Characterization of a New *Streptomyces* Strain, DS3024, That Causes Potato Common Scab

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**ABSTRACT**

A novel strain of *Streptomyces* (named DS3024) was isolated from a potato field in Michigan in 2006. The taxonomy of the organism was determined by morphology, biochemistry, and genetic analysis. Analysis of the 16S ribosomal RNA gene sequence indicated that the organism was most similar to an isolate of *Streptomyces* sp., ME02-6979.3a, which is not pathogenic to potato tubers but is distinct from other known pathogenic *Streptomyces* spp. Strain DS3024 has genes that encode thaxtomin synthetase (*txtAB*), which is required for pathogenicity and virulence, and tomatinase (*tonA*), which is a common marker for many pathogenic *Streptomyces* spp. However, the *nec1* gene (associated with virulence in most pathogenic *Streptomyces* spp.) was not detected. The new strain was capable of growth at pH 4.5, caused necrosis on potato tuber slices, and produced thaxtomin A. In greenhouse experiments, DS3024 caused scab symptoms on potato tubers similar to those caused by *Streptomyces scabies* on tubers of potato cv. Atlantic, which is scab susceptible. We propose that DS3024 is a new strain of *Streptomyces* capable of causing common scab on potato tubers. The prevalence of this strain of *Streptomyces* in potato-producing areas in the north-central United States has not been determined.

Potato common scab (PCS), caused by *Streptomyces scabies* (Thaxt.) Waksman & Henrici, is indigenous in all potato-growing areas in the world (26,33) and is ranked the fourth most important potato disease in the United States (24). Several species of *Streptomyces* can cause PCS but *Streptomyces scabies* is considered to be predominant (18). The disease has little impact on total potato yield but spoils the appearance, quality, and marketability of the tubers (13).

At least 13 different *Streptomyces* spp. have been found to cause PCS on potato worldwide (1,6,17,24). For the past 50 years, studies have mainly focused on *S. scabies* as the prevalent pathogen causing PCS. However, other *Streptomyces* spp. have been reported to be casual agents of PCS, and each differs from *S. scabies* in their response to cultural practices (e.g., irrigation management). In the United States, several strains of *Streptomyces* spp. have been characterized as pathogens in addition to the four officially reported species in the United States: *S. scabies*, *S. acidiscabies*, *S. europaeiscabies*, and *S. stelliscabies* (34). It is likely that more species capable of causing PCS will be found as detection technologies advance. The number of pathogenic *Streptomyces* spp. may be increasing due to horizontal gene transfer, genetic mechanisms by which bacterial genes can be transferred from one species to another (4,26,27). Clusters of genes that are associated with pathogenicity are referred to as pathogenicity islands (PAIs). The transfer of a PAI from pathogenic *Streptomyces* spp. into a nonpathogenic species may result in the emergence of a new pathogen (26,34).

The identification and taxonomy of *Streptomyces* spp. has been based on morphological and physiological characteristics combined with thaxtomin production and pathogenicity tests in vitro and in vivo (33); however, recently, molecular techniques have become more important. Ability to produce thaxtomin toxin is strongly correlated with the pathogen’s pathogenicity. Thaxtomin production can be easily detected with bioassays (23) measured by using liquid chromatography/mass spectrometry (LC/MS) or high-performance liquid chromatography (HPLC) analysis. More importantly, DNA-based methods enable fast and accurate detection of thaxtomin and identification of various *Streptomyces* spp. Although *Streptomyces* taxonomy is complex, phylogenetic analysis using 16S ribosomal (r)RNA gene sequencing and DNA-DNA hybridization have also been used successfully for species-level identification (17,21,27,31).

The highly conserved genes responsible for pathogenicity among genetically diverse *Streptomyces* isolates are ideal molecular markers for determination of pathogenicity. The *nec1* gene, for example, has been used as a genetic marker for potential pathogenicity in breeding programs. This gene encodes a secreted necrogenic protein (14), Nec1, and ectopic expression of *nec1* is sufficient to convert nonpathogenic *S. lividans* into a pathogen (2.3). The *nec1* gene is thought to be highly conserved in both structure and function among unrelated pathogenic *Streptomyces* spp., and is not known to be present in nonpathogenic species (2,17,27). Pathogenic strains of *S. scabies*, *S. acidiscabies*, and *S. turgidiscabies* may produce both Nec1 and thaxtomin A but *nec1* is not required for thaxtomin A production (2,3).

The role of *nec1* is not conclusive. Joshi et al. (14) showed that *nec1* was necessary for pathogenicity but others suspected it to have a subsidiary role in pathogenicity, because it is missing from some other pathogenic strains of *Streptomyces* (2,17,33).

Studies have shown a correlation between pathogenicity and thaxtomin production (2,20,24,25). Mutant *S. scabies* isolates with reduced or undetectable levels of thaxtomin were nonpathogenic or had reduced virulence (10). The *txtA* and *txtB* genes in *Streptomyces* spp. encode a nonribosomal peptide synthase responsible for synthesis of thaxtomin A toxin. They are arranged in an operon and translationally coupled. This operon can be used as a molecular marker for pathogenicity. Loria et al. (26) reported a strong correlation between thaxtomin production and plant pathogenicity in diverse *Streptomyces* spp. In addition to *nec1* and *txtB*, no other pathogenicity factors have been found, although some pathogenic isolates may not produce thaxtomin (27,29,33).

Hence, the *txtAB* and *nec1* genes are presently the best molecular markers for detection of pathogenic *Streptomyces* spp. using polymerase chain reaction (PCR). The appearance of new taxa of scab-causing *Streptomyces* spp. is most likely dependent on horizontal gene transfer, as described above (4,26,27). We isolated genetically and phenotypically distinct *Streptomyces* spp. as part of a preliminary survey in Michigan (7). Although some isolates were distinct from the known ref-
ence strains of pathogenic Streptomyces spp., they were closely related to other nonpathogenic species. For example, some were tolerant to pH 4.5 but, according to 16S sequence analysis, were not *S. acidiscabies*, which is tolerant to low pH conditions (18). In addition, some pathogenic Streptomyces isolates that we obtained lacked either nec1 or toma genes but were pathogenic on potato tubers. One strain of Streptomyces (named DS3024), which was represented by a single isolate, was distinct both genetically and phenotypically from known pathogenic Streptomyces spp. This strain did not belong to any of the known pathogenic Streptomyces spp. but still caused typical PCS symptoms on potato tubers.

Currently, the statewide distribution of DS3024 in Michigan is unknown. The basic biology and genetics of this pathogen should be addressed before working on large-scale distribution surveys and analyses to enable accurate diagnostics. Our objectives in this study were to characterize the Streptomyces strain DS3024 isolated from soil collected from a potato field in Michigan using morphological, physiological, and molecular methods; determine its taxonomic position; and test its pathogenicity on potato in laboratory and greenhouse conditions. A preliminary report of this study was published (7) and the sequence of DS3024 16S rDNA has been submitted to the National Center for Biotechnology Information GenBank (accession FJ238114).

**MATERIALS AND METHODS**

**Isolates of Streptomyces spp.** Streptomyces strain DS3024 was isolated from soil in a potato field in central Michigan. Plants in this field showed symptoms of PCS and the level of disease incidence was high. After harvest on 16 September 2006, a bulk soil sample (2.0 liters total) was collected from 10 sites within the field. Individual samples of about 0.2 liter were collected with an auger to a depth of 15 cm and placed in a paper bag. The samples were combined in the laboratory by hand-shaking in the bag. Plant debris and stones were manually removed and the remaining soil sample was air dried at 20°C for 1 week at 15% relative humidity. Drying eliminates other microorganisms in the soil that require moisture and potentially compete with Streptomyces spp. because the spores of Streptomyces spp. can survive in dry soil. The soil sample (10 g) was suspended in 90 ml of sterile distilled water and shaken at 200 rpm for 10 min at 20°C. The soil solution was serially diluted and spread onto Streptomyces selective medium STR (5). The plates were incubated at 28°C for 14 days. Colonies of characteristic Streptomyces spp. were picked and serially transferred until a pure culture was obtained for further analysis. Reference strains of *S. scabies* (American Type Culture Collection [ATCC] 48173) and *S. acidiscabies* (ATCC 49003) were obtained from the ATCC and Leslie Wanner, United States Department of Agriculture—Agricultural Research Service, Beltsville, MD. The isolates were maintained in long-term storage as spore suspensions in 20% glycerol at −80°C. Strain DS3024 and several other isolates, which survived low pH value on agar media, were selected.

**Morphological and physiological characterization.** The isolates of Streptomyces were maintained on International Streptomyces Project (ISP) Medium 2 (yeast malt extract [YM]) agar plates. The sample used for scanning electron microscopy. The sample for scanning electron microscopy was prepared by the method of Eguchi et al. (8). Characteristics of the spores from 14-day-old cultures on YM agar at 28°C were determined by microscopy. Cultural characteristics of cells in ISP media 2, 6, and 7 were recorded after incubation for 14 days at 28°C by the methods recommended by Shirling and Gottlieb (30). Production of dark, diffusible melanoid pigment was determined on peptone-yeast extract iron agar (ISP medium 6) and in tyroside agar (ISP medium 7). The plates were incubated at 28°C and the morphological descriptions of colonies were recorded after 14 days. Cultures of the isolates were grown on STR medium without antibiotics at pH 4.0, 4.2, 4.5, 5.0, 6.0, and 7.2.

**Molecular characterization.** **Extraction of genomic DNA.** Streptomyces isolates were grown on YME at 28°C for 7 days, and the colonies were removed from the plate with a toothpick and resuspended in 500 µl of lysis buffer (1.86 g of KCl, 0.605 g of Tris-HCl, 500 µl of Tween 20, and 450 ml of water, adjusted to pH 8.3; the final volume was brought up to 500 ml with water). The suspension was boiled for 15 min at 100°C and centrifuged for 10 min at 9,391 × g. The supernatant containing genomic DNA was used for the molecular analysis. The amount and purity of the DNA preparations was checked by a NanoDrop ND-1000 (Thermal Scientific, Wilmington, DE).

**Sequencing of the 16S rDNA gene.** The 16S rDNA gene was amplified from DNA of Streptomyces isolates by PCR using the primers 16S-1F (5′-GATCTCACGGA GGTGATCC-3′) and 16S-1R (5′-AGAAGAGGGTGTACCCAGG-3′) (32). Amplification was carried out in 20 µl of reaction system containing 10 mM Tris-HCl (pH 9.0), 50 mM KCl, 0.1% Triton X-100, 1.0 to 2.0 mM MgCl2, 250 µM each dNTP, 10 pmol each primer, Taq DNA polymerase at 5 U/ml, 2 to 25 ng DNA template, and MilliQ water to bring the volume to 20 µl. The thermal cycler was set to the following conditions: initial denaturation at 95°C for 3 min; followed by 40 cycles of denaturing at 95°C for 20 s, annealing at 60°C for 30 s, and extension at 72°C for 2 min; and ending with a 4°C hold. The PCR product of 1,500 bp was verified by agarose gel electrophoresis and purified by using the QIAquick PCR product purification kit (Qiagen Science, MD). The PCR product was sequenced at Michigan State University Genomic Technology Support Facility (East Lansing, MI). The 16S rDNA gene sequences were aligned with the MEGALIGN program of DNASTAR (DNASTAR Inc., Madison, WI). A phylogenetic tree (an unrooted tree and all branch lengths) was constructed using ClustalX version 2 (19,28), which was based on the neighbor-joining (NJ) method (28). To generate the tree, sequences of 16S rDNAs of all the collected isolates were aligned. The distance (percent divergence) was calculated between all pairs of sequences based on the alignment. A distance matrix was generated by the NJ method. The phylogenetic tree was displayed with the aid of Dendroscope software (12) with 1,000 iterations.

**PCR detection of marker genes.** Strain DS3024 and other Streptomyces spp. isolates were tested by PCR for the presence of genes characteristic of the *S. turgidiscabies* PAI as described by Kers et al. (15) using specific primers: TxB1B (5′-CCA CCAGGACCTGCTCT-3′) and TxB2B (5′-TCAGTTGACCTCAAGAT-3′) to the toma operon, Tom3 (5′-GAGCGGT TTGTTGAGTCTCA-3′) and Tom4 (5′-TTGGGTTGTGACTCTTCGT-3′) to the tomA gene (34), and Nf (5′-ATGACG GCGAACAGGAGCGCGGAG-3′) and Nrf (5′-ATGACG CGGCAGCAG-3′) to the nec1 gene (2). The amplified DNA was separated on a 1.5% agarose gel to verify the size and presence of the expected 385-bp txAB, 398-bp toma, and 700-bp nec1 amplicons. Additional primers for Streptomyces spp. identification based on 16S rDNA were also tested on DS3024. Primers used (34) for Streptomyces spp. included S. scabies and *S. europaeaschabiei* (scab1m: 5′-CGAAGCTT GCGGGATCCGA-3′ and scab2m: 5′-TTCT GACAGCTCCTCCTTAC-3′), *S. stel liscabiei* (Stel3: 5′-GAAACAGCATTAGATGTTGCC-3′ and T2r2: 5′-CGA CGACTCTCCTCCCCCCGAAG-3′), *S. botropensis* (Stel3 and Aciz 2: 5′-CGAGCTCGCTCCTCCCCCCCAAG-3′), *S. acidiscabies* (Aciz 1: 5′-TCTAAGCTTGCAGCA TGCGG-3′ and Aciz 2), and *S. turgidiscabies* (Tur1m: 5′-CCCTCCTGGGATGGTTGGG TCC-3′ and Turg2m: 5′-CGACACGCTT CTCCTCCCCTGGAG-3′), and *S. auroraficiens* (Aur1: 5′-TCCGCATGGGGTGG TG-3′ and Aur2: 5′-TGTAAGTCCCCG ACATTACT-3′).

**Dot blot analysis on nec1 gene.** Dot blot analysis was performed to confirm the absence of the nec1 gene in strain DS3024.
using strain ATCC 49173 as a control. The \textit{nce1} PCR amplicon (700 bp) was labeled as a probe using PCR digoxigenin (DIG) probe synthesis kit (Roche Applied Science, Mannheim, Germany). Total genomic DNA was extracted and purified using a UltraClean DNA Purification Kit (Mo Bio Laboratories, Inc, Carlsbad, CA). The quantity and quality of the eluted DNA were measured using a NanoDrop.

The DNA was diluted in 50 µl of 2× SSC buffer (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate) and denatured by heating at 100°C for 5 min, immediately followed by 5 min of chilling on ice. Two µg of each DNA sample was spotted onto an Amersham Hybond-N+ membrane (GE Healthcare Bio-Sciences Corp., Piscataway, NJ), which was placed on a Whatman 3MM chromatography paper with a vacuum system underneath. The membrane was baked for 2 h at 80°C. Prehybridization (4 h) and hybridization (22 h) were carried out using Roche’s blotting kits (Roche Applied Science). The blot was placed into a hybridization bag containing prewarmed hybridization buffer and incubated for 30 min at 65°C with gentle agitation. After the incubation, the hybridization buffer was poured out and replaced with new hybridization buffer containing DIG-labeled \textit{nce1} probe, which was prepared by heating the probe in hybridization buffer for 5 min and chilling on ice for 5 min. The blot was incubated for 20 h at 65°C with gentle agitation. After the hybridization, the membrane was washed for 1 min in 2× SSC/1% sodium dodecyl sulfate (SDS) followed by 60 min at 68°C in 0.1× SSC/0.1% SDS for two times. The membrane was then incubated in blocking solution for 30 min at room temperature, incubated in antibody solution for 30 min, and washed with washing buffer for two times of 15 min each. It was then equilibrated for 5 min in detection buffer. The CSPD solution (1:100 in detection buffer) was added to the detection buffer and incubated for 5 min. The membrane was dried, autoradiography was performed with Blue Ultra Autorad Film (ISCBioExpress, Kaysville, UT), and film was exposed for 2 min to detect hybridization. This experiment was repeated once.

\textbf{Pathogenicity assay.} \textit{Thaxtomin assay.} The thaxtomin production of isolate DS3024 was examined and compared with \textit{S. scabies} ATCC 49173 using HPLC. The assay was conducted following the method described by Loria et al. (23) with slight modifications. Briefly, 200 ml of oatmeal broth with trace elements was inoculated with 200 µl of a concentrated spore suspension of \textit{Streptomycyes} strains. The cultures were incubated at 28°C at 180 rpm for 7 days. Cultures were centrifuged to remove the cells and particulate materials. The cell pellet was dried and the total weight was measured. Culture supernatants (15 ml) were then extracted twice with an equal volume of ethyl acetate. The combined ethyl acetate extracts were evaporated to dryness and the yellow residue was dissolved in 3 ml of methanol. Thaxtomin A was purchased from Axxora LLC (San Diego, CA) and used as a control standard. Extracts and the thaxtomin A standard were analyzed by LC/MS using a Waters QToF Ultima API mass spectrometer coupled to a Waters 2795 HPLC (Waters, Milford, MA), using electrospray ionization in positive ion mode. A Supelco Ascentis Express C18 column (2.1 by 50 mm, 2.7-µm particle) was used for separation, using a flow rate of 0.3 ml/min and a binary linear gradient consisting of solvent A = 0.1% acetic acid and B = acetoni- trile, programming from 95%A to 10%A from 0 to 7 min and a hold at 10%A.

\textit{Tuber slice assay.} Potato tubers (cv. Atlantic, obtained from Dr. D. Douches in the Crop and Soil Sciences Department at Michigan State University, East Lansing, were disinfected by surface sterilization with 15% bleach (0.94% NaClO) for 2 min followed by rinsing in sterile distilled water; seed pieces were then planted at a 15-cm depth. Three replicate plants (pots) were included in each treatment. Plants were watered every 2 to 3 days and fertil- izer was applied as needed. Potato plants were grown in the greenhouse for 112 days before harvesting and then scored for disease severity. Scab symptoms were evaluated based the following scale: 0 = no symptoms, 1 = 1 to 10% surface area with superficial or raised lesions, 2 = 11 to 25% surface area with superficial or raised lesions, 3 = 26 to 50% surface area with superficial or raised lesions or 6 to 25% pitted lesion area, and 5 = >50% surface area with superficial or raised lesions or >25% pitted area. The experiment was repeated once. Pathogens were isolated from the tuber lesions, purified on STR medium, and genomic DNA of the culture was extracted. PCR was conducted by using primer pair DSF (forward) 5'-TAACACTCGTGTCGCGATGG-3' and DSR (reverse) 5'-TGTCTCCATACGTTTACG-3', following the procedures described above. The primers are specific to DS3024. All statistical analysis was conducted using SAS (version 9.1.3; SAS Institute Inc. Cary, NC).

\textbf{RESULTS}

\textbf{Morphological and physiological characteristics of the strain DS3024.} The morphology of strain DS3024 was consistent with its assignment to the genus \textit{Streptomycyes}. After 14 days of growth, DS3024 produced abundant aerial and vegetative hyphae, which were well developed and exhibited fragmentation. Rough, cylindri- cal spores were borne in long, flexuous chains (Fig. 1). Growth of aerial and sub- strate mycelium was found on all media tested. The aerial mycelium of DS3024 varied from white to dark brown in different media (Table 1). The substrate mycel- ium was colored from yellow to brown. The diffusible pigment melanin was observed on ISP-6 medium. DS3024 can survive at pH 4.5, slightly higher than the minimum of \textit{S. acidiscabies} (pH 4.2) but much lower than that of \textit{S. scabies} (pH 6.0) on STR medium.
Molecular characterization. The PCR analysis of isolate DS3024 revealed the presence of a homolog of the tomatinase gene, tomA. However, the primer pair amplifying the nec1 gene yielded no PCR product (Table 2). Seven different sets of PCR primers were used to amplify the 16S rDNA sequence characteristic of seven plant-pathogenic Streptomyces spp.: S. scabiei, S. europaeiscabiei, S. stelliscabiei, S. bottropensis, S. acidiscabiei, S. turgisiscabiei, and S. aureofaciens. No PCR product was obtained when DS3024 DNA was used as template with those primers, indicating that DS3024 does not belong to any of these species. Dot blot analysis showed that a nec1 DNA probe hybridized with ATCC49173 genomic DNA but not with DS3024 genomic DNA. This result confirmed that DS3024 lacks a nec1 homolog.

The complete sequence of the 16S rRNA gene from DS3024 was determined (submitted to the GenBank, accession number FJ238114) and it had less than 97% similarity to the 16S rRNA gene from the above Streptomyces spp., but was 99.7% identical to the 16S rRNA sequence of Streptomyces sp. ME02-6979.3a (accession EU080943). Both DS3024 and Streptomyces sp. ME02-6979.3a were clearly separated from the other pathogenic Streptomyces spp. but they were more closely related to S. caviscabies and S. luridisca-biei in terms of distance (Fig. 2).

Pathogenicity and thaxtomin production of the strain DS3024. Necrosis was visible 36 h after inoculation on potato tuber disks inoculated with either DS3024 or ATCC 49173. Necrotic lesions continued to expand over a period of 7 days. No necroses or any color changes of the tuber disks were observed when noninoculated agar was used (Fig. 3A). In greenhouse experiments, scab lesions averaging 5 mm in depth were observed in potato tubers grown in soil infested with ATCC 49173 and with DS3024 (Fig. 3B). Typical symptoms caused by DS3024 on potato tubers were deep-pitted scabs, with slight variations on the borders of the lesions on the skin, which showed either a clear cut (Fig. 3B, middle) or a gradual change (Fig. 3B, right) from healthy to diseased tissue. The disease severity of ATCC49173 ranged from 0 to 4, with an average of 0.7 ± 1.16, whereas the disease severity of DS3024 ranged from 0 to 3 with an average of 1.0 ± 1.11; however, there was no significant difference between these two, examined with a t test. The result was consistent in all three experiments. A positive PCR result was shown for strain DS3024 on the reisolated culture from the tuber lesion, indicating that the pathogen that caused scab symptom was DS3024. In HPLC analysis, an abundant ion peak for prototoxin thaxtomin A was detected in extracts of DS3024 at m/z 439.19, with a retention time of 4.37 min. Compared with the standard sample, the actual concentration of thaxtomin in the extract solution was calculated as 10.99 µg/ml or 628 µg/g of cells. The extract of ATCC 49173 yielded a barely detectable signal at the same mass and retention time. This indicates that DS3024 produces more thaxtomin than ATCC 49173.

**DISCUSSION**

This study demonstrated that DS3024 produced a greater amount of thaxtomin, a required pathogenicity factor in potato scab-causing *Streptomyces* isolates (16,23,24), than the ATCC 49173 strain. The presence of the thaxtomin synthesis genes was also confirmed by PCR using txtAB primers. Strain DS3024 was aggressive and caused PCS symptoms on potato tubers in the greenhouse. Morphologically, DS3024 did not look like any of the known pathogenic *Streptomyces* spp. This evidence collectively suggested that DS3024 is a new potato pathogen in the genus *Streptomyces*.

**Table 1.** Morphological and physiological characteristics of strain DS3024 grown on different nutrient media

<table>
<thead>
<tr>
<th>Medium*</th>
<th>Color of aerial mycelium</th>
<th>Color of vegetative mycelium</th>
<th>Production of soluble pigment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yeast-malt extract agar (ISP medium 2)</td>
<td>Brown</td>
<td>Yellow</td>
<td>Yellow</td>
</tr>
<tr>
<td>Oatmeal agar (ISP medium 3)</td>
<td>White</td>
<td>White</td>
<td>None</td>
</tr>
<tr>
<td>Inorganic salts-starch agar (ISP medium 4)</td>
<td>White</td>
<td>White</td>
<td>None</td>
</tr>
<tr>
<td>Glycerol-asparagine agar (ISP medium 5)</td>
<td>White</td>
<td>White</td>
<td>None</td>
</tr>
<tr>
<td>Peptone-yeast extract-iron agar (ISP medium 6)</td>
<td>Blackish brown</td>
<td>Brown</td>
<td>Melanin</td>
</tr>
<tr>
<td>Tyrosine agar (ISP medium 7)</td>
<td>Yellowish white</td>
<td>White</td>
<td>None</td>
</tr>
<tr>
<td>Czapek’s agar</td>
<td>White</td>
<td>White</td>
<td>None</td>
</tr>
<tr>
<td>Nutrient agar</td>
<td>White</td>
<td>White</td>
<td>None</td>
</tr>
</tbody>
</table>

*ISP = International Streptomyces Project.

**Table 2.** Comparison of strain DS3024 with *Streptomyces scabies* American Type Culture Collection (ATCC) 49173

<table>
<thead>
<tr>
<th>Criteria for comparisons</th>
<th>ATCC 49173</th>
<th>DS3024</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location</td>
<td>New York</td>
<td>MI</td>
</tr>
<tr>
<td>Sub mycelium color</td>
<td>Light brown</td>
<td>Reddish brown</td>
</tr>
<tr>
<td>Spore color</td>
<td>Gray</td>
<td>Light brown</td>
</tr>
<tr>
<td>Color on yeast malt extract</td>
<td>Orange brown</td>
<td>Whitish yellow</td>
</tr>
<tr>
<td>Thaxtomin</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>16S ribosomal RNA similarity</td>
<td>S. scabiei</td>
<td>Streptomyces sp. ME02-6979.3</td>
</tr>
<tr>
<td>txtAB gene</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>TomA gene</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>nec1 gene</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Pathogenicity</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

**Fig. 1.** Morphological properties of *Streptomyces* strain DS3024. Colonies (left), chain morphology (middle), and ornamentation (right) of spores.
The nec1 gene has been reported to be a virulence factor and it has never been found in nonpathogenic strains of Streptomyces. However, in some pathogenic Streptomyces isolates, nec1 is not required for pathogenicity (29,33). Strain DS3024 has both txtAB and tomA genes and produces scab lesions on potato tubers despite lacking the nec1 gene. Similar situations have been reported in Korea (27). Bukhalid et al. (2) reported that, out of 43 tested isolates, a South African S. scabies isolate (CEK 018) produced thaxtomin A but lacked the nec1 gene (15). Wanner (35) reported a new streptomycete isolated from southeastern Idaho that causes common scab in potato but lacks the nec1 gene. Isolates of S. scabies from Michigan have been previously reported as having txtAB but lacking nec1 and tomA (34). Some of our other isolates of S. scabies from the same location as DS3024, which are distinct from DS3024 but have not been tested for pathogenicity, also showed no nec1. In a preliminary survey in Michigan soil for Streptomyces spp., some isolates from potato tubers had a variation associated with PAI: some had two but some had all three of the marker genes (txtAB, tomA, and nec1; J. J. Hao, unpublished). This observation supports other reports that Streptomyces spp. can be pathogenic without the nec1 gene (33).

Although DS3024 is pathogenic on potato tubers, it does not belong to any of the known pathogenic species of Streptomyces. The morphological properties of isolate DS3024 are consistent with its classification within the genus Streptomyces. Strain DS3024 differs in morphology and production of pigmentation in different ISP media (Tables 1 and 2). Some Streptomyces spp., including DS3024, survived lower pH (as low as 4.2), a condition that was thought to be tolerated only by S. acidiscabies.

A phylogenetic tree constructed on the basis of 16S rRNA gene sequence showed that the Streptomyces spp. that cause potato scab, including S. acidiscabies, S. scabies, and S. turgidiscabies, constitute unique branches. It is evident that strain DS3024 forms a distinct phyletic line from known pathogenic strains and was most closely related to strain ME02-6979.3a, which was described as a potential biological control agent in the GenBank. Strain DS3024 did not look like any of the pathogenic Streptomyces spp. reported earlier. Based on literature and the phylogenetic tree, DS3024 was close to the clade that had S. caviscabies (S. griseus), S. setonii, and S. argenteolus (22); however, none are known in the United States.

Studies have shown some diversity within strains morphologically identified as S. scabies, as well as other phytopathogenic streptomycetes, and provided evidence that potato scab is caused by a polyphyletic group of Streptomyces spp. (11).
These *Streptomyces* spp. are different in ecology, host range, virulence, pH optima, toxin production, and the mechanisms of pathogenicity (18, 27). The case of DS3024 could be an example of the occurrence of horizontal gene transfer within these strains (2). It is possible that strain DS3024 was a nonpathogenic soil inhabitant that received a partial PAI cluster (*txaAB* and *toma*) from another pathogenic *Streptomyces* spp. and then became pathogenic. Genetically, this is very similar to the Idaho strain that Wanner described in terms of PAI genes (35). The absence of a *necl* homolog in DS3024 indicates that *necl* is not directly involved in thaxtomin A production and is not a required pathogenicity factor for all streptomycetes. Hence, our study supports the hypothesis that thaxtomin A production is a pathogenicity determinant in plant-pathogenic *Streptomyces* spp. This finding, along with the apparent lack of *necl* gene in the isolates from Wanner’s (30) study, suggests that North American isolates either contain rearrangements or deletions within the PAI.

In conclusion, isolate DS3024 could be a new pathogenic common scab species in potato. There is need to further investigate whether it is widely distributed. If so, potential risk should be assessed and disease management strategies for this pathogen will require further investigation.

**ACKNOWLEDGMENTS**

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**LITERATURE CITED**


