1	Curtobacterium glycinis sp. nov. from Glycine max, Curtobacterium gossypii sp.
2	nov. from Gossypium hirsutum and Curtobacterium oryzae sp. nov. from Oryza sativa,
3	three new Curtobacterium species and endophytes from agricultural crops
4	
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14	
15	ABSTRACT
16	Three new Curtobacterium species from healthy tissues of agricultural crop plants
17	in the United States are reported. They are Curtobacterium glycinis sp. nov. from soybean
18	in Missouri, Curtobacterium gossypii from cotton in Puerto Rico and Curtobacterium
19	oryzae sp. nov. from rice in Texas.
20	
21	Keywords: novel species, taxonomy, bacteria, prokaryotes, endophytes, agriculture,
22	Actinobacteria, Microbacteriaceae
23	
24	
25	INTRODUCTION
26	Bacterial endophytes of plants have many beneficial interactions with their hosts
27	and play a crucial role in promoting plant health (Finkel et al. 2017; Schenk et al. 2012).
28	Microbiomes of the major agricultural crops including Cotton (Gossypium hirsutum L.),
29	rice (Oryza sativa L.) and soybean (Glycine max (L.) Merr.) have been characterized
30	(Edwards et al. 2015; Liu et al. 2019; Longley et al. 2020; Qiao et al. 2017; Roman-Reyna
31	et al. 2020; Ullah et al. 2019), and beneficial endophytes have been detected (Bertani et al.
32	2016; de Almeida Lopes et al. 2018; Zhou et al. 2018). Plant-associated microorganisms
33	are increasingly used for biotechnological applications, including biological control of

34 plant pathogens, plant growth promotion, or isolation of active compounds (Ryan et al.

35 2008; Glick 2012; Bouizgarne 2013; Dey et al. 2014).

36 Curtobacterium is a genus of Actinobacteria in the family Microbacteriaceae 37 comprising eight validly published species (Parte 2018). Several of the species have been 38 isolated from agricultural plants. These include Curtobacterium albidum shown to 39 counteract salt stress in rice (Vimal et al. 2019), Curtobacterium flaccumfaciens a 40 pathogen of bean, beat, tulip and poinsetta (Collins and Jones 1983), Curtobacterium 41 herbarum from grasses (Behrendt et al. 2002), and Curtobacterium plantarum from corn 42 and soybean seed (Dunleavy 1989). 43 In this study, three *Curtobacterium* strains were isolated from healthy plant tissues 44 in the United States. Strain OG107 from soybean in Missouri, strain VK105 from cotton in 45 Puerto Rico, and strain SS108 from rice in Texas. The strains were characterized using 46 molecular and phenotypic tests to determine their taxonomic placement. Our results 47 indicate that the strains represent three new species of *Curtobacterium*, and we propose the 48 names Curtobacterium glycinis sp. nov. (OG107), Curtobacterium gossypii sp. nov. 49 (VK105) and Curtobacterium oryzae sp. nov. (SS108). These strains are taxonomically 50 separated from the plant pathogens found within this genus.

51 52

METHODS

53 Isolation. Strains OG107 and SS108 were collected from the roots of healthy field-54 grown Glycine max in Missouri and Oryza sativa seedlings in Texas, United States, 55 respectively. Plant tissue was washed with a mild detergent to remove particulates, 56 surface-sterilized with bleach (1% v/v sodium hypochlorite) and ethanol (70% v/v), and 57 homogenized. Serial dilutions of tissue homogenate were plated on a panel of media types 58 for endophyte cultivation. Strain OG107, a small (0.7 mm diameter), pale yellow colony, 59 arose on R2A agar after 5 days of incubation at 24°C, and strain SS108 arose as a non-60 descript colony on Actinomycete Isolation Agar. Both strains were streaked to purity and 61 stored in glycerol (20% v/v) at - 80° C until subjected to further testing.

Strain VK105 was isolated as described in Irizarry and White (2017). Briefly, seeds
of wild, non-cultivated *G. hirsutum* plants were collected roadside in Guayama, Puerto
Rico. Seeds were inoculated on Potato Dextrose Agar (PDA) and incubated at room
temperature (25°C). Strain VK105 arose as an irregular yellow-orange pigmented colony.
The colony was streaked to purity and stocked in 25% glycerol at -80°C until further
analysis.

Motility. Strains were tested for flagellar-dependent swimming and swarming
motility on R2A plates solidified with 0.3% and 0.6% agar, respectively. Three
independent colonies were inoculated onto R2A broth and grown for 36 hr at 24°C. Broth
cultures were normalized to an OD600 of 0.1, and 1.5 μl of culture was spotted directly
onto the surface of the motility agar. The diameter of colony expansion was measured for
5 days.

Carbon source utilization. Substrate utilization was assessed using Biolog GenIII
Microplates (Catalogue No. 1030) (Biolog Inc., Hayward, CA). Each bacterium was
inoculated in duplicate plates using Protocol A described by the manufacturer, with the
exception that plates were incubated at 30°C for 36 hr. Respiration leading to reduction of
the tetrazolium indicator was measured by absorbance at 590 nm.

79 Biochemical analyses. Catalase activity was evaluated by immediate effervescence 80 after the application of 3% (v/v) hydrogen peroxide solution via the tube method, a 81 positive reaction was indicated by the production of bubbles. *Staphylococcus aureus* 82 NCIMB 12702 and Streptococcus pyogenes ATCC 19615 were used as positive and 83 negative controls, respectively. Oxidase activity was evaluated via the oxidation of Kovács 84 oxidase reagent, 1% (w/v) tetra-methyl-p-phenylenediamine dihydrochloride in water, via 85 the filter-paper spot method. A positive reaction was indicated when the microorganism's 86 color changed to dark purple. Pseudomonas aeruginosa NCIMB 12469 and Escherichia 87 coli ATCC 25922 were used as positive and negative controls, respectively.

88 Phylogenetic and genomic analyses. DNA was extracted from pure cultures using 89 the Omega Mag-Bind Universal Pathogen Kit according to manufacturer's protocol with a 90 final elution volume of 60µl (Omega Biotek Inc., Norcross, GA). DNA samples were 91 quantified using Qubit fluorometer (ThermoFisher Scientific, Waltham, MA) and 92 normalized to 100 ng. DNA was prepped using Nextera DNA Flex Library Prep kit 93 according to manufacturer's instructions (Illumina Inc., San Diego, CA). DNA libraries 94 were quantified via qPCR using KAPA Library Quantification kit (Roche Sequencing and 95 Life Science, Wilmington, MA) and combined in equimolar concentrations into one 24-96 sample pool. Libraries were sequenced on a MiSeq using pair-end reads (2x200bp). Reads 97 were trimmed of adapters and low-quality bases using Cutadapt (version 1.9.1) and 98 assembled into contigs using MEGAHIT (version 1.1.2) (Li et al. 2015). Reads were 99 mapped to contigs using Bowtie2 (version 2.3.4) (Langmead and Salzberg 2012), and 100 contigs were assembled into scaffolds using BESST (2.2.8) (Sahlin et al. 2014).

101	Average nucleotide identity analyses were performed using the pyani ANIm
102	algorithm (Richter and Rosselló-Móra 2009) implemented in the MUMmer package
103	(Kurtz et al. 2004) retrieved from https://github.com/widdowquinn/pyani.
104	16S rRNA gene sequences were extracted from genome assemblies using barrnap
105	(Seemann 2019) and submitted to GenBank. Phylogenetic analyses based on the 16S
106	rRNA gene were performed using FastTree (Price et al. 2010) with a General Time
107	Reversible substitution model. Taxon sampling for each species is described in the
108	respective phylogenetic tree figure legend.
109	
110	RESULTS
111	Phylogenetic and genomic analyses
112	Curtobacterium glycinis sp. nov. strain OG107
113	Strain OG107 shared 98.9% 16S rRNA gene sequence identity with
114	Curtobacterium citreum DSM 20528^{T} and less with the remaining Curtobacterium
115	species. A phylogenetic tree using FastTree (Price et al. 2010) confirmed the affiliation of
116	strain OG107 with the genus Curtobacterium. OG107 was most closely related to C.
117	herbarum P 420/07 ^T with 84% bootstrap support (Figure 1). The top average nucleotide
118	identity (ANI) value of OG107 was 89.2% with C. luteum ATCC 15830 ^T . This value was
119	well below the threshold for species demarcation (Richter and Rosselló-Móra 2009; Chun
120	et al. 2018) providing further genomic support that strain OG107 represents a new
121	genomic species of Curtobacterium.
122	
123	Curtobacterium gossypii sp. nov. strain VK105
124	Strain VK105 shared 99.8% 16S rRNA gene sequence identity with
125	Curtobacterium citreum DSM 20528 ^T and less with the remaining Curtobacterium
126	species. A phylogenetic tree using FastTree (Price et al. 2010) confirmed the affiliation of
127	strain VK105 with the genus Curtobacterium. VK105 formed a monophyletic group with
128	the species C. albidum and C. citreum supported by high bootstrap support (Figure 1) and
129	was equally related to either of the two species. Average nucleotide identity (ANI) values
130	of C. albidum DSM 20512^{T} and C. citreum DSM 20528^{T} to VK105 were both 87.7%.
131	These values are well below the threshold for species demarcation (Richter and Rosselló-
132	Móra 2009; Chun et al. 2018) providing further genomic support that strain VK105
133	represents a new genomic species of Curtobacterium.

135	Curtobacterium oryzae sp. nov. strain SS108
136	Strain SS108 shared 99.6% 16S rRNA gene sequence identity with Curtobacterium
137	luteum strain DSM 20542 ^T and less with the remaining Curtobacterium species. A
138	phylogenetic tree using FastTree (Price et al. 2010) confirmed the affiliation of strain
139	SS108 with the genus Curtobacterium. SS108 was most closely related to C. luteum with
140	79% bootstrap support (Figure 1). The top average nucleotide identity (ANI) value of
141	SS108 was 89.2% with C. luteum DSM 20542 ^T . This value was well below the threshold
142	for species demarcation (Richter and Rosselló-Móra 2009; Chun et al. 2018) providing
143	further genomic support that strain SS108 represents a new genomic species of
144	Curtobacterium.
145	
146	Morphology, physiology and biochemical characteristics
147	Curtobacterium glycinis sp. nov. strain OG107
148	Strain OG107 stained Gram-positive. Cells had a slightly curved rod shape (0.2-0.4
149	μm in width and 1-3 μm in length); V and Y shaped forms were observed (Figure 2). They
150	were aerobic, non-sporulating and showed motility. On nutrient agar, colonies were cream,
151	entire, punctiform and convex. After 48 hr, good growth is observed on TSA, R2A and NA
152	at 22 and 30°C. Growth was poor to moderate on the same media and incubation times at
153	37°C. Growth is observed when media are supplemented with 4% NaCl (w/v). Strain OG
154	107 was positive for catalase, but not for oxidase or urease. Being aerobic, the strain has an
155	oxidative metabolism and according to the Biolog system, it is capable of using a range of
156	substrates as carbon sources that include sugars, alcohol sugars and organic acids.
157	Compared to its closest phylogenetic neighbor, C. citreum JCM 1345 ^T , OG17 uses N-
158	Acetyl-D-Galactosamine, D-arabitol, gentobiose, myo-inositol, mannitol and turanose as
159	carbon sources. These tests help to differentiate between the two strains. A detailed
160	description is given in Table 1 and the species description.
161	Curtobacterium gossypii sp. nov. strain VK105
162	The cells of strain VK105 stained Gram-positive, formed short irregular rods (0.4-0.6 μ m
163	in width and 2-4 μ m in length), with cells diving by bending (Figure 3). Y shaped-forms
164	were sometimes visible. Cells were non-motile and did not produce spores. Strain VK105
165	showed good aerobic growth on R2A and TSA agars. On both media, yellow to orange
166	colonies were observed after 2 days incubation at 22 and 30°C. At 7 days, colonies were
167	circular, smooth, glistening and slightly convex. The new strain produced catalase but not
168	oxidase. Strain VK105 has an oxidative metabolism and uses a range of substrates as

169 carbon sources that are also useful for to differentiate against its closest phylogenetic 170 neighbors, Curtocabterium citreum and C. albidum, namely D-arabitol, L-fucose, D-171 glucuronic acid, and myo-inositol. These and other physiological characteristics are 172 presented in Table 1 and the species description. 173 174 Curtobacterium oryzae sp. nov. strain SS108 175 Cells stained-Gram positive, are non-motile, non-spore-forming and rod shaped or 176 slightly curved. In many cases, two cells remained together and formed V shapes (Figure 177 4). Average cell size ranged from 0.2-0.4 µm in width and 1.5 x 2 µm in length. Colonies 178 of strain SS108 were light yellow, entire, round and raised when grown on nutrient agar. 179 After 2 days, good growth was observed on TSA and R2A agars at 22, 30 and 37°C. On 180 nutrient agar, good growth was obtained at 30°C after 2 days and was moderate at 22 and 181 37°C. However, after 7 days, good growth was also seen at 22°C. Aerobic growth. 182 Tolerates 4% NaCl (w/v). Enzyme activity was detected for catalase, but not for oxidase or 183 urease. Several carbon sources are useful to differentiate between the new strain and C. 184 luteum JCM 1480^T and include the utilization of N-Acetyl-D-Galactosamine, D-arabitol, 185 gentobiose, myo-inositol and D-mannitol among others (Table 1). Other phenotypic 186 characteristics are given in the species description. 187 188 189 **DESCRIPTION OF CURTOBACTERIUM GLYCINIS SP. NOV.** 190 191 Curtobacterium glycinis (gly.ci'nis. N.L. fem. gen. n. glycinis of Glycine max, the 192 soybean, referring to the origin of the type strain) 193 Cells are Gram-stain positive, non-spore forming, motile and rod-shaped with V 194 195 and Y shapes produced. Aerobic and chemoorganotrophic. Colonies are circular, smooth, 196 glistening and slightly convex. On R2A and nutrient agar, colonies are cream color and 197 entire, punctiform and convex. Growth is abundant at 22 and 30°C but moderate at 37°C. 198 Grows in the presence of 4% NaCl. Growth is observed between pH 5-7. Positive for 199 catalase; negative for oxidase and urease. The following substrates are used as carbon 200 sources: N-acetyl-D-glucosamine, N-acetyl-D-galactosamine, D-arabitol, D-cellobiose, L-201 fucose, gentiobiose, *myo*-inositol, α -D-lactose, D-mannitol, D-melibiose, β -methyl-D-

202	glucoside, D-raffinose, turanose, D-gluconic acid, D-glucuronic acid, L-alanine, L-
203	glutamic acid (weak), L-rhamnose, D-Glucose-6-PO4.
204	The type strain is resistant to nalidixic acid and aztreonam.
205	
206	The type strain OG 17 ^T was isolated from the roots of healthy field-grown <i>Glycine</i>
207	max in Missouri, USA.
208	
209	
210	DESCRIPTION OF CURTOBACTERIUM GOSSYPII SP. NOV.
211	
212	Curtobacterium gossypii (gos.sy'pi.i. N.L. gen. n. gossypii of Gossypium, the
213	generic name of cotton, referring to the origin of the type strain)
214	
215	Cells are Gram-stain positive, non-motile, non-spore-forming, coccoid or rod-
216	shaped. Aerobic and chemoorganotrophic. Yellow to orange colonies on R2A and TSA
217	agars. Growth on these media is observed after 48 h. Catalase is produced but not oxidase.
218	Grows in the presence of 1 - 4 % NaCl; weak at 8% and at pH 5-7. The following
219	substrates are used as sole carbon sources: dextrin, D-maltose, D-trehalose, D-cellobiose,
220	gentiobiose, sucrose, D-turanose, stachyose, D-raffinose, α -D-lactose, D-melibiose, β -
221	methylD-glucoside, D-salicin, N-acetyl-D-glucosamine, α -D-glucose, D-mannose, D-
222	fructose, D-galactose, inosine, sodium lactate, D-mannitol, D-arabitol, myo-inositol,
223	glycerol, L-aspartic acid, L-glutamic acid, pectin, D-arabitol, D-gluconic acid, D-
224	glucuronic acid and glucuronamide.
225	The type strain is resistant to nalidixic acid and aztreonam.
226	
227	The type strain, VK105 ^T , was isolated from seeds of wild cotton collected in Puerto
228	Rico, USA.
229	
229	
230	DESCRIPTION OF CURTOBACTERIUM ORYZAE SP. NOV.
231	
232	Curtobacterium oryzae (o.ry'zae. N.L. gen. n. oryzae of rice, referring to the origin of the
233	type strain)
234	

235	Cells are Gram-stain positive, non-motile, non-spore-forming and rod shaped or
236	slightly curved; V shapes are formed. Aerobic and chemoorganotrophic. Colonies are light
237	yellow, entire, round and raised on nutrient agar. Good growth is obtained at 22, 30 and
238	37°C on the media tested. Grows on media supplemented with 4% NaCl and at pH 5-7.
239	Positive for catalase, but negative for oxidase or urease. The following substrates are used
240	as carbon sources: N-Acetyl-D-Glucosamine, N-Acetyl-D-Galactosamine, N-Acetyl
241	Neuraminic Acid, D-Arabitol, D-Cellobiose, L-Fucose, Gentiobiose, myo-Inositol, α -D-
242	Lactose, D-Mannitol, D-Melibiose, β -Methyl-D-Glucoside, D-Raffinose, D-Sorbitol,
243	Turanose, D-Gluconic Acid, D-Glucuronic Acid, L-Alanine, L-Glutamic Acid, L-
244	Rhamnose, D-Glucose-6-PO4 and Acetic Acid.
245	The type strain is resistant to nalidixic acid and aztreonam.
246	The type strain, SS108 ^T , was isolated from <i>Oryza sativa</i> seedlings collected in
247	Texas, USA.
248	
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253	
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- 339
- 340

341 TABLES

- 342
- 343 **Table 1.** Physiological characteristics of newly described *Curtobacterium glycinis*, *C*.
- 344 gossypii and C. oryzae species and related Curtobacterium type strains. All strains were
- 345 positive for the assimilation of dextrin, D-fructose, D-galactose, α-D-glucose, D-mannose
- 346 and sucrose (data not shown).

	C. gossypii VK105	C. oryzae OG107	C. glycini s SS108	С. citreu m JCM 1345 ^т	C. pusillu m JCM 1350 ^T	C. luteum JCM 1480 ^T	C. albidum NBRC 15078 ^T	C. herbaru m DSM 14013 ^T
General characteristics:								
Colony pigment		light yellow	ivory	yellow	yellow	yellow	ivory	orange
Motility	-			+	+	+		+
Motility	-	W	W				-	
Oxidase	-	-	-	-	-	-	-	-
Catalase	+	+	+	+	+	+	+	+
Urease	-	-	-	-	-	-	-	nr
Growth with 4% NaCl	+	+	+	nr	nr	nr	nr	nr
Growth with 8% NaCl	+	-	-	nr	nr	nr	nr	nr
Utilization of:								
N-Acetyl-D- Glucosamine	+	+	+	+	-	+	W	nr
N-Acetyl-D- Galactosamine	-	+	+	-	-	-	-	nr
N-Acetyl Neuraminic Acid	-	+	-	nr	nr	nr	nr	nr
D-Arabitol	+	+	+	-	W	-	-	nr
D-Cellobiose	+	+	+	+	-	+	+	nr
L-Fucose	w	+	+	+	-	+	-	nr
Gentiobiose	+	+	+	-	+	-	+	nr
myo-Inositol	+	+	+	-	-	-	-	nr
α-D-Lactose	+	+	+	+	-	+	+	nr
D-Mannitol	+	+	+	-	+	-	W	+
D-Melibiose	+	+	+	+	+	+	W	+
β-Methyl-D- Glucoside	+	+	+	-	+	-	W	nr

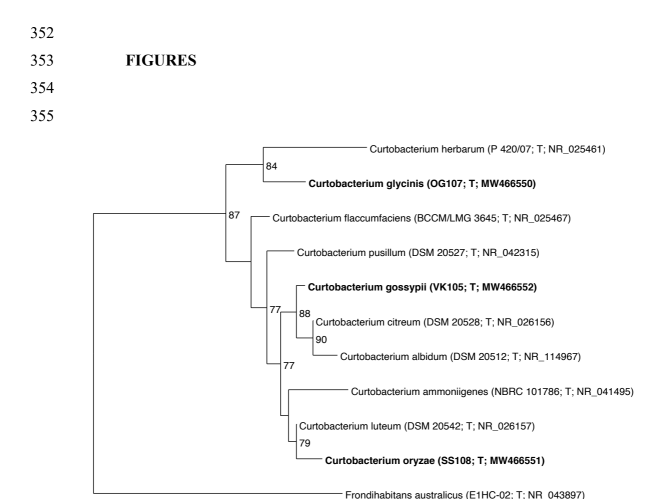
					1			1
D-Raffinose	+	+	+	-	-	-	+	nr
D-Sorbitol	W	+	-	-	+	-	+	+
Turanose	+	+	+	-	+	-	+	nr
D-Gluconic Acid	+	+	+	-	+	-	+	nr
D-Glucuronic Acid	+	+	+	-	-	-	-	nr
L-Alanine	+	+	+	-	-	-	W	nr
L-Glutamic Acid	+	+	W	+	-	-	-	nr
L-Rhamnose	W	+	+	-	-	-	-	+
D-Glucose-6- PO4	-	+	+	-	-	-	-	nr
Acetic Acid	W	+	-	-	-	-	-	nr

347 348

349	+, Positive; w, Weakly positive; -, Negative; nr, Not reported.

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351 Data from this study, Aizawa et al. (2007) and Behrendt et al. (2002).

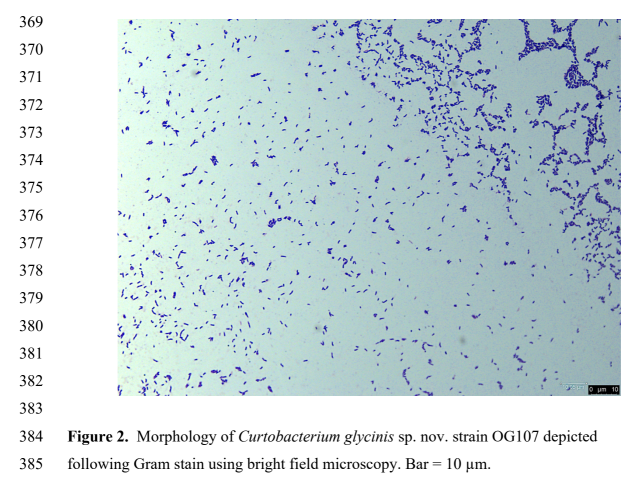


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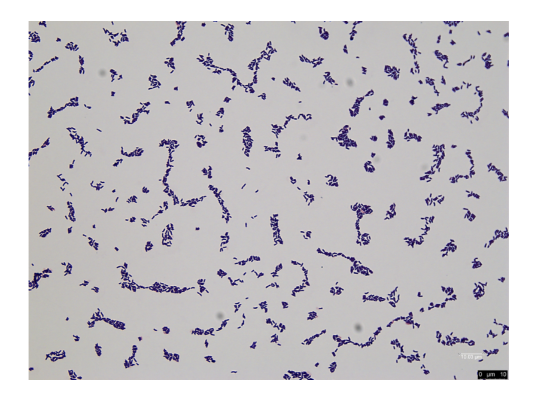
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358 Figure 1. Phylogenetic 16S rRNA gene tree of *Curtobacterium glycinis* sp. nov. 359 strain OG107, Curtobacterium gossypii sp. nov. strain VK105, Curtobacterium oryzae sp. 360 nov. strain SS108 and relatives generated using FastTree (Price et al. 2010). All validly 361 published Curtobacterium species were included in the tree, the tree is rooted with 362 Frondihabitans australicus. New species are in bold. Strain identifiers and GenBank 363 accession numbers follow species names, T stands for 'type'. Support values above 70% 364 are given by the branches. *Curtobacterium glycinis* is most closely related to *C. herbarum* 365 with 84% support, Curtobacterium gossypii to C. albidum and C. citreum with 82% 366 support and Curtobacterium oryzae to C. luteum with 79% support. Branch lengths are 367 proportional to the changes along the branches, a scale bar is provided. 368



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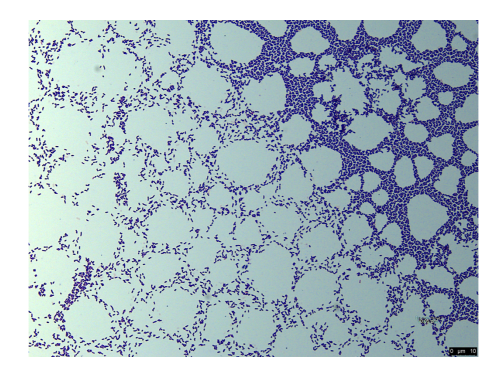


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Figure 3. Morphology of *Curtobacterium gossypii* sp. nov. strain VK105 depicted

- 390 following Gram stain using bright field microscopy. Bar = $10 \mu m$.
- 391



- 392 393
- 394 Figure 4. Morphology of *Curtobacterium oryzae* sp. nov. strain SS108 depicted following
- 395 Gram stain using bright field microscopy. Bar = $10 \mu m$.