Phenylobacterium immobile gen. nov., sp. nov., a Gram-Negative Bacterium That Degrades the Herbicide Chloridazon

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Bacteria which utilize the xenobiotic compounds chloridazon, antipyrin, and pyramidon as sole carbon sources were isolated from various soil samples. The 22 strains isolated are similar with respect to morphological, physiological, biochemical, serological, and genetic properties. These bacteria are aerobic gram-negative rods or coccoid rods (0.7 to 1.0 by 1.0 to 2.0 μm) that occur singly, in pairs, or in short chains and are nonmotile and nonsporforming. Physiological and biochemical characteristics and susceptibility to antibiotics were determined. The strains need vitamin B12 as a growth factor; they are catalase positive and weakly oxidase positive and show slight H2S production. All of the other tests which we performed were negative. The nutritional spectrum is extraordinarily limited, with optimal growth on chloridazon, antipyrin, pyramidon, and 1-phenylalanine. Most sugars, alcohols, amino and carboxylic acids, and ordinary complex molecules are not utilized. The bacteria are serologically uniform group of organisms, which are harmless to rats and rabbits. Their guanine-plus-cytosine contents range between 65 and 68.5 mol%. The chloridazon-degrading bacteria are characterized as a new genus, Phenylobacterium, with a single species, Phenylobacterium immobile. The type strain Phenylobacterium immobile strain E (= DSM 1986), is not closely related to any other gram-negative bacterium, as shown by a 16s ribosomal ribonucleic acid partial sequence analysis. This organism is a member of group I of the purple nonsulfur bacteria, but is phylogenetically isolated in this group. Phenylobacterium immobile is remotely related to Pseudomonas diminuta, Rhizobium leguminosarum, rhodopseudomonads, and Aquaspirillum itersonii. Like other members of this group, Phenylobacterium immobile contains 2,3-diamino-2,3-dideoxy-α-glucose in its lipopolysaccharide. The murine equals a normal murine from a gram-negative bacterium. All citric acid cycle enzymes are detectable in Phenylobacterium immobile.

Herbicides, fungicides, and other agrochemicals are distributed in large amounts, and microorganisms which are able to degrade synthetic organic molecules play an important role in the elimination of these chemicals from soil and water. Chloridazon [5-amino-4-chloro-2-phenyl-3(2H)-pyridazinone] is the active ingredient of the herbicide Pyramin, which has been used for more than 20 years for the control of weeds in sugar beet and beet root culture. The decomposition of chloridazon (formerly called pyrazon) is a microbial process, as demonstrated by Frank and Switzer on heat-sterilized soil (17) and by the experiments of Drescher and Otto (13), who studied the breakdown of the herbicide in soil samples under various conditions. In 1969 Engvild and Jensen described the isolation of bacteria with the ability to grow on chloridazon as a sole carbon source (15). At the same time and independently, Fröhner et al. (18) isolated chloridazon-degrading bacteria with properties very similar to those of the organisms described by Engvild and Jensen. From soils from various places in the world additional chloridazon-degrading bacterial strains have been isolated (3, 28, 33; H. Blecher, Ph.D. thesis, University of Tübingen, Tübingen, Federal Republic of Germany, 1980). At the moment, 22 different isolates which can grow on the herbicide chloridazon or the structurally related analgesics antipyrin and pyramidon are known.

The chloridazon-degrading bacteria can be characterized as small rods, coccid rods, or cocci. They are aerobic, gram negative, nonmotile, nonpigmented, and nonsporeforming, and they are remarkable for their high nutritional specialization; i.e., they grow optimally on chloridazon, antipyrin (3, 38), pyramidon (4), or L-phenylalanine (7, 42).

None of the previous efforts to identify the chloridazon-degrading bacteria have been successful. In the first attempt, Fröhner et al. tentatively identified chloridazon-degrading bacteria as Acinetobacter species, based mainly on common negative reactions for metabolic properties (18). However, the deoxyribonucleic acid (DNA), guanine-plus-cytosine (G+C) contents of the herbicide-degrading organisms (65 to 68.5 mol%) compared with the values of 38 to 47 mol% for Acinetobacter strains (24) rule out a close relationship between these two groups of gram-negative bacteria. Regarding their preference for unusual carbon sources, the chloridazon-degrading bacteria exhibit a certain degree of similarity to members of the genus Pseudomonas. However, since pseudomonads are defined by motility, the nonmotile chloridazon-degrading bacteria cannot be allocated to this genus.

Recent data on the partial sequence of the 16s ribosomal ribonucleic acid (rRNA) of a chloridazon-degrading bacterium have confirmed the view that these organisms represent a novel, as-yet-undescribed bacterial genus (31). Phylogenetically, the chloridazon-degrading organism is a member of group I of the purple nonsulfur bacteria, showing a remote relationship to Pseudomonas diminuta, Rhizobium leguminosarum, rhodopseudomonads, and Aquaspirillum itersonii, with binary similarity coefficients (SAB values) ranging between 0.51 and 0.46.

In addition, these findings on the phylogenetic position of the chloridazon-degrading bacteria are supported by the results of studies on the lipopolysaccharide composition of this organism (43) and by serological investigations (J. Dorfer, G. Layh, J. Eberspächer, and F. Lingens, unpublished data).
In this paper we present relevant data for the strains of Engvild and Jensen, the strains of Fröhner et al., and 16 additional bacterial strains that are able to degrade the xenobiotic compounds chloridazon, antipyrin, and pyramidon. These bacteria are classified as a new genus and species, for which the name *Phenylobacterium immobile* is proposed.

**MATERIALS AND METHODS**

**Isolation of bacteria.** In our laboratory isolation was achieved as described below. A 300-g sample of soil was mixed with 0.5 g of chloridazon, and the preparation was incubated at 30°C in a flower pot and regularly moistened with deionized water. For the isolation of antipyrin- or pyramidon-degrading bacteria these chemicals instead of chloridazon were added to the soil sample. Degradation of the xenobiotic compounds was followed by thin-layer chromatography. When decomposition was complete (usually in 1 to several weeks, depending on the soil) 5-g samples of the active soil were placed into Erlenmeyer flasks containing 50 ml of mineral salts medium supplemented with the carbon source required at a concentration of 0.4%. This culture was incubated on a rotary shaker at 30°C, and degradation was monitored by thin-layer chromatography. The mixed culture which developed in this culture fluid was subcultured (investigated, 1%) when the breakdown of the xenobiotic compound was complete. After several transfers a sample of the liquid culture was streaked onto agar plates containing the same medium. Single colonies, which normally appeared after 10 days to 3 weeks, were picked and again streaked onto agar. Usually after 5 to 10 transfers pure cultures were obtained. The isolation of strains N and C2 proved to be more difficult, since both of these chloridazon-degrading organisms were closely associated with other bacteria. In the case of strain C2 the other bacterium was identified as *Pseudomonas cepacia*.

**Media.** Two mineral salts media, minimal media A and B, were used. In the first years of study the organisms were cultured with minimal medium A (18). Minimal medium B, with a higher buffer capacity, was developed (37) because when chloridazon-degrading bacteria are grown on chloridazon or antipyrin, they acidify the culture fluid. Minimal medium A contains (per 1,000 ml of deionized water) 0.875 g of KH2PO4, 0.125 g of K2HPO4, 0.5 g of MgSO4·7H2O, 0.1 g of NaCl, 0.1 g of CaCl2·2H2O, 1.0 g of (NH4)2SO4, and the following sources of trace elements (supplied from a stock solution): 0.1 mg of H3BO3, 0.04 mg of CuSO4·5H2O, 0.01 mg of KI, 0.2 mg of FeCl2·4H2O, 0.4 mg of MnSO4·4H2O, 0.4 mg of ZnSO4·7H2O, 0.2 mg of (NH4)2MoO4·4H2O, 0.1 mg of biotin, and 0.03 mg of vitamin B12. The pH value of this medium was adjusted to 6.8 to 7.0 by adding 1 N NaOH. Minimal medium B contains (per 1,000 ml of deionized water) 0.3 g of KH2PO4, 0.7 g of Na2HPO4·12H2O, 0.3 g of (NH4)2HPO4, 0.7 g of (NH4)2HPO4, 0.1 g of (NH4)2SO4, 0.05 g of CaCl2·2H2O, 0.25 g of MgSO4·7H2O, and the same trace elements as minimal medium A. The pH value of minimal medium B is 7.0. The minimal media were supplemented with the required carbon sources at concentrations of 0.04 to 0.1%.

Complex medium A contains (per 1,000 ml of deionized water) 5 g of yeast extract, 10 g of nutrient broth (dehydrated), and 5 g of NaCl. Complex medium B contains (per 1,000 ml of deionized water) 1 g of yeast extract and 1 g of peptone (both from Difco Laboratories, Detroit, Mich.). Medium C is an enriched mineral salts medium and contains (per 1,000 ml of minimal medium A) 1 g of yeast extract (Difco), 1 g of peptone (Difco), and 0.5 g of chloridazon or antipyrin. Complex medium D contains (per 1,000 ml of deionized water) 2.5 g of yeast extract (Difco), 2.5 g of peptone (Difco), 1.5 g of meat extract, 2.5 g of NaCl, and 1 g of chloridazon or antipyrin.

Agar cultures were prepared with 1.5% Bacto-Agar (Difco). Media were sterilized at 121°C for 20 min, and heat-labile compounds were added by sterile filtration.

**Chemicals.** Chloridazon and chloridazon analogs, which were used in growth tests, were a gift from BASF, Ludwigshafen, Federal Republic of Germany. The crude product of chloridazon was purified by repeated washing with acetone. Antipyrin was purchased from Riedel de Haen, Seelze-Hannover, Federal Republic of Germany.

Protease K, pronase, adenosine triphosphate, adenosine diphosphate, adenosine monophosphate, nicotinamide adenine dinucleotide, reduced nicotinamide adenine dinucleotide (NADH), nicotinamide adenine dinucleotide phosphate, reduced nicotinamide adenine dinucleotide phosphate, and coenzyme A were obtained from Boehringer, Mannheim, Federal Republic of Germany. Lysozyme, pancreatic ribonuclease, and lauroylsarcosine were obtained from Serva, Heidelberg, Federal Republic of Germany. Cesium chloride was obtained from Roth, Karlsruhe, Federal Republic of Germany, and 3H-labeled adenine was obtained from Amersham Buchler, Braunschweig, Federal Republic of Germany. The Sephadex G-25 used was from Pharmacia, Uppsala, Sweden. API 20E test systems were obtained from apiobio Merieux, Nürtlingen, Federal Republic of Germany.

**Maintenance of strains.** The isolated bacteria were regularly transferred as agar cultures at intervals of 2 weeks. The purity of the strains was tested at each transfer. Seven strains were stored at the Deutsche Sammlung für Mikroorganismen, Göttingen, Federal Republic of Germany (see Table 1).

**Purity of cultures.** None of the chloridazon-degrading bacterial strains grew on complex medium A; thus, growth on complex medium A plates indicated contamination of the culture. Purity was also checked by light microscopic examination of the bacterial cultures.

**Physiological tests.** The physiological tests were performed for all 22 isolates. The incubation temperature used throughout was 30°C. The nutritional specialization and osmotic sensitivity of the strains did not allow the use of routine media. Therefore, special media on which the bacteria were able to multiply were used.

For catalase and oxidase tests the strains were cultivated on agar plates containing minimal medium B supplemented with chloridazon or antipyrin and on agar plates containing complex medium B. After 7 days of incubation, catalase activity was determined by dropping an H2O2 solution (3%) onto the colonies. Oxidase activity was tested after 5 and 20 days of cultivation on agar plates by using a reagent consisting of solution A [1% naphthol-(1) in 96% ethanol] and solution B (1% N,N,N',N'-tetramethyl-1,4-phenylenediamine dichloride in deionized water); 2 parts of solution A and 3 parts of solution B were mixed and then immediately dropped onto the colonies.

The urease reaction was tested on agar plates containing medium C supplemented with 12 mg of phenol red per liter. The concentration of (NH4)2SO4 in medium C was reduced to 0.01%, and 0.1% urea was added after sterilization. Arginine decarboxylase, lysine decarboxylase, and ornithine decarboxylase activities were tested on agar plates containing minimal medium B supplemented with 1.0 g of antipyrin.
per liter, 1.0 g of the required amino acid per liter, and 20 mg of bromothymol blue per liter (pH 7.0).

Fermentation reactions were tested by using the API 20E system, microtubes containing different carbohydrates, and phenol red as the indicator. The bacteria were suspended in medium C, which for this purpose was adjusted to pH 7.4. Medium C per se becomes neither alkaline nor acidic during growth.

Voges-Proskauer and methyl red tests were performed in tubes containing minimal medium B supplemented with 1.0 g of yeast extract per liter, 1.0 g of peptone per liter, and 2.0 g of glucose per liter (pH 7.0). For the Voges-Proskauer reaction 0.6 ml of 5% naphthol-(1) in 96% ethanol and 0.2 ml of a 40% KOH solution in water were added to 1 ml of the culture.

The indole reaction was tested in flasks containing minimal medium B supplemented with 0.5 g of antipyrin per liter, 0.5 g of yeast extract per liter, 0.5 g of peptone per liter, and 0.5 g of tryptophan per liter. The test reagent was 6-dimethyl-aminobenzaldehyde (5.0 g) in a solution containing 75 ml of amylalcohol and 25 ml of concentrated HCl.

H₂S formation was tested on agar plates containing medium C supplemented with (NH₄)₂Fe(III) citrate and either 2.0 g of Na₂S₂O₃ per liter or 2.0 g of L-cysteine per liter.

The formation of nitrite from nitrate was tested in tubes containing medium C supplemented with 1.0 g of NaN₂O₃ per liter; (NH₄)₂SO₄ was omitted. The test reagents used were 0.5 g of sulfuric acid in 15 ml of 5 N acetic acid and 0.1 g of α-naphthylamine dissolved in 20 ml of boiling water to which 130 ml of 5 N acetic acid was added. The occurrence of fluorescent pigments was tested on agar plates containing modified King medium B (1.0 g of yeast extract, 1.0 g of peptone, 1.0 g of glycerol, 1.0 g of MgSO₄·7H₂O, and 1.8 g of KH₂PO₄·7H₂O in 1 liter of deionized water). The pH value was adjusted to 7.5.

Gelatin hydrolysis was tested in gelatin stabs containing medium C supplemented with 12% gelatin.

Starch hydrolysis was tested on agar plates containing medium C supplemented with 0.5 g of starch per liter. After 3 weeks of cultivation, the dishes were flooded with iodine solution and examined for clearing zones.

Esculin hydrolysis was tested on agar plates containing minimal medium B supplemented with 0.5 g of antipyrin per liter, 0.5 g of yeast extract per liter, 0.5 g of peptone per liter, and 1.0 g of (NH₄)₂Fe(III) citrate per liter (pH 7.0). Esculin (0.1%) was added after sterilization.

Cellulose hydrolysis was tested in tubes containing medium C and filter paper strips rising above the culture liquid. The tubes were observed for 12 weeks.

Casein hydrolysis was tested on agar plates containing medium C supplemented with 0.2% casein. After 4 weeks of incubation the dishes were flooded with 20% acetic acid and examined for clearing zones.

**Growth tests.** The growth of chloridazon-degrading bacteria on different carbon sources was tested either on agar plates containing minimal medium A supplemented with the compound being tested at a concentration of 0.05% or in liquid cultures in 100-ml Erlenmeyer flasks containing 50 ml of minimal medium B supplemented with the carbon source at a concentration of 0.1% or 0.04%. Agar plates were inoculated with a loopful of chloridazon-degrading bacteria suspended in saline, and incubation was for 5 weeks. Liquid cultures were inoculated with 0.5 ml from a logarithmic culture of chloridazon-degrading bacteria in minimal medium B supplemented with either 0.1% chloridazon, 0.1% antipyrin, or 0.1% L-phenylalanine as the carbon source.

Erlenmeyer flask were shaken at 30°C, and growth was followed turbidimetrically. For determinations of doubling times and cell yields, cell numbers were evaluated microscopically with a cytometer chamber.

**Antibiotic susceptibility.** The bacteria were plated onto agar plates containing minimal medium B supplemented with chloridazon or antipyrin. The plates were incubated at 30°C for 20 h, and then disks and multidisks (Oxoid Ltd., London, England) containing the amount of antibiotic desired were placed onto the agar surfaces. After incubation for 3 days at 30°C, inhibition zones were measured.

**Determination of DNA G+C contents.** Using the method of Flossdorf (16), we subjected a total of 2.5 × 10⁷ cells from a chloridazon-degrading strain together with the same number of Escherichia coli cells (for reference DNA) to lysis at 56°C for 10 min with Na-lauroyl sarcosine and ethylenediaminetetraacetate in citrate buffer (pH 7.0). Then the protein was digested by proteinase K treatment at 56°C for 1 h. The lysate was mixed with a 58.1% cesium chloride solution and centrifuged at 44,000 rpm and 25°C for 24 h in an analytical ultracentrifuge (model E; Beckman Instruments, Munich, Federal Republic of Germany) equipped with a type An-FTi rotor.

**Isolation of DNA and hybridization experiments.** Bacteria (2 g) grown in minimal medium B containing antipyrin or chloridazon were suspended in 75 ml of 0.1 M tris(hydroxymethyl)aminoenethane hydrochloride buffer (pH 7.0). Lysozyme [60 mg in 5 ml of tris(hydroxymethyl)aminoenethane hydrochloride buffer] and 10 ml of 0.25 M ethylenediaminetetraacetate (pH 8.0) were added, and the suspension was incubated at 37°C for 15 min, cooled in an ice bath, and treated with 0.5 g of sodium dodecyl sulfate in 10 ml of tris(hydroxymethyl)aminoenethane hydrochloride buffer. The cell lysate, together with 60 ml of chlorororm and 120 ml of phenol solution, was shaken for 45 min at 30°C. To prepare the phenol solution freshly distilled phenol was saturated with 0.03 M tris(hydroxymethyl)aminoenethane-0.005 M ethylenediaminetetraacetate-0.05 M NaCl (pH 8.0). After shaking, the aqueous layer was separated by centrifugation, and 3 parts of 96% ethanol was added. The precipitated DNA was collected with a glass rod. The isolated DNA was dissolved in sodium citrate buffer (0.015 M sodium citrate, 0.15 M NaCl, pH 7.0) and treated with 5 mg of pancreatic ribonuclease per mg of DNA. The DNA concentration was determined by the method of Burton (8). Finally, the solution was again deproteinized with phenol-chloroform. The DNA was precipitated with ethanol and dissolved in citrate buffer. For hybridization experiments the DNA was concentrated in a cesium chloride gradient. Hybridization was performed by the membrane filter method, as described by Denhardt (11) and Goebel and Schrempf (20). For DNA labeling bacteria were grown in minimal medium B containing antipyrin or chloridazon, and during the exponential growth phase [³H]adenine (0.1 mCi) was added.

**Isolation and analysis of murein.** Murein was isolated by the method of Braun and Sieglin (9). Frozen cells were thawed and broken by shaking with glass beads. The cell envelopes obtained after differential centrifugation were treated by boiling in 4% sodium dodecyl sulfate, followed by pronase treatment. For amino acid analyses, the samples were hydrolyzed in 4 N HCl for 15 h at 100°C and analyzed with a Unichrom amino acid analyzer containing a long column packed with type M82 resin (Beckman Instruments). Under these conditions muramic acid is well separated from glutamic acid without a temperature change. For lysozyme
digested murein (about 2 mg/ml) was shaken in 0.1 M ammonium acetate buffer supplemented with 75 μg of lysozyme per ml for 15 h at 37°C. To remove undegraded murein, the mixture was centrifuged for 1 h at 40,000 × g. The supernatant was concentrated in a rotary evaporator and applied to a Sephadex G-25 column for the removal of enzymes. The resulting viscous mixture was taken up in 6 ml of 0.01 M potassium phosphate buffer (pH 7.0) and centrifuged for 20 min at 20,000 × g. The supernatant was used as a crude extract; this crude extract was used for most tests. For molecular weight determinations and for kinetic studies the crude extract was partially purified by fractionated ammonium sulfate precipitation. The activities of the citric acid cycle enzymes were tested by previously described methods (30).

RESULTS

Isolation of strains. Since 1969 22 bacterial strains have been isolated by using the xenobiotic compounds chloridazon, antipyrin, and pyrimidin as sole carbon sources. These bacteria were isolated from all over the world. The isolation procedure was performed at 30°C. Most soil samples have never been treated with the herbicide chloridazon before they were incubated in the laboratory. All of the chloridazon-degrading bacterial strains are listed in Table 1.

Attempts to isolate microorganisms which degrade chloridazon or antipyrin under anaerobic conditions have not been successful (B. Schink and N. Pfennig unpublished data; E. Keller, unpublished data). All efforts to isolate chloridazon-degrading bacteria which grow at 37°C have also failed. For this purpose soil and water samples originating from hot springs, from near-volcanic regions, and from regions with tropical climates were subjected to chloridazon enrichment.

Morphology. (i) Colonies. The chloridazon-degrading bacteria show optimal growth on agar plates containing minimal media and chloridazon, antipyrin, or L-phenylalanine as the only carbon source. With incubation of 30°C, single colonies are not visible before 1 week, and after 3 weeks the colonies are 1 to 2 mm in diameter, circular with entire edges, slightly raised, and not adherent to the agar. The colonies of strains K2, K3, K5, J1, L, M11, N, R, and A12 are smooth and shiny and can readily be emulsified in saline or water; all other strains have rough and dry colonies which cannot be emulsified in saline, where clumping occurs. In water after vigorous agitation emulsification is possible.

(ii) Cells. The chloridazon-degrading bacteria are rods, coccid, rods, or cocci (0.7 to 1.0 by 1.0 to 2.0 μm) and occur singly, in pairs, in short chains, or in clumps. The cells of all strains are nonmotile, nonpigmented, and nongranule and do not produce sheaths or pseudohyphae.

Strains A6, A11, A13, A14, ET, M13, M15, ZS, Z6, and Z7 form exclusively coccal rods, which tend to clump, and in liquid cultures growth is sometimes flocculent. All other strains have short rod-shaped cells which occur singly, in pairs, or in short chains. These strains never clump in liquid cultures. For the most part chains are found in cultures of strains which form smooth colonies on agar plates (strains K2, K3, K5, J1, J2, L, M11, N, R, and A12) and are smooth and shiny and can readily be emulsified in saline or water; all other strains have rough and dry colonies which cannot be emulsified in saline, where clumping occurs. In water after vigorous agitation emulsification is possible.

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In old cultures, especially if the bacteria are cultivated in medium D, which allows only poor growth, pleomorphic forms such as long rods (1.0 by 2.0 to 4.0 μm), long chains (10 to 50 μm) of cells connected by small filaments, or club-shaped and elliptical forms, may occur.

Gram staining, Ziehl-Neelsen staining, and capsule staining are negative. Electron microscopy of thin sections of chloridazon-degrading bacteria also reveals the typical gram-
FIG. 1. Phase-contrast photomicrographs of *Phenylobacterium immobile* strains E<sup>T</sup>, K<sub>2</sub>, N, and A<sub>13</sub>.

negative cell wall pattern (Fig. 2). Figure 3 shows thin sections of ruthenium red-treated bacteria, which reveal the presence of a microcapsule surrounded by a slime layer of acid polysaccharides for strain K<sub>2</sub>, one of the strains which form smooth colonies. Following the definition of Costerton et al. (10), the chloridazon-degrading bacteria possess a "flexible" capsule, which can be made visible only by electron microscopy, but no "rigid" capsule, which represents the typical Indian ink-excluding capsule type.

**Physiological and biochemical characteristics.** The chloridazon-degrading bacteria are aerobic with respiratory (never fermentative) metabolism. The temperature for optimum
growth is 28 to 30°C. The strains do not grow at 4 and 37°C. At 37°C the bacteria die within a few days. Growth occurs between pH 6.5 and 8.0; no growth occurs at pH 4.0 and 9.0. Growth is optimal between pH 6.8 and 7.0. Minimal medium A containing chloridazon, antipyrin, or pyramidon becomes acidic (pH 6 in the stationary phase); complex medium B becomes alkaline (pH 8.2). Vitamin B12 is required as a growth factor.

All strains produce a greenish-yellow nonfluorescent pigment in liquid media supplemented with chloridazon or antipyrin. In liquid cultures containing L-phenylalanine as the sole carbon source a yellowish-green fluorescent pigment is formed. When L-phenylalanine is added in concentrations of 3 to 5 g/liter, the yield of the fluorescent pigment increases. Strain C2 is the only strain to form dark brown colonies on agar plates together with dark coloration of the surrounding agar, especially in L-phenylalanine-containing cultures.

The nutritional specialization and osmotic sensitivity of the herbicide-degrading bacteria did not allow use of routine media for biochemical characterization. Therefore, the tests were performed in special media, as described above. The biochemical characteristics, which were determined for all strains, and the physiological characteristics are as follows: strictly anaerobic; metabolism respiratory, never fermentative; optimum temperature for growth, 28 to 30°C; optimum pH for growth, 6.8 to 7.0; vitamin B12 required as growth factor; NH4+ and NO3− used as nitrogen sources; N2 not used as a nitrogen source; negative for denitrification; catalase positive; weakly oxidase positive; urease negative; no hydrolysis of gelatin, casein, starch, agar, or esculin; no indole reaction; methyl red and Voges-Proskauer tests negative; weak H2S production from thiosulfate or L-cysteine; litmus milk negative; no acid or gas produced from N-ace-
tyl-glucosamine, adonitol, amygdalin, amylose, D(-)-arabinose, L(-)-arabinose, dextrin, dulcitol, erythritol, D(-)-fructose, galactose, D(+) glucose, glycerol, glycogen, m-inositol, inulin, lactose, maltose, mannitol, D(-)-melezitose, D(+)melibiose, methyl-D-glucopyranoside, methyl-D-mannopyranoside, methyl-D-xylopyranoside, D(-)-raffinose, rhamnose, ribose, saccharose, salicin, sorbitol, starch, D(-)-trehalose, D(+)xylose, or L(-)-xylose.

FIG. 2. Ultrathin section of a cell from Phenyllobacterium immobile type strain E (= DSM 1986). cm, Cytoplasmic membrane; p, periplasm; m, outer membrane; c, capsule (flexible capsule).

Utilization of carbon sources. The different strains were isolated in mineral salts media containing either chloridazon, antipyrin, or pyramidon as the sole carbon source (Table 1). All 22 strains do not utilize all three substrates; the differences are summarized in Table 3. Chloridazon and antipyrin are well utilized by most strains; pyramidon is utilized by only 7 of the 22 isolates. When pyramidon is added to media containing antipyrin or chloridazon, an inhibitory effect is observed.

A number of aromatic or heterocyclic compounds that are structurally related to chloridazon or antipyrin were tested as possible carbon sources. Some of these compounds and the results of growth tests with them are shown in Fig. 4. Most chloridazon analogs with altered heterocyclic moieties were good growth substrates; however, substitution at the aromatic nucleus made the compound inaccessible for the bacteria (o-, m-, p-methylchloridazon, o-nitrochloridazon).

If the pyridazinone molecule bears two chlorine substituents without an amino group, it is no longer a growth substrate for chloridazon-degrading bacteria, and when added to chloridazon-containing media, this compound acts as a growth inhibitor. Only poor growth was observed with compounds structurally related to antipyrin, like isopropylphenazon, amino-antipyrin, and N-phenylpyrrolidinon.

Of the aniline derivatives tested, N-methylacetanilide and N-methylformanilide were poor growth substrates, and acetanilide, N-dimethylaniline, N-methylaniline, and aniline were not utilized. In addition, the following aromatic com-

- fructose, galactose, D(+) glucose, glycerol, glycogen, m-inositol, inulin, lactose, maltose, mannitol, D(-)-melezitose, D(+)-melibiose, methyl-D-glucopyranoside, methyl-D-mannopyranoside, methyl-D-xylopyranoside, D(-)-raffinose, rhamnose, ribose, saccharose, salicin, sorbitol, starch, D(-)-trehalose, D(+)xylose, or L(-)-xylose.

The susceptibilities of the 22 strains to 17 different antibiotics are given in Table 2.

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- fructose, galactose, D(+) glucose, glycerol, glycogen, m-inositol, inulin, lactose, maltose, mannitol, D(-)-melezitose, D(+)-melibiose, methyl-D-glucopyranoside, methyl-D-mannopyranoside, methyl-D-xylopyranoside, D(-)-raffinose, rhamnose, ribose, saccharose, salicin, sorbitol, starch, D(-)-trehalose, D(+)xylose, or L(-)-xylose.

The susceptibilities of the 22 strains to 17 different antibiotics are given in Table 2.

Utilization of carbon sources. The different strains were isolated in mineral salts media containing either chloridazon, antipyrin, or pyramidon as the sole carbon source (Table 1). All 22 strains do not utilize all three substrates; the differences are summarized in Table 3. Chloridazon and antipyrin are well utilized by most strains; pyramidon is utilized by only 7 of the 22 isolates. When pyramidon is added to media containing antipyrin or chloridazon, an inhibitory effect is observed.

A number of aromatic or heterocyclic compounds that are structurally related to chloridazon or antipyrin were tested as possible carbon sources. Some of these compounds and the results of growth tests with them are shown in Fig. 4. Most chloridazon analogs with altered heterocyclic moieties were good growth substrates; however, substitution at the aromatic nucleus made the compound inaccessible for the bacteria (o-, m-, p-methylchloridazon, o-nitrochloridazon).

If the pyridazinone molecule bears two chlorine substituents without an amino group, it is no longer a growth substrate for chloridazon-degrading bacteria, and when added to chloridazon-containing media, this compound acts as a growth inhibitor. Only poor growth was observed with compounds structurally related to antipyrin, like isopropylphenazon, amino-antipyrin, and N-phenylpyrrolidinon.

Of the aniline derivatives tested, N-methylacetanilide and N-methylformanilide were poor growth substrates, and acetanilide, N-dimethylaniline, N-methylaniline, and aniline were not utilized. In addition, the following aromatic com-
pounds which usually are degraded by various species of bacteria were not utilized by the chloridazon-degrading bacteria: benzene, toluene, biphenyl, phenol, catechol, resorcinol, pyrogallol, benzoic acid, benzoic acid amide, benzoic acid amidine, 2-aminobenzoic acid, 2-,3- and 4-hydroxybenzoic acids, and 2,4-, 2,5-, and 3,5-dihydroxybenzoic acids.

Strain N was the only strain which grew on L-phenylalanine with a normal lag phase of 1 day. All other strains had lag phases of at least 2 to 3 weeks. The long lag phases occurred only when the strains were transferred from chloridazon or antipyrin to L-phenylalanine for the first time; after additional transfers the organisms grew immediately. Several aromatic compounds that are structurally related to L-phenylalanine, like phenylpropionate, phenylpyruvate, and phenylacetate, are also good growth substrates (42). Less effective are phenylacetate, phenylbutyrate, cinnamate, and mandelate, and no growth was observed with L-phenylserine.

The chloridazon-degrading bacteria can use relatively few simple carbon compounds as sole sources of carbon and...
energy. Growth which was much poorer than growth on chloridazon or 1-phenylalanine was observed on L-glutamate, pyruvate, fumarate, succinate, and malate (Table 4). No growth occurred on fructose, galactose, mannose, ribose, xylose, arabinose, saccharose, lactose, maltose, trehalose, inositol, mannitol, erythritol, glycerol, methanol, ethanol, and glycolate, as well as all amino acids except L-phenylalanine and L-glutamate. A mixture of carbon compounds yielded better growth than single carbon compounds, for example, a combination of L-glutamate plus pyruvate, fumarate, succinate, and malate (Table 4). When added to L-glutamate, vitamin B<sub>2</sub>, which in the first studies (15, 18) was found to be a growth factor for chloridazon-degrading bacteria, was routinely added to the mineral salts media at a concentration of 30 µg/liter. For Pseudomonas diminuta and Pseudomonas vesicularis strains a mixture containing pantothenate (400 µg/liter), biotin (400 µg/liter), vitamin B<sub>12</sub> (400 µg/liter), and cystine (50 mg/liter) was necessary for growth (2). The addition of this mixture, however, did not stimulate growth of the chloridazon-degrading bacteria.

Complex media in concentrations which are usual for routine cultivation (10 to 20 g of peptone, or meat extract plus yeast extract per liter) did not support growth of the chloridazon-degrading bacteria. These bacteria were found to be osmotically sensitive, as demonstrated by NaCl addition to the chloridazon-mineral salts medium. Considerable growth inhibition was observed at an NaCl concentration of 5 to 7 g/liter, and total inhibition was observed at an NaCl concentration of 10 g/liter. The chloridazon-degrading bacteria did grow on complex media supplemented with 0.5 to 2 g of peptone plus yeast extract per liter, although growth was considerably slower than growth on chloridazon. Growth was observed but was very poor in medium D containing 6.5 g of yeast extract plus peptone per liter and 2.5 g of NaCl per liter.

**Degradation of chloridazon.** The pathway for the degradation of chloridazon was elucidated by metabolic and enzymatic studies (Fig. 5). In the first step O<sub>2</sub> is incorporated into the benzene nucleus in the 2,3 position by the action of a dioxygenase (37). The resulting cis-dihydro-dihydropyridine derivative is oxidized by a dehydrogenase, yielding a catechol derivative (14). By the action of a meta-cleaving enzyme (34) a yellow compound is formed, which is cleaved by an amidease to form the heterocyclic moiety of chloridazon and 2-hydroxyxuconate (35). 2-Hydroxyxuconate is further degraded via 2-oxo-4-hydroxyvalerate (5), yielding pyruvate and acetaldehyde, two compounds of intermediary metabolism, which are potential sources of carbon and energy for the bacterial cells.

The pathway for the degradation of antipyrin (38) and pyridon (4) closely resembles the pathway summarized above, and in the pathway for the degradation of L-phenylalanine (7) and cinnamate (40) the first steps are the same as those for chloridazon degradation.

**G+C content.** Determinations of DNA G+C molar ratios for the different strains yielded values between 65 and 66.5 mol%. For strain C<sub>2</sub> a slightly higher G+C ratio, 68.5 mol%, was found. If the G+C determination was run with a mixture of DNAs from strain C<sub>2</sub> and strain E<sub>1</sub>, a distinct shoulder was obtained in the ultraviolet scan.

**DNA hybridization.** Radioactive DNA from chloridazon-degrading bacterial strain R was bound to filters containing homologous unlabeled DNA. The effects of excessive amounts of DNAs isolated from different organisms on the binding of radioactive DNA were tested. Chloridazon-degrading bacterial strains E<sub>1</sub>, J<sub>2</sub>, L, N, and R gave 100% competition with strain R. No competition was found with DNAs from Agrobacterium tumefaciens, Bacillus subtilis, Escherichia coli, Pseudomonas fluorescens, and calf thymus.

**Plasmids.** Dye-buoyant density equilibrium centrifugation demonstrated that all strains of chloridazon-degrading bacteria harbor plasmid DNA (23). Agarose gel electrophoresis revealed the presence of at least two different plasmids in all strains, except strain C<sub>2</sub>, which contained only one large plasmid (170 megadaltons) (33). Several strains possessed three, four, or even six plasmids. Regarding plasmid size, considerable variation within the different plasmids of one strain and among the different strains was observed; molecular weights between 8 × 10<sup>6</sup> and 300 × 10<sup>6</sup> were found.
TABLE 4. Doubling times and cell yields in the stationary growth phase of the type strain of *Phenylobacterium immobile* in minimal medium B containing different carbon sources

<table>
<thead>
<tr>
<th>Growth substrate</th>
<th>Conc (g/liter)</th>
<th>Doubling time (h)</th>
<th>Cell yield (cells/ml x 10^6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chloridazon</td>
<td>0.4</td>
<td>7.5</td>
<td>3.2</td>
</tr>
<tr>
<td>Antipyrin</td>
<td>0.4</td>
<td>7</td>
<td>3.8</td>
</tr>
<tr>
<td>Antipyrin</td>
<td>1.0</td>
<td>7.5</td>
<td>9.5</td>
</tr>
<tr>
<td>t-Phenylalanine</td>
<td>4.0</td>
<td>13</td>
<td>13.0</td>
</tr>
<tr>
<td>Phenylpropionate</td>
<td>0.4</td>
<td>8.5</td>
<td>12.5</td>
</tr>
<tr>
<td>l-Glutamate</td>
<td>1.0</td>
<td>32</td>
<td>5.6</td>
</tr>
<tr>
<td>Pyruvate</td>
<td>1.0</td>
<td>44</td>
<td>5.0</td>
</tr>
<tr>
<td>Succinate</td>
<td>1.0</td>
<td>75</td>
<td>2.4</td>
</tr>
<tr>
<td>l-Glutamate + pyruvate</td>
<td>1.0 (each)</td>
<td>20</td>
<td>9.4</td>
</tr>
<tr>
<td>Yeast extract + peptone</td>
<td>1.0 (each)</td>
<td>29</td>
<td>6.0</td>
</tr>
</tbody>
</table>

**Partial sequence analysis of 16S rRNA.** The 16S rRNA from chloridazon-degrading bacterial strain E was isolated and digested with ribonuclease T1. The 5' ends of the oligonucleotides were labeled in vitro with [32P]ATP and phosphatase-free T4 polynucleotide kinase. Fingerprinting and sequence analysis of the labeled oligonucleotides of hexamer size and larger yielded an oligonucleotide catalog (31). This catalog was compared with the oligonucleotide catalogs of more than 400 microorganisms which were analyzed previously by this method. S_{AB} values were calculated from a comparison of the catalogs. On the basis of these results, the chloridazon-degrading organism is not closely related to any bacterium investigated so far with this method. The highest S_{AB} values (0.51), indicating only a remote relationship, were found with *Pseudomonas diminuta* and *Rhizobium leguminosarum*.

**FIG. 4.** Compounds that are structurally related to chloridazon and their utilization as sole sources of carbon and energy.
**Lipopolysaccharide.** The lipopolysaccharide of strain E^{T} was obtained by two-stage extraction with phenol-ethylene-diaminetetraacetate, and its composition was analyzed by a combination of chemical procedures, gas chromatography, mass spectrometry, paper electrophoresis, and enzymatic tests (43). The carbohydrate moiety was found to consist of heptose, 3-deoxyoctulosonic acid, and β-glucose in a molar ratio of 1:2:2.3. Lipid A is composed of 1 mol of 2,3-diamino-2,3-dideoxy-o-glucose and 2 mol of amide-bound and 2.6 mol of ester-bound fatty acids per mol. The amide-bound fatty acids were identified as 3-hydroxydodecanoic acid and 3-hydroxyhexadecanoic acid. R(-)-3-hydroxydodec-5-enoic acid and dodecanoic acid were found to be ester linked. Dodecanoic acid was demonstrated to be linked additionally with the hydroxy groups of the amide-bound fatty acids.

**Murein analysis.** The composition of the murein from strain E^{T}, which was obtained by differential centrifugation of cell envelopes and treatment with sodium dodecyl sulfate, is shown in Table 5. The contents of glutamic acid and alanine were higher than expected for the murein of a gram-negative bacterium. A lipoprotein attached to the murein as described for E. coli was not present in the chloridazon-degrading organism. The murein was further purified by pronase digestion, and the values for amino acids, which were assumed not to be constituents of the murein, dropped considerably. The value for glutamic acid, however, was still too high for a normal murein of a gram-negative bacterium. A comparison of the lysozyme degradation products with those of E. coli by thin-layer chromatography revealed that the main component from the chloridazon-degrading bacterium was identical to the C₇ muropeptide (GlcNAc-MurNAc-Ala-Glu-A2pm-Ala) of E. coli. The amino acid composition of the muropeptides isolated after Sephadex G-25 chromatography of lysozyme-treated murein was also almost identical to that of E. coli (Table 5). The high glutamic acid content have been due to an impurity, probably polyglutamic acid.

**Serology.** By injecting Formalin-treated bacteria intravenously into rabbits, antisera against chloridazon-degrading bacterial strains E^{T}, K₃, M₁₁, and N were prepared. Agglutination tests and immunofluorescence tests revealed the serological uniformity of the 22 chloridazon-degrading strains (25). On the basis of differences in agglutination reactions, antibody titers, and immunofluorescence data, the bacterial strains were classified into five serological subgroups. No serological relationship was found between chloridazon-degrading bacteria and 40 representative gram-negative bacteria, including Acetobacter aceti, Achromobacter parvulus, Acinetobacter calcoaceticus, Agrobacterium tumefaciens, Alcaligenes faecalis, Azospirillum brasilense, Caulobacter sp., E. coli, Flavobacterium sp., Paracoccus denitrificans, Pseudomonas acidivorans, Pseudomonas aeruginosa, Pseudomonas diminuta, Pseudomonas fluorescens, Pseudomonas lemoignei, Pseudomonas putida, Pseudomonas sp., Pseudomonas stutzeri, Pseudomonas testosteroni, Rhizobium sp., Rhodobacterium vannieli, Rhodopseudomonas capsulata, and Salmonella typhimurium. A slight but significant immunofluorescence reaction was observed when antisera against chloridazon-degrading bacteria were allowed to react with Pseudomonas vesicularis, Gluconobacter oxydans, and Aquaspirillum itersonii (Dorfer et al., unpublished data).

**Enzymatic investigations.** (i) Citric acid cycle. The presence and some properties of the enzymes that catalyze citric acid cycle reactions and anaplerotic sequences were tested in a cell-free extract of strain E^{T}. This organism was found to possess all of the enzymes of the citric acid cycle and

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**TABLE 5.** Composition of the murein from the type strain of *Phenylobacterium immobile*^{a}\(^{b}\)

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Murein after differential centrifugation and sodium dodecyl sulfate treatment</th>
<th>Murein after differential centrifugation, sodium dodecyl sulfate treatment, and pronase digestion</th>
<th>Muropeptides from lysozyme-treated murein after gel chromatography</th>
<th>E. coli murein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Muramic acid</td>
<td>1.16</td>
<td>0.82</td>
<td>1.02</td>
<td>0.85</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>3.79</td>
<td>2.49</td>
<td>1.03</td>
<td>1.10</td>
</tr>
<tr>
<td>Alanine</td>
<td>2.49</td>
<td>1.78</td>
<td>1.80</td>
<td>1.77</td>
</tr>
<tr>
<td>Diaminopimelic acid</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Glucosamine</td>
<td>1.08</td>
<td>0.97</td>
<td>0.98</td>
<td>1.0</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>0.15</td>
<td>0.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Threonine</td>
<td>0.14</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serine</td>
<td>0.11</td>
<td>0.04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glycine</td>
<td>0.32</td>
<td>0.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leucine</td>
<td>0.08</td>
<td>0.07</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* The values are related to the concentration of diaminopimelic acid. Other amino acids in acid protein hydrolysates not listed had values of less than 0.03.

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**FIG. 5.** Pathway for degradation of chloridazon.
enzymatic activities for at least four anaplerotic routes (Table 6).

Several properties of citrate synthase, an enzyme assumed to be of taxonomic significance (44), were studied; by using gel filtration on Sephadex G-200, a molecular weight of 120,000 was found, and the pH optimum was determined to be 8.0. The addition of 0.02 M NaCl and 0.02 M adenosine monophosphate had no influence on the enzyme activity. KC1 (0.02 M) slightly stimulated (125%), NH4Cl (0.2 M) inhibited (48%), and MgCl2 (0.2 M) and CaCl2 (0.2 M) completely inhibited the enzyme activity. Adenosine triphosphate (0.01 M) caused 30% inhibition, and 2-ketoglutarate (0.01 M) and NADH (0.25 mM) resulted in complete inhibition of enzyme activity. The addition of KCl did not influence 2-ketoglutarate inhibition, and the addition of adenosine monophosphate to NADH had no influence on the inhibitory effect of NADH.

(ii) Rhodanese. Rhodanese has been demonstrated in various organisms and has been discussed as a tool which has potential taxonomic significance (41).

Rhodanese activity has been found in all strains of the chloridazon-degrading bacteria, and the specific activities, the pH optima, and molecular weights of the enzymes have been determined (26). The enzymes had a pH optimum of 11; molecular weights between 14,000 and 16,000 were found. Specific activities ranged between 0.16 and 1.9 U/mg.

(iii) Arogenate dehydrogenase. The existence of one or two pathways (namely, the 4-hydroxy-phenylpyruvate or arogenate pathway or both) for tyrosine biosynthesis and the properties of the enzymes which catalyze these routes have been used for comparative taxonomic studies (29).

Chloridazon-degrading bacterial strain ET was found to synthesize tyrosine via arogenate exclusively (22). Arogenate dehydrogenase utilized either oxidized nicotinamide adenine dinucleotide or oxidized nicotinamide adenine dinucleotide phosphate as a hydrogen acceptor. This enzyme did not show significant regulation by repression and induction. The activity of the enzyme was not influenced by L-tyrosine or L-phenylalanine.

(iv) meta-Cleaving enzymes. Chloridazoncatechol, an intermediate in chloridazon degradation, is cleaved in meta-proximal position by a chloridazoncatechol dioxygenase. The properties of this enzyme were investigated in 12 strains of chloridazon-degrading bacteria (strains A11, A13, A14, ET, J1, J2, K1, L, M13, M13, M15, and N) and compared with the properties of meta-cleaving enzymes from seven other bacteria (39). These bacteria belonged to various Pseudomonas, Acinetobacter, and Nocardia species, and were isolated with aromatic compounds such as toluene, benzoate, 4-chlorobenzoate, 3-phenylpropionate, and papaverine. Double-diffusion experiments with antibodies prepared against chloridazoncatechol dioxygenase from strain ET and the kinetic behavior of the meta-cleaving enzymes (substrate specificity and requirement for Fe2+ ions for activity) revealed the uniformity of the chloridazon catechol dioxygenases, which seem to form a distinct group among the nonheme iron-containing dioxygenases.

Toxicity of chloridazon-degrading bacteria. Cells of strains ET and N were orally administered to rats for 7 days. No adverse reactions were observed (21). Exposure by air in inhalation experiments did not lead to specific pulmonary changes. Intracutaneous injection of cells did not lead to adverse skin reactions, and intraperitoneal injections of cells did not kill the rats, although bacteria entered the blood. Intravenous injections of living and Formalin-killed bacteria into rabbits during immunization procedures did not lead to toxic effects.

**DISCUSSION**

Homogeneity of the strains. Morphological, physiological, biochemical, and nutritional studies, serological and enzymological data, and the results of DNA G+C determinations and DNA hybridizations have demonstrated the high degree of similarity among the 22 isolates of chloridazon-degrading bacteria. It seems reasonable to group these strains together in a single species. On the other hand, minor but significant differences exist in serological and enzyme properties, in plasmid pattern, and in the utilization of the xenobiotic compounds chloridazon, antipyrine, and pyramidion; this allows us to propose a subgroup classification of the isolates (25, 39).

Failure to identify chloridazon-degrading bacteria. All attempts to identify the chloridazon-degrading bacteria on the basis of routine characters led to unsatisfactory results. The high nutritional specialization of these bacteria, which grow optimally on man-made compounds and utilize poorly only a few normal carbon sources, and the fact that nearly all biochemical tests give negative results are their most distinguishing features. Considering the 19 parts of Bergey's Manual of Determinative Bacteriology, 8th ed. (6), part 7 (gram-negative aerobic rods and cocci) and part 10 (gram-negative cocci and coccobacilli) seem to be most appropriate. Part 10 includes the genus Acinetobacter, which in the first attempt to identify the chloridazon-degrading bacteria was taken into consideration. This genus must be excluded, since the G+C values of the herbicide-degrading organism is not in accordance with Acinetobacter values. More morphological, physiological, biochemical, and nutritional features are shared with the genus Pseudomonas. The general morphological characteristics of Pseudomonas, however, exclude nonmotile organisms. All other genera of part 7 and part 10 show fewer similarities with the organisms of this study.

The view that chloridazon-degrading bacteria are not related to Acinetobacter or Pseudomonas or to numerous other species which were studied for comparison is supported by serological and enzymological investigations. Thus, the results of studies on the molecular and kinetic properties of rhodanese (26), the biosynthetic route for tyrosine synthesis (22), and the substrate specificity, kinet-
ics, and immunology of meta-cleaving enzymes (39) demonstrated that the chloridazon-degrading bacteria have enzyme properties which are clearly different from those of *Pseudomonas* and *Acinetobacter*.

The regulatory properties of the citrate synthase from strain E1 are similar to those of the enzymes from facultatively anaerobic gram-negative bacteria, like members of the *Enterobacteriaceae*. The low molecular weight (120,000) of the citrate synthase and the fact that it was impossible to abolish 2-ketoglutarate inhibition by adding KCl and NADH inhibition by adding adenosine monophosphate are different than the properties described for members of the *Enterobacteriaceae*, thus indicating the exceptional properties of the citrate synthase from the herbicide-degrading organism (44).

**Phylogenetic position of the chloridazon-degrading bacteria.** Serological investigations on the chloridazon-degrading bacteria also demonstrated that these organisms are not closely related to a variety of gram-negative bacteria (25). Only slight immunological relatedness to *Pseudomonas vesicularis*, *Gluconobacter suboxydans*, and *Aquaspirillum iteronii* was detected (Dorfer et al., unpublished data).

A partial sequence analysis of 16S rRNA, which is a powerful method for the elucidation of even remote relationships among microorganisms, also revealed the isolated position of the chloridazon-degrading bacteria (31). The highest *S*AB values (0.51) were found with *Pseudomonas diminuta* and *Rhizobium leguminosarum*. *S*AB values of this magnitude indicate a rather remote relationship, which would not be detectable by DNA-DNA hybridization. A comparison of 16S rRNA nucleotide catalogs showed that the chloridazon-degrading bacteria are members of group I of the purple nonsulfur bacteria (19). *S*AB values ranging from 0.46 to 0.49 were detected with *Rhodopsseudomonas viridis*, *Rhodopsseudomonas capsulata*, *Rhodopsseudomonas palustris* (36), *Rhodopsseudomonas sulfoviridis* (1), *Pseudomonas diminuta* (45), and *Nitrobacter* species (32).

Most of the remote relatives of the chloridazon-degrading bacteria exhibit features which clearly distinguish them from the organisms described in this paper. The rhizobia are described as motile, and they utilize a great variety of carbon sources and have the capacity to nodulate leguminous hosts. Also, the members of the genera *Rhodopsseudomonas* and *Rhodomicrobium*, with their photosynthetic capacity, and *Aquaspirillum*, with its unique morphology, are phenotypically quite different from the chloridazon-degrading bacteria. *Pseudomonas diminuta* was originally defined by its unusual flagellar structure. This organism is mononitricous, and the flagella is very small and tightly coiled (27). Investigations of *Pseudomonas diminuta* and *Pseudomonas vesicularis* strains showed that these organisms utilize a quite limited range of carbon sources and require pantothenate, biotin, and cyanocobalamin as growth factors (2). The nutritional spectrum, the requirement of growth factors, and the DNA G+C values of 66.3 to 67.3 mol% for *Pseudomonas diminuta* and 65.8 mol% for *Pseudomonas vesicularis* are similar to the properties of chloridazon-degrading bacteria. Therefore, the ability of *Pseudomonas diminuta* and *Pseudomonas vesicularis* to utilize chloridazon or antipyrin was tested. The result was negative, and also negative was an experiment to stimulate growth of chloridazon-degrading bacteria by adding the *Pseudomonas diminuta* growth factors. Interestingly, with *Pseudomonas vesicularis* but not with *Pseudomonas diminuta* a weak immunofluorescence reaction with antisera against the chloridazon-degrading bacteria was found (Dorfer et al., unpublished data). Based on the DNA-rRNA hybridization experiments of De Vos and De Ley (12) and a 16S rRNA analysis (C. R. Woese et al., manuscript in preparation), *Pseudomonas diminuta* and *Pseudomonas vesicularis* are misnamed species and not members of the genus *Pseudomonas*. As nonmotile *Pseudomonas*-like organisms, the chloridazon-degrading bacteria show the highest degree of relationship to these misnamed pseudo-pseudomonads, but it is quite clear that they cannot be identified as *Pseudomonas vesicularis* or *Pseudomonas diminuta*.

**Ecological considerations.** The technique for the enrichment of chloridazon-degrading bacteria is based on selective pressure exerted on a microbial population. This technique leads to the isolation of bacteria which are able to utilize synthetic molecules not normally encountered in nature. We suppose that these organisms have sufficient selective advantage only under the conditions of the special isolation process. Obviously, the chemical nature of the selective agent (heterocyclic plus phenyl moiety) is a special challenge for microbial cells, for which other well-described bacterial taxa do not have the right response. This hypothesis is supported by unsuccessful efforts to isolate the same organisms with L-phenylalanine as the selective agent (U. Tittmann and F. Lingens, unpublished data).

Enrichment technique and nutrition characteristics raise the question of which substrates are used by these bacteria in their natural environment. Chloridazon, antipyrin, and pyramidon do not occur in nature; nevertheless, we have shown that these bacteria occur all over the world. Recent experiments with an immunofluorescence membrane filter technique demonstrated that these organisms, or serologically closely related organisms, also occur in soils which have never been treated with the herbicide (Layh and Lingens, unpublished data). This means that these bacteria utilize some other substrate(s) in nature, like L-phenylalanine or structurally related molecules or perhaps mixtures of various organic compounds, like pyruvate, succinate, and L-glutamate, which allow slow growth.

In the last two decades in our laboratory the breakdown of numerous organic molecules, including 4-chlorobenzoate, 4-chlorophenylacetate, 4-toluidine, 4-toluidine, biphenyl, carboxanilide fungicides, caffeine, ephedrine, papaverine, 4-chlorophenylacetate, o-toluidine, p-toluidine, biphenyl, carboxanilide fungicides, caffeine, ephedrine, papaverine, and juglone, has been studied. In all cases and in many other examples described in the literature, bacteria with degradative capacity were isolated and could be identified without difficulty. So far, chloridazon seems to be an exception rather than the standard case.

We propose to introduce the chloridazon-degrading bacteria as a new genus with the name *Phenylobacterium* (Phe.ny.lo.bac'ter.i.um. G. n. phen phen benzene; L. n. bacterium bacterium; *Phenylobacterium* benzene bacterium) due to the unique growth characteristics of these bacteria, which predominantly utilize compounds with a phenyl moiety. All of the chloridazon-degrading bacteria
belong to one species, *Phenylobacterium immobile* (im.mo'-bi.e. L. adj. immobile, nonmotile).

**Description of Phenylobacterium immobile** gen. nov., sp. nov. Cells are rods, coccal rods, or cocci, 0.7 to 1.0 by 1.0 to 2.0 μm, and occur singly, in pairs, or in short chains. Some strains tend to clump. In old cultures pleomorphic forms, such as long rods (1.0 by 2.0 to 4.0 μm), long chains (10 to 50 μm) connected by filaments, and elliptical forms, may occur. Nonmotile. Do not produce sheaths or prosthecae. Resting stages not known. Capsule stain negative; flexible capsule present. Not acid fast. Gram negative.

Growth on glutamate, pyruvate, fumarate, succinate, and 3-amino, antipyrin, pyramidon, and L-phenylalanine. Slow growth on glutamate, pyruvate, fumarate, succinate, and L-phenylalanine. Slow growth with NZ. Osmotically sensitive. Vitamin requirement for optimum growth, 6.8 to 7.0. pH value for optimum growth, 6.8 to 7.0. Isolated from soil after enrichment in mineral media containing chloridazon, antipyrin, or pyrimidin.

Do not denitritify; do not produce nitrite from nitrate. NH₄⁺ and NO₃⁻ are used as sole sources of nitrogen. No acid and no gas produced from sugars and alcohols. Methyl red and Voges-Proskauer negative. Litmus milk negative. Catalase positive. Weakly oxidase positive. Gelatin, casein, starch, and esculin not hydrolyzed. Urease negative. Litmus milk negative. Weak H₂S production from thiosulfate or cysteine. Methyl red and Voges-Proskauer negative. No indole reaction. No acid and no gas produced from sugars and alcohols.

All enzymes of citric acid cycle are present. G+C content: 65.5 mol%.

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