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### Title

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# Complete genome sequence of *Saccharomonospora viridis* type strain (P101<sup>T</sup>)

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## Keywords

thermophile, hot compost, Gram-negative actinomycete, farmer's lung disease, bagassosis, humidifier fever, pentachlorophenol metabolism, *Pseudonocardiaceae* 

## Abstract

Saccharomonospora viridis (Schuurmans et al. 1956) Nonomurea and Ohara 1971 is the type species of the genus Saccharomonospora which belongs to the family Pseudonocardiaceae. S. viridis is of interest because it is a Gram-negative organism classified amongst the usually Gram-positive actinomycetes. Members of the species are frequently found in hot compost and hay, and its spores can cause farmer's lung disease, bagassosis, and humidifier fever. Strains of the species S. viridis have been found to metabolize the xenobiotic pentachlorophenol (PCP). The strain described in this study has been isolated from peat-bog in Ireland. Here we describe the features of this organism, together with the complete genome sequence, and annotation. This is the first complete genome sequence of the family Pseudonocardiaceae, and the 4,308,349 bp long single replicon genome with its 3906 protein-coding and 64 RNA genes is part of the Genomic Encyclopedia of Bacteria and Archaea project.

## Introduction

Saccharomonospora viridis strain  $P101^{T}$  (DSM 43017 = ATCC 15386 = JCM 3036 = NCIMB 9602) is the type strain of the species, which represents the type species of the genus Saccharomonospora [1, 2], which presently contains nine species [3]. Although phylogenetically a member of the Gram-positive actinomycetes, already the initial report on S. viridis strain P101<sup>T</sup> noticed the astonishing feature of the organism to be Gram-negative,

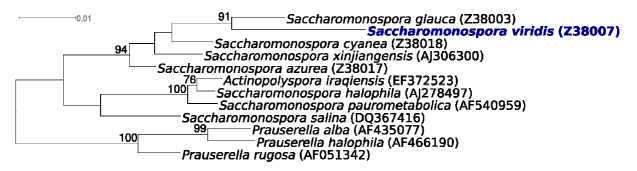
despite showing the typical mycelium morphology of *Saccharomonospora* [2]. Like in other actinomycetes, spores of *S. viridis* are readily dispersed in air, and apparently the prolonged exposure to the antigens of spores can result in acute respiratory distress (farmer's lung disease) which may lead to irreversible lung damage [4, 5].

Here we present a summary classification and a set of features for *S. viridis*  $P101^{T}$  (Table 1), together with the description of the complete genomic sequencing and annotation.

## **Classification and features of organism**

Members of the species *S. viridis* have been isolated or molecularly identified on several occasions from hot composts in Europe and USA, [6-9], and also from soil in Japan [1]. One novel, yet unpublished, cultivated member of the species has been reported by Lu and Liu from Chinese soil (AF127525). Uncultured clone sequences with significant (99%) sequence similarity were observed from composting mass in China (AM930281 and AM930338). Screening of environmental genomic samples and surveys reported at the NCBI BLAST server indicated no closely related phylotypes that can be linked to the species or genus, with the closest matches (about 90% sequence similarity) to strain P101<sup>T</sup> 16S rRNA identified in a marine metagenome from the Sargasso Sea [10].

Figure 1 shows the phylogenetic neighborhood of *S. viridis* strain P101<sup>T</sup> in a 16S rRNA based tree. The sequences of all three copies of the 16S rRNA gene are identical and perfectly match the previously published 16S rRNA sequence generated from NCIMB 9602 (Z38007).

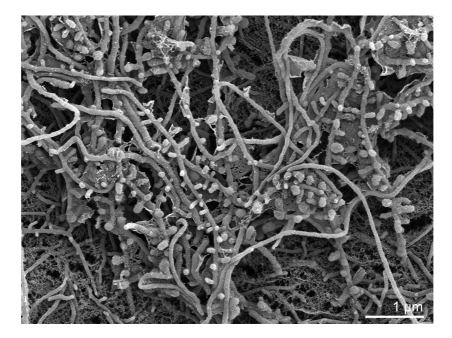


**Figure 1.** Phylogenetic tree of *S. viridis* strain  $P101^{T}$  and all type strains of the genus *Saccharomonospora* inferred from 1474 aligned characters [11, 12] of the 16S rRNA gene under the maximum likelihood criterion [13]. The tree was rooted with all type strains of the members of the genus *Prauserella*, another genus in the family *Pseudonocardiaceae*. The branches are scaled in terms of the expected number of substitutions per site. Numbers above branches are support values from 1000 bootstrap replicates if larger than 60%. Lineages with type strain genome sequencing projects registered in GOLD [14] are shown in blue, published genomes in bold.

The hyphae of the vegetative mycelium of strain P101<sup>T</sup> are branched and sometimes show curved endings [6]. Single spores are observed only on the aerial mycelium either directly on the hyphae or on short sporophores (Figure 2). The spores are oval, 0.9-1.1 $\mu$ m × 1.2-1.4  $\mu$ m in size. Only very occasionally two spores are observed. The aerial mycelium is either greyish green in colour, or turns from white to greenish as on Czapek Agar. The optimal temperature for growth is 55°C, but 45°C for aerial mycelium formation and pigment production. At 37°C and 60°C the growth is very limited and without aerial mycelium. No growth occurs at 27°C and 70°C [6].

Strain P101<sup>T</sup> has been observed to be sensitive to a variety of phages [15]. Members of S. *viridis* are apparently able to metabolize pentachlorophenol but not other chlorophenols [9]. It was suggested that S. viridis metabolizes PCP by conjugation to form a more polar transformation product, but unlike other PCP-degrading bacteria, is incapable of effecting total degradation of the xenobiotic [9]. Microorganisms such as S. viridis may therefore contribute to PCP removal by microbial communities in situ, despite being unable to completely mineralize chlorophenols in pure culture [9]. S. viridis produces a thermostable  $\alpha$ amylase which forms 63% (w/w) maltose on hydrolysis of starch [16]. Maltotriose and maltotetraose are the only intermediate products observed during this reaction, with maltotriose accumulating to 40% (w/w). Both unimolecular and multimolecular mechanisms (transfers and condensation) have been shown to occur during the concentration-dependent degradation of maltotriose and maltotetraose. Such reactions result in the almost exclusive formation of maltose from maltotriose at high initial concentration [16]. S. viridis produces thermoviridin, an antibiotic that is primarily active against the Gram-positive bacteria (growth inhibition) [15, 2]. At higher concentrations, also Gram-negative bacteria were growthinhibited [2].

**Figure 2.** Scanning electron micrograph of *S. viridis* P101<sup>T</sup>



*Chemotaxonomy.* The murein of  $P101^{T}$  is of cell wall type IV. It contains mesodiaminopimelic acid in the peptidoglycan and arabinose and galactose in whole-cell hydrolysates (sugar type A). Mycolic acids and teichonic acids were not reported. Strain  $P101^{T}$  contains menaquinones MK-9(H<sub>4</sub>) (60%) and MK-8(H<sub>4</sub>) (20 to 30%). The combination of the tetrahydromultiprenyl menaquinones MK-9(H<sub>4</sub>) and MK-8(H<sub>4</sub>) is characteristic for the genus *Saccharomonospora* [15]. The major cellular fatty acids are saturated, iso-branched acids with 16 and 18 carbon atoms, and 2-hydroxydodecanoic acids. Details are described in the Compendium of *Actinobacteria* [17]. Phosphatidylethanolamine, hydroxy-phosphatidylethanolamine, and lyso-phosphatidyl-ethanolamine were identified as the main phospholipids.

**Table 1.** Classification and general features of *S. viridis* P101<sup>T</sup> in accordance with the MIGS recommendations [18]

MIGS ID	Property Term		Evidence code <sup>a,b</sup>
		Domain Bacteria	
		Phylum Actinobacteria	
		Class Actinobacteria	TAS [19]
	Current classification	Order Actinomycetales	TAS [19]
	Current classification	Suborder Pseudonocardineae	TAS [19]
		Family Pseudonocardiaceae	TAS [19]
		Genus Saccharomonospora	TAS [1]
		Species Saccharomonospora viridis	TAS [2]
		Type strain P101	
	Gram stain	negative	TAS [2]
	Cell shape	variable	TAS [17]
	Motility	nonmotile	NAS
	Sporulation	single spores on mainly aerial mycelium	TAS [1]
	Temperature range	thermophile, 37-60°C	TAS [15]
	Optimum temperature	55°C for growth, 45°C for aerial mycelium	TAS [1, 6,
	• •	formation	15]
	Salinity	7% NaCl	TAS [15]
MIGS-22	Oxygen requirement	aerobic; nor reported if essential	TAS [15]
	Carbon source	D-glucose, sucrose, dextrin	TAS [15]
	Energy source	carbobydrates	TAS [15]
MIGS-6	Habitat	peat and compost (species occurrence)	TAS [1, 4,
	Haunai	pear and compose (species occurrence)	6, 8, 9]
MIGS-15	Biotic relationship	Free living	
MIGS-14	Pathogenicity	Lung damage	TAS [4]
	Biosafety level	1	TAS [20]
	Isolation	peat-bog at 250 cm depth	TAS [6]
MIGS-4	Geographic location	Irish peat	
MIGS-5	Sample collection time	before 1963	TAS [6]
MIGS-4.1 MIGS-4.2	Latitude – Longitude	not reported	
MIGS-4.3	Depth	not reported	
MIGS-4.4	Altitude	not reported	

a) Evidence codes - IDA: Inferred from Direct Assay (first time in publication); TAS: Traceable Author Statement (i.e., a direct report exists in the literature); NAS: Non-traceable Author Statement (i.e., not directly observed for the living, isolated sample, but based on a generally accepted property for the species, or anecdotal evidence). These evidence codes are from <u>http://www.geneontology.org/GO.evidence.shtml</u> of the Gene Ontology project [21]. If the evidence code is IDA, then the property should have been directly observed, for the purpose of this specific publication, for a live isolate by one of the authors, or an expert or reputable institution mentioned in the acknowledgements.

## Genome sequencing and annotation information

#### **Genome project history**

This organism was selected for sequencing on the basis of each phylogenetic position, and is part of the *Genomic Encyclopedia of Bacteria and Archaea* project. The genome project is deposited in the Genome OnLine Database [14] and the complete genome sequence in GenBank NOT YET. Sequencing, finishing and annotation were performed by the DOE Joint Genome Institute (JGI). A summary of the project information is shown in Table 2.

 Table 2. Genome sequencing project information

MIGS ID	Property	Term	
MIGS-31	Finishing quality	Finished	

MIGS-28	Libraries used	Two Sanger libraries - 8 kb pMCL200 and fosmid pcc1Fos
MIGS-29	Sequencing platforms	ABI3730
MIGS-31.2	Sequencing coverage	12.9x Sanger
MIGS-30	Assemblers	phrap
MIGS-32	Gene calling method	Genemark 4.6b, tRNAScan-SE- 1.23, infernal 0.81
	INSDC / Genbank ID	not yet available
	Genbank Date of Release	not yet available
	GOLD ID	<u>Gi02228</u>
	NCBI project ID	<u>20835</u>
	Database: IMG-GEBA	<u>2500901760</u>
	Project relevance	Tree of Life, GEBA

#### Growth conditions and DNA isolation

S. viridis strain P101<sup>T</sup>, DSM 43017, was grown in DSMZ medium 83 (CZAPEC PEPTONE Medium) at 45°C. DNA was isolated from 1-1.5 g of cell paste using Qiagen Genomic 500 DNA Kit (Qiagen, Hilden, Germany) with a modified protocol for cell lysis including overnight incubation with lysozyme, mutanolysine, lysostaphine, achromopeptidase, and proteinase K at  $35^{\circ}$ C.

#### Genome sequencing and assembly

The genome was sequenced using Sanger sequencing platform only. All general aspects of library construction and sequencing performed at the JGI can be found at <u>http://www.jgi.doe.gov</u>. The Phred/Phrap/Consed software package (<u>www.phrap.com</u>) was used for sequence assembly and quality assessment. After the shotgun stage reads were assembled with parallel phrap (High Performance Soft ware, LLC). Possible mis-assemblies were corrected with Dupfinisher [22] or transposon bombing of bridging clones (Epicentre Biotechnologies, Madison, WI). Gaps between contigs were closed by editing in Consed, custom primer walk or PCR amplification (Roche Applied Science, Indianapolis, IN). A total of 354 finishing reactions were produced to close gaps and to raise the quality of the finished sequence. The completed genome sequences of *S. viridis* contains 66,210 Sanger reads, achieving an average of 12.9x sequence coverage per base, with an error rate less than 1 in 100,000.

#### Genome annotation

Genes were identified using GeneMark [23] as part of the genome annotation pipeline in the Integrated Microbial Genomes Expert Review (IMG-ER) system (http://img.jgi.doe.ogv/er) [24], followed by a round of manual curation using JGI's GenePRIMP pipeline (http://geneprimp.jgi-psf.org) [25]. The predicted CDSs were translated and used to search the National Center for Biotechnology Information (NCBI) nonredundant database, UniProt, TIGRFam, Pfam, PRIAM, KEGG, COG, and InterPro databases. The tRNAScanSE tool [26] was used to find tRNA genes, whereas ribosomal RNAs were found by using the tool RNAmmer [26]. Other non coding RNAs were identified by searching the genome for the Rfam profiles using INFERNAL (v0.81) [28]. Additional gene prediction analysis and manual functional annotation was performed within the Integrated Microbial Genomes (IMG) platform (http://img.jgi.doe.gov/) [29].

#### Metabolic network analysis

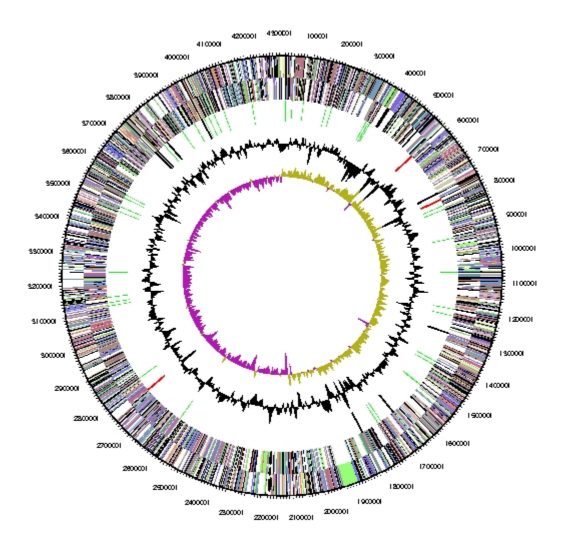
The metabolic Pathway/Genome Database (PGDB) was computationally generated using Pathway Tools software version 12.5 [30] and MetaCyc version 12.5 [31], based on annotated EC numbers and a customized enzyme name mapping file. It has undergone no subsequent manual curation and may contain errors, similar to a Tier 3 BioCyc PGDB [32].

## **Genome properties**

The genome is 4,308,349 bp long and comprises one main circular chromosome with a 67.3% GC content (Table 3). Of the 3970 genes predicted, 3906 were protein coding genes, and 64 RNAs. 78 pseudogenes were also identified. 71.2% of the genes were assigned with a putative function while the remaining are annotated as hypothetical proteins. The properties and the statistics of the genome are summarized in Table 3. The distribution of genes into GOGs functional categories is presented in Table 4.

Attribute	Value	% of Total
Genome size (bp)	4,308,349	
DNA Coding region (bp)	3,805,483	88.33%
DNA G+C content (bp)	2,900,171	67.32%
Number of replicons	1	
Extrachromosomal elements	0	
Total genes	3970	
RNA genes	64	1.61%
rRNA operons	3	
Protein-coding genes	3906	98.39%
Pseudo genes	78	1.96%
Genes with function prediction	2828	71.23%
Genes in paralog clusters	534	13.45%
Genes assigned to COGs	2709	68.24%
Genes assigned Pfam domains	2845	71.66%
Genes with signal peptides	725	18.26%
Genes with transmembrane helices	880	22.17%
CRISPR repeats	9	

 Table 4. Genome Statistics

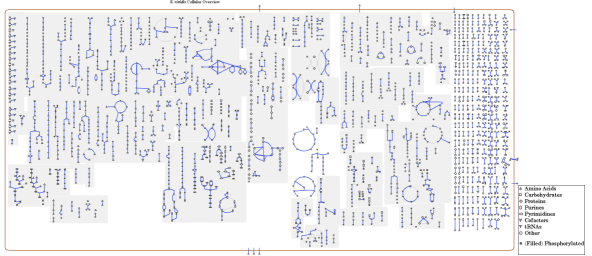


**Figure 3.** Graphical circular map of the genome. From outside to the center: Genes on forward strand (color by COG categories), Genes on reverse strand (color by COG categories), RNA genes (tRNAs green, sRNAs red, other RNAs black), GC content, GC skew.

Code	COG counts and percentage of protein-coding genes Description		
Genome			
	value	% of	
	value	total	
J	158	4.0	Translation, ribosomal structure and biogenesis
А	1	0.0	RNA processing and modification
Κ	276	7.1	Transcription
L	125	3.2	Replication, recombination and repair
В	1	0.0	Chromatin structure and dynamics
D	25	0.6	Cell cycle control, mitosis and meiosis
Y	0	0.0	Nuclear structure

Table 4. Number of genes associated with the 21 general COG functional categories

V	44	1.1	Defense mechanisms
Т	146	3.7	Signal transduction mechanisms
Μ	125	3.2	Cell wall/membrane biogenesis
Ν	2	0.1	Cell motility
Ζ	0	0.0	Cytoskeleton
W	0	0.0	Extracellular structures
U	27	0.7	Intracellular trafficking and secretion
0	107	2.7	Posttranslational modification, protein turnover, chaperones
С	214	5.5	Energy production and conversion
G	214	5.5	Carbohydrate transport and metabolism
Е	293	7.5	Amino acid transport and metabolism
F	85	2.2	Nucleotide transport and metabolism
Η	175	4.5	Coenzyme transport and metabolism
Ι	189	4.8	Lipid transport and metabolism
Р	146	3.7	Inorganic ion transport and metabolism
Q	139	3.6	Secondary metabolites biosynthesis, transport and catabolism
R	389	10.0	General function prediction only
S	182	4.7	Function unknown
-	1197	30.6	Not in COGs



**Figure 4.** Cellular overview diagram. This diagram provides a schematic of all pathways of *Saccharomonospora viridis* strain P101 metabolism. Nodes represent metabolites, with shape indicating class of metabolite (see key to right). Lines represent reactions.

Attribute	Value
Total genes	3970
Enzymes	880
Enzymatic reactions	1155
Metabolic pathways	244
Metabolites	863

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