Nocardiopsis mutabilis, a New Species of Nocardioform Bacteria Isolated from Soil

MARCIA C. SHEARER,1* PAULA M. COLMAN,1 AND CLAUDE H. NASH III2

Department of Natural Products Pharmacology, Smith Kline & French Laboratories, Philadelphia, Pennsylvania 191011 and Sterling-Winthrop Research Institute, Rensselaer, New York 121442

A nocardioform bacterium isolated from soil was studied. On the basis of cell wall composition and physiological characteristics, this organism was placed in the genus Nocardiopsis. This organism differed from the only previously described species of this genus by a number of morphological and biochemical characteristics, including inability to decompose adenine and xanthine, fragmentation in submerged cultures, growth in lysozyme broth, and possession of type PIV phospholipids. It also produced a novel antibiotic, polynitroxin. This organism is regarded as a new species, for which we propose the name Nocardiopsis mutabilis. The type strain of this species is SK&F-AAA025 (= ATCC 31520).

Since instituting analysis of whole-cell hydrolysates as a routine part of our preliminary taxonomic studies, we have found a number of antibiotic-producing actinomycetes which contain meso-diaminopimelic acid and have a type C sugar pattern (13). Analyses of purified cell wall preparations from these organisms have revealed type III cell walls with no characteristic sugars (13). A few of these cultures have very distinct morphological characteristics which make them easy to assign to a genus. Others, however, have morphologies which fall somewhere in that continuous spectrum of morphological characteristics that links Streptomyces with Nocardia. At present, these actinomycetes are very difficult to classify because a very limited number have been isolated and adequately characterized. In this paper we describe the taxonomic characteristics of one of these actinomycetes, strain SK&F-AAA025T (type strain), which produces a novel antibiotic (polynitroxin). The isolation and characterization of the antibiotic have been described elsewhere (T. C. Jain, D. J. Newman, and M. C. Shearer, U.S. patent 4,317,812, March 1982).

MATERIALS AND METHODS

Bacterial strain and culture methods. Strain SK&F-AAA025T was isolated by standard soil dilution and plating techniques from a soil sample taken from a cultivated field in Gujarat, India.

Stock cultures were grown on medium 172 (M172) (1). Slants of this medium were also used for determining the growth temperature range. The media used for morphological observations were M172, thin potato-carrot agar (TPC) (6), and soil extract agar. The soil extract agar contained 1.0 g of dextrose, 0.2 g of K2HPO4, 1,000 ml of soil extract, and 20 g of agar (pH 7.0). The soil extract was prepared by adding 500 g of fertile field soil to 1,200 ml of tap water, autoclaving this preparation for 30 min, and filtering it through paper. Additional media used to characterize strain SK&F-AAA025T were yeast extract-malt extract agar (21), oatmeal agar (21), inorganic salts-starch agar (21), glycerol-asparagine agar (21), tyrosine agar (21), peptone-yeast extract-iron agar (21), tryptone-yeast extract broth (21), defined agar (9), glucose-yeast extract agar (24), gelatin medium 19 (24), Czapek-sucrose agar (Czapek-Dox broth [Difco Laboratories] solidified with agar), nutrient agar (Difco), litmus milk medium (Difco), and D Nase Test Agar (BBL Microbiology Systems).

All tests were performed at 28°C. For growth tests under anaerobic conditions, the GasPak system (BBL) was used.

Microscopy. Whole mounts of sporophores were prepared by the method of Lechevalier and Lechevalier (12) and were viewed with a Siemens model 1A transmission electron microscope.

Chemotaxonomy. Purified cell walls and whole-cell hydrolysates were analyzed by the methods of Becker et al. (2) and Lechevalier (10), respectively. Cell wall phospholipid and mycolic acid analyses were performed by the methods of Lechevalier et al. (11, 14).

Physiological tests. The physiological tests used to characterize strain SK&F-AAA025T were those of Gordon (3, 4) and Gordon and Mihm (5). In the tests for acid production from carbohydrates and utilization of organic acids, all results were confirmed by subculturing onto fresh medium. The tests used for melanoid pigments were those of Shirling and Gottlieb (21).

Susceptibility to penicillin and rifampin. The susceptibility of strain SK&F-AAA025T to penicillin and rifampin was examined by the following method: susceptibility disks (BBL) were placed on M172 plates seeded with a suspension of strain SK&F-AAA025T.
RESULTS

General description. Strain SK&F-AAA025\textsuperscript{T} was gram positive and not acid-fast. It was a filamentous organism that formed a mycelium differentiated into (i) a substrate mycelium that penetrated the agar and formed a compact layer on top of the agar and (ii) an aerial mycelium originating from the substrate mycelium. No motile elements were observed.

Colony morphology. A great variety of colony topography and texture was observed, particularly on rich organic media. There were two basic colony types, but a continuous spectrum of intermediate colony types was also usually present. One basic colony type was light yellow, butyrous, and convex with a smooth surface; this type of colony never produced any visible aerial mycelium. The second basic colony type was yellowish brown, leathery, and conical crateriform; this second colony type became completely covered with a white aerial mycelium. When well-isolated colonies of each type were carefully removed from a plate and transferred to fresh plates, both colony types, along with intermediate colony types, were again observed.

Vegetative mycelium. The vegetative mycelium was well developed. Individual hyphae were long, moderately to densely branched, and about 0.5 to 1.0 \( \mu \text{m} \) in diameter. On any one plate of TPC there were usually areas where the vegetative mycelium had fragmented into short rods and coccolid elements, as well as other areas where the vegetative mycelium appeared to be relatively unfragmented and stable (Fig. 1 through 3).

Extensive microscopic observations of the development of the two basic colony types on plates containing M172, TPC, and soil extract agar revealed that the vegetative mycelia of the yellow, butyrous colonies began to fragment after approximately 4 days on all three media. On plates containing TPC and soil extract agar the vegetative mycelia of the yellowish brown, leathery colonies appeared to be relatively unfragmented after 21 days; a few zig-zag hyphae and numerous gaps of the types described by Uesaka (23) were present. However, on plates...
Nitrate was reduced to nitrite. Milk was peptonized. Gelatin was both hydrolyzed and liquefied. Growth occurred in lysozyme broth. No hydrogen sulfide or melanoid pigments were produced. Esclulin and hippurate were decomposed, but urea and allantoic were not. Catalase and deoxyribonuclease were produced. No growth occurred on 4% NaCl. Acid was produced from L-arabinose, D-cellulbiose, dextrin, dextrose, D-fructose, D-galactose, glycerol, glycogen, L-inositol, lactose, maltose, D-mannose, D-melezitose, melibiose, α-methyl-D-mannoside, D-ribose, salicin, starch, sucrose, trehalose, and D-xylene. No acid was produced from D-erythritol, inulin, or L-sorbose. Acid production during an initial culture in medium containing adonitol, dulcitol, D-mannitol, α-methyl-D-glucoside, raffinose, rhamnose, or D-sorbitol was variable. However, when the strain was subcultured in media containing these carbohydrates, the results were consistent; acid was produced from α-methyl-D-glucoside and raffinose but not from adonitol, dulcitol, D-mannitol, rhamnose, or D-sorbitol.

containing M172, the vegetative mycelia of the yellowish brown, leathery colonies usually fragmented after 4 to 6 days.

**Aerial mycelium.** The aerial hyphae were either unbranched or moderately and irregularly branched; they were either straight or irregularly curved. These hyphae frequently showed nocardoid zig-zags during sporulation (8) (Fig. 4) and completely fragmented into spores (Fig. 5), which were somewhat irregular in length (approximately 0.7 by 1.0 to 2.0 μm). The spores had smooth surfaces (Fig. 6).

**Growth in submerged cultures.** Strain SK&F-AAA025T fragmented when it was grown in a submerged culture. Growth harvested from shaken flasks of tryptone-yeast extract broth on day 4 consisted largely of very short to relatively long rods and coccobacillary units. Branched hyphae were rarely observed.

**Physiological and biochemical characteristics.** Strain SK&F-AAA025T did not grow under anaerobic conditions. The temperature range for growth was 15 to 45°C. Starch, casein, L-tyrosine, and hypoxanthine were hydrolyzed, but adenine and xanthine were not. Nitrate was reduced to nitrite. Milk was peptonized. Gelatin was both hydrolyzed and liquefied. Growth occurred in lysozyme broth. No hydrogen sulfide or melanoid pigments were produced. Esclulin and hippurate were decomposed, but urea and allantoic were not. Catalase and deoxyribonuclease were produced. No growth occurred on 4% NaCl. Acid was produced from L-arabinose, D-cellulbiose, dextrin, dextrose, D-fructose, D-galactose, glycerol, glycogen, L-inositol, lactose, maltose, D-mannose, D-melezitose, melibiose, α-methyl-D-mannoside, D-ribose, salicin, starch, sucrose, trehalose, and D-xylene. No acid was produced from D-erythritol, inulin, or L-sorbose. Acid production during an initial culture in medium containing adonitol, dulcitol, D-mannitol, α-methyl-D-glucoside, raffinose, rhamnose, or D-sorbitol was variable. However, when the strain was subcultured in media containing these carbohydrates, the results were consistent; acid was produced from α-methyl-D-glucoside and raffinose but not from adonitol, dulcitol, D-mannitol, rhamnose, or D-sorbitol.

**FIG. 3.** Micrograph of vegetative mycelium of strain SK&F-AAA025T at intersection of colonies (4-day-old culture on TPC). Bar = 20 μm.

**FIG. 4.** Micrograph of aerial hyphae of strain SK&F-AAA025T showing the nocardoid zig-zag which occurs frequently during spore formation. Bar = 20 μm.
Citrate, malate, succinate, acetate, pyruvate, propionate, and lactate were utilized. Oxalate, tartrate, and benzoate were not utilized. During an initial culture in medium containing formate, utilization was variable; when the strain was subcultured, the results were consistently negative.

**Chemotaxonomy.** Purified cell wall preparations of strain SK&F-AAA025T contained *meso*-diaminopimelic acid, alanine, glutamic acid, glucosamine, and muramic acid; no diagnostic sugars were present. Whole-cell hydrolysates contained galactose, glucose, mannose, ribose, and rhamnose; madurose was absent. The phospholipids found were cardiolipin, diphosphatidylglycerol, lysocardiolipin, phosphatidylethanolamine, phosphatidylethanolamine acylated with hydroxy fatty acids, phosphatidylinositol, and unknown glucosamine-containing phospholipids. Thus strain SK&F-AAA025T has a type III cell wall (13) with a type C whole-cell sugar pattern (13) and a type PIV phospholipid composition (11). No mycolic acids of any type were found in the cell extracts.

Whole-cell hydrolysates were prepared from relatively stable cultures of each of the two basic colonial morphology variants. Both of these hydrolysates contained *meso*-diaminopimelic acid and had a type C sugar pattern (13).

**Appearance on various media.** All plates were incubated for 3 weeks at 28°C in the dark. The colors of the cultures were determined by comparison with chips from the ISCC-NBS Centroid...
NOCARDIOPSIS MUTABILIS SP. NOV.

Color Charts (7, 18) or the Munsell Book of Color (17). The characteristics of the culture on the media tested are given below.

Yeast extract-malt extract agar: growth excellent, raised and wrinkled; vegetative mycelium yellow (ISCC-NBS 89, pale yellow); reverse yellow brown; aerial mycelium abundant, white (ISCC-NBS 263, white); no soluble pigment present.

Inorganic salts-starch agar: growth good; vegetative mycelium yellow (ISCC-NBS 82, vivid yellow); reverse yellow brown; aerial mycelium abundant but thin, white (ISCC-NBS 263, white); light brown soluble pigment variably present.

Oatmeal agar: growth fair to good; vegetative mycelium yellow (ISCC-NBS 89, pale yellow); reverse yellow brown; aerial mycelium abundant but thin, white (ISCC-NBS 263, white); no soluble pigment present.

Glycerol-asparagine agar: growth good; vegetative mycelium yellow (ISCC-NBS 67, brilliant orange yellow); reverse yellow brown; aerial mycelium moderate, white (ISCC-NBS 263, white); light brown soluble pigment variably present.

M172: growth excellent, raised and wrinkled; vegetative mycelium yellow (ISCC-NBS 72, dark orange yellow); reverse yellow brown; aerial mycelium abundant, white (ISCC-NBS 263, white); yellow brown soluble pigment present.

Soil extract agar: growth fair, flat; vegetative mycelium yellow (ISCC-NBS 89, pale yellow); reverse yellow brown; aerial mycelium abundant but thin, white (ISCC-NBS 263, white); no soluble pigment present.

Tyrosine agar: growth good; vegetative mycelium yellow (ISCC-NBS 86, light yellow); reverse yellow brown; aerial mycelium abundant, white (ISCC-NBS 263, white); no soluble pigment present.

Defined agar: growth good, raised and wrinkled; vegetative mycelium yellow (ISCC-NBS 71, moderate orange yellow); reverse yellow brown; aerial mycelium none to sparse, white (ISCC-NBS 263, white); yellow brown soluble pigment present.

Glucose-yeast extract agar: growth good, raised and wrinkled; reverse yellow brown; aerial mycelium none to sparse, white; light yellow brown soluble pigment present.

Nutrient agar: growth fair, thin; reverse light yellow brown; aerial mycelium none to sparse, white; no soluble pigment present.

Czapek-sucrose agar: growth fair, flat; reverse light yellow brown; aerial mycelium sparse, white; light yellow brown soluble pigment present.

TPC: growth fair to good; vegetative mycelium yellow (ISCC-NBS 86, light yellow); reverse yellow brown; aerial mycelium abundant, white (ISCC-NBS 263, white); no soluble pigment present.

Strain SK&F-AAA025T was photochromogenic. The aerial mycelium was white on all media when the cultures were grown in the dark. However, on at least three of the media used in the description of this strain (TPC, tyrosine agar, and oatmeal agar) the aerial mycelium was pale orange yellow (Munsell 10YR 9/2) when cultures were grown in the light.

Susceptibility to penicillin and rifampin. Strain SK&F-AAA025T was resistant to penicillin (10U). Rifampin (5 μg) produced zones of inhibition varying from 10 to 20 mm in diameter. However, these zones always contained satellite colonies, indicating that resistance developed readily.

DISCUSSION

Strain SK&F-AAA025T belongs to the group of organisms described as nocardioform by Prauser (19). Nocardioform actinomycetes reproduce solely by fragmentation of either all of their hyphae, or more or less accidently involved parts of their hyphae, into irregular to rodlike to coccolid elements. However, the generic placement of strain SK&F-AAA025T presents problems. Before the introduction of chemotaxonomy, this organism would most likely have been placed in the genus Nocardiopsis because of the extensive fragmentation which occurs in its vegetative mycelium, but its type III cell wall and complete lack of mycolic acids now preclude this grouping.

On the basis of chemotaxonomy alone, the creation of a new genus might have been justified, as no nocardioform organism with the combination of a type III cell wall and a type PIV phospholipid pattern has been described previously. This was considered unwise because only one strain was available for description and this one strain possesses no distinctive reproductive structures or other morphological features that would make the new genus either easy to describe definitively or easy to recognize readily. However, strain SK&F-AAA025T can be accommodated in the genus Nocardiopsis as described by Meyer (15), and it is this more conservative approach that we have taken. Strain SK&F-AAA025T was compared with the one species previously placed in this genus (22), Nocardiopsis dassonvillii, by using the neotype strain, N. dassonvillii ATCC 23218 (15). Although the most obvious difference between these two organisms was the variety of colony morphologies that were usually present in plate cultures of strain SK&F-AAA025T, they also differed in several other morphological, physiological, and chemotaxonomic characteristics (Table 1). Therefore, we regard strain SK&F-
Growth at 45°C
Decomposition of:
Adenine
Xanthine
Acid from:
i-Inositol
Lactose
Melezitose
Resistance to lysozyme
Resistance
Growth in submerged culture
Phospholipid type
Almost complete
PIV

AAA025T as a new species, for which we propose the name Nocardiopsis mutabilis (muta'bi'lis. L. adj. mutabilis changeable, variable, inconstant, referring to the variety of colony morphology observed, particularly on rich organic media). Strain SK&F-AAA025, the type strain of N. mutabilis, has been deposited in the American Type Culture Collection, Rockville, Md., as strain ATCC 31520.

ACKNOWLEDGMENTS

We are deeply indebted to Mary P. Lechevalier for the electron microscopy, as well as the phospholipid and pure cell wall analyses. Of even greater value was her advice and encouragement while we were working with this very interesting but difficult organism.

LITERATURE CITED