and analyzed with the use of *t* test. The relationship between the recovery of *P. sojae* and residue dry weight at each depth was determined with the use of Pearson's correlation. All statistical analyses were conducted with SAS software (SAS Institute, Cary, NC).

**RESULTS**

**Regional samples.** In the spring 1994 samples, *P. sojae* was detected in 77.4% of the total locations comprising 58.1% of the fields sampled across the four states. Fields in Minnesota and Iowa had the greatest prevalence of *P. sojae* (77.3 and 75.0%, respectively), followed by Indiana (53.1%) and Illinois (26.7%) (Table 1). Of the fields in which *P. sojae* was detected, 33.1% and 25% were no-till and conventional-till fields, respectively. Overall, the prevalence of *P. sojae* was greater in no-till than in conventional-till fields, except in Minnesota. However, McNemar's test showed no significant difference (*P* values ranging from 0.16 to 0.56) among the matched

![Fig. 2. Recovery of Phytophthora sojae (as determined by a leaf-disk bioassay) in no-till and conventional-till fields at depths of 0 to 7.5 cm (A) and 7.5 to 15 cm (B) in samples collected from Illinois, Indiana, Iowa, and Minnesota in spring 1994. Bars with the same letter within a state are not significantly different according to paired *t* test (*P* ≤ 0.05).]
pairs of no-till and conventional-till fields for the individual states (Table 1).

The recovery of *P. sojae*, expressed as the percentage of leaf disks colonized, at 0 to 7.5 cm depth were greater in no-till samples than in conventional-till samples for all states (Fig. 2A). The paired *t* test analysis showed that the difference was significant for samples from Illinois, Indiana, and Iowa (*P* = 0.02, *P* = 0.01, *P* = 0.01, respectively), but not from Minnesota (*P* = 0.19). Samples from the 7.5- to 15-cm depth showed a similar trend in recovery, with samples from the no-till fields having greater recovery than those from the conventional-till fields. However, the difference was significant only in samples from Iowa (*P* = 0.05, Fig. 2B).

Samples collected from Illinois and Iowa during the fall of 1995 had greater prevalence of *P. sojae* than those collected in the spring of 1994 (Table 1). The difference in incidence between no-till fields and conventional-till fields was not significant. In samples from Illinois at the 0- to 7.5-cm depth, recovery of *P. sojae* was significantly greater in no-till fields than in conventional-till fields (*P* = 0.05, Fig. 3A). However, there was no significant difference in *P. sojae* recovery at this depth between the two tillage systems in samples from Iowa. Samples from both states showed no significant difference between the two tillage systems at the 7.5- to 15-cm depth (Fig. 3B).

*Nashua samples.*** *P. sojae* was recovered from the upper 10 cm of the soil from the no-till fields at both sites more frequently than from their respective adjacent conventional-till fields (Figs. 4A and 4B). Recovery of *P. sojae* was greatest near the soil surface in the no-till fields and declined with depth in the first 10 to 15 cm of the soil. Within this depth range, recovery of the pathogen was two to three times greater in the no-till fields than in the conventional-till fields. Recovery of the pathogen in the conventional-till fields peaked at 20 cm, where it was comparable to that of the no-till fields. Overall, there was greater recovery of *P. sojae* in the no-till fields than in the conventional-till fields at both sites (*P* = 0.05 and *P* = 0.06 for sites 1 and 2, respectively).

In the no-till fields, the vertical distribution of the pathogen followed the vertical distribution of residues (Fig. 5). There was a significant correlation between the percentage of leaf disks colonized and residue dry weight in no-till fields at both sites (r = 0.84, *P* = 0.04, and r = 0.86, *P* = 0.03, for sites 1 and 2, respectively). However, in the conventional-till fields, the relationship was less clear than in the no-till fields (r = −0.06, *P* = 0.90, and r = −0.64, *P* = 0.17 for sites 1 and 2, respectively).

**DISCUSSION**

This study demonstrated that at soil depths close to the surface, there was greater recovery of *P. sojae* in no-till fields than in conventional-till fields. This trend was generally evident in most of the regional samples and in all of the samples from the Nashua Research Farm in Iowa.

The presence of *P. sojae* in greater abundance near the soil surface in no-till fields than in conventional-till fields may be attributed partially to the absence of vertical soil inversion in no-till fields. *P. sojae* is primarily a root-infecting, soil-borne pathogen. However, lesions from root infection can extend on the stem up to the tenth node (23), and abundant oospores are produced in infected stems (24). In no-till fields, such stems remain on or near the soil surface after harvest as part of the residue. When the infested stem residues decompose, oospores may be released to the immediate vicinity (near the soil sur-

**Fig. 3.** Recovery of *Phytophthora sojae* (as determined by a leaf-disk bioassay) in no-till and conventional-till fields at depths of 0 to 7.5 cm (A) and 7.5 to 15 cm (B) in samples collected from Illinois and Iowa in fall 1995. Bars with the same letter within a state are not significantly different according to paired *t* test (*P* ≤ 0.05).
The survival of plant pathogens in crop residues has been well-documented (27), and for many plant pathogens, surface crop residues, especially in no-till fields, have been associated with increase in diseases (13,16,21,22,29). In samples from the no-till fields at the Nashua research farm, the percentage of leaf disks colonized by *Phytophthora sojae* significantly correlated with residue dry weight. However, in the conventional-till fields, the relationship between residue dry weight and recovery of *P. sojae* was not as clear. The lack of a clear relationship may be attributed to the fact that buried residues decompose faster than surface residues (9), and the survival ability of *P. sojae* in fragmented and decomposing soybean tissues in the soil may have been reduced.

Generally, soybean seeds are planted within the top 5 to 6 cm of the soil. The relative abundance of propagules of *P. sojae* in no-till fields may have important implications for damping-off of soybean. In no-till fields, germinating seeds may be exposed to greater risk of pre- and post-emergence damping-off than in the conventional-till fields in which *P. sojae* densities were lower in the seed placement zone. Meyer and Sinclair (15), in working with the same pathogen and host, showed that stem lesions develop when the inoculum is placed at 1 or 4 cm below the soil surface, but not at 9 cm. They further demonstrated that root and shoot dry weights of soybean increase with increase in depth of inoculum placement.

No-till fields maintain more surface residues and are more compact than conventional-till fields, at least early in the season (12). Consequently, no-till fields retain more moisture (3,19) than conventional-till fields, a condition that favors development of damping-off (24). Compaction (17), long saturation periods, and high propagule densities of *P. sojae* may increase the risks of damping-off in no-till fields.

The leaf-disk bioassay is a semiquantitative technique, and there are advantages and disadvantages to its use in estimating propagule densities in the soil (11,31). In this method, the production of sporangia and zoospores is enhanced through a wetting and drying cycle. The enhancement of spore production may lead to colonization of all or most of the leaf disks if there are initially high population densities of the fungus in the soil. Consequently, the method may fail to detect differences between treatments. However, this technique has been successfully used in the past for other *Phytophthora* spp. when the fungus was at low population densities (18,33). In fact, the percentage of leaf disks colonized was shown to be correlated with disease severity in the field better than the conventional dilution plating method (18). In our study, we had a high frequency of recovery of *P. sojae* only in samples collected from Iowa during the fall of 1995 (nearly 100% of the leaf disks were colonized in approximately 50% of the samples), and that may have resulted in the lack of difference observed between the tillage systems.

In the regional samples, the history of the individual fields, such as the number of years under a particular tillage management regime, might have helped refine our results, but such information was not available. However, by pairing no-till and conventional-till fields at each location, consistent results across the states were obtained. The level of inoculum in the soil is only one aspect of the disease triangle, especially for diseases caused by *Phytophthora* species, in which environmental factors play a major part. In this investigation, *P. sojae* was recovered in greater frequency near the soil surface in no-till fields than in conventional-till fields. In light of this result, we believe that the potential for disease caused by *P. sojae* in no-

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**Fig. 4.** Relationships between soil depth and recovery of *Phytophthora sojae* in no-till (continuous lines) and conventional-till fields (dotted lines) at sites 1 (A) and 2 (B) at the Nashua Research Farm. Data point from each depth of no-till fields is paired with that of conventional-till fields to test the difference between the tillage systems; no-till fields had greater recovery of *Phytophthora sojae* in greater abundance further down than near the soil surface in conventional-till fields.

<table>
<thead>
<tr>
<th>Depth (cm)</th>
<th>% leaf discs colonized</th>
<th>No-till</th>
<th>Conv.-till</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>80</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>60</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>40</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25</td>
<td>20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The leaf-disk bioassay is a semiquantitative technique, and there are advantages and disadvantages to its use in estimating propagule densities in the soil (11,31). In this method, the production of sporangia and zoospores is enhanced through a wetting and drying cycle. The enhancement of spore production may lead to colonization of all or most of the leaf disks if there are initially high population densities of the fungus in the soil. Consequently, the method may fail to detect differences between treatments. However, this technique has been successfully used in the past for other *Phytophthora* spp. when the fungus was at low population densities (18,33). In fact, the percentage of leaf disks colonized was shown to be correlated with disease severity in the field better than the conventional dilution plating method (18). In our study, we had a high frequency of recovery of *P. sojae* only in samples collected from Iowa during the fall of 1995 (nearly 100% of the leaf disks were colonized in approximately 50% of the samples), and that may have resulted in the lack of difference observed between the tillage systems.

In the regional samples, the history of the individual fields, such as the number of years under a particular tillage management regime, might have helped refine our results, but such information was not available. However, by pairing no-till and conventional-till fields at each location, consistent results across the states were obtained. The level of inoculum in the soil is only one aspect of the disease triangle, especially for diseases caused by *Phytophthora* species, in which environmental factors play a major part. In this investigation, *P. sojae* was recovered in greater frequency near the soil surface in no-till fields than in conventional-till fields. In light of this result, we believe that the potential for disease caused by *P. sojae* in no-

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**Fig. 4.** Relationships between soil depth and recovery of *Phytophthora sojae* in no-till (continuous lines) and conventional-till fields (dotted lines) at sites 1 (A) and 2 (B) at the Nashua Research Farm. Data point from each depth of no-till fields is paired with that of conventional-till fields to test the difference between the tillage systems; no-till fields had greater recovery of *Phytophthora sojae* than conventional-till fields according to t test ($P = 0.05$ and $P = 0.06$ for sites 1 and 2, respectively).
Fig. 5. Relationships between recovery of *Phytophthora sojae* and residue dry weights in no-till fields at sites 1 (diamonds) and 2 (circles) at the Nashua Research Farm.

till fields is greater than in conventional-till fields.

**ACKNOWLEDGMENTS**
This project was supported by the Iowa Soybean Promotion Board, the Northcentral Soybean Research Program, and Hatch Act and State of Iowa Funds. We thank T. S. Abney for valuable information on soybean production for Indiana. We also thank J. J. Nelson, K. Elbashar, and H. A. Roozen for technical assistance. P. Lundeen and A. Bower helped locate fields in Iowa and Illinois.

**LITERATURE CITED**