# Antonie van Leeuwenhoek Journal of Microbiology Kroppenstedtia pulmonis sp. nov. and Kroppenstedtia sanguinis sp. nov., isolated from human patients --Manuscript Draft--

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Abstract:	Three human clinical strains (W9323T, X0209T and X0394) isolated from lung biopsy, blood and cerebral spinal fluid, respectively, were characterized using a polyphasic taxonomic approach. Comparative analysis of the 16S rRNA gene sequences showed the three strains belonged to two novel branches within the genus Kroppenstedtia: 16S rRNA gene sequence analysis of W9323T showed closest sequence similarity to Kroppenstedtia eburnea JFMB-ATE T (95.3 %), Kroppenstedtia guangzhouensis GD02T (94.7 %) and strain X0209T (94.6 %); sequence analysis of strain X0209T showed closest sequence similarity to K. eburnea JFMB-ATE T (96.4 %) and K. guangzhouensis GD02T (96.0 %). Strains X0209T and X0394 were 99.9 % similar to each other by 16S rRNA gene sequence analysis. The DNA-DNA relatedness was 94.6 %, confirming that X0209T and X0394 belong to the same species. Chemotaxonomic data for strains W9323T and X0209T were consistent with those described for the genus Kroppenstedtia: whole-cell peptidoglycan contained LL-diaminopimelic acid; the major cellular fatty acids were iso-C15 and anteiso-C15; and the major menaquinone was MK-7. Different endospore morphology and carbon utilization profiles of strains W9323T and X0209T supported by phylogenetic analysis enabled us to conclude that the strains represent two new species within the genus Kroppenstedtia, for which the names Kroppenstedtia pulmonis sp. nov. (type strain W9323T =DSM 45752 T) and Kroppenstedtia sanguinis sp. nov. (type strain X0209T =DSM 45749T=CCUG 38657 T) are proposed.	

Responses to reviewer's comments:

Editors comment: it appears from title page that Catherine Spröer and Peter Schumann should have been included as authors but their names are missing from the author list. Please correct.

Response: The author list has been corrected.

Reviewer #1:

Due to strains X0209T and X0394 belong to the same species, I would suggest the strain X0394 will be deleted from the manuscript.

[Editors comment: please retain consideration of strain X0394 - descriptions of taxa are better when based on more than one strain]

Response: We are agreement with the Editor and will leave the X0394 in the manuscript.

The abstract is too rough; please give the key characteristic differences between w9323T and X0209T.

Response: A key characteristic for differentiation between the two novel species is 16S rRNA gene sequence similarity which is stated in the abstract. In the manuscript, the details of phenotypic differences between the strains is discussed in depth. Due to word limitation in the abstract, we have elected not add additional differential phenotypic characteristics to the abstract.

#### Reviewer #3

Author's need to ensure all culture collection certificates are available for the two new species. [Editors comment: please supply these on resubmission]

Response: The certificates are provided here with the revised manuscript.

This is correct in line 60 "plastic surface in a contract manufacturing"

Response: This line is correct.

The main phylogenetic tree should also have the maximum parsimony algorithm included. In addition in the figure legend the references for the algorithms used should be included. Finally in instead of using black dots the initials of the algorithms which the branches were recovered would be more appropriate.

Response: We have edited the tree accordingly and added the reference to the figure legend.

It is not evident if the phenotypic analysis was carried out with type strains from the genus Kroppenstedtia or the work was carried out on just the novel taxa. If the later happened then the work needs to be repeated alongside a type strain from the genus.

Response: As noted in the footnote of Table 2, all data is from the present study except for K. gangzhouensis. This is due to the inability of our lab to acquire the type strain from the culture collections of China and Korea. Unfortunately, the type strain is unavailable from any other resource.

Several references in the reference list are not present in the manuscript please remove: Addou et al., 2012; Felsenstein, 1981; Kampfer et al., 2004; Klude et al., 1969; Mesbah et al., 1989; Rhuland et al., 1955; Saitou & Nei, 1989; Sneath, 1989; Tamura et al., 2011

Response: The references have been edited accordingly.

Antonie van Leeuwenhoek

To Whom It May Concern,

Please find enclosed our manuscript entitled "*Kroppenstedtia pulmonis* sp. nov. and *Kroppenstedtia sanguinis* sp. nov., isolated from human patients" for consideration for publication as an original article in the journal Antonie van Leeuwenhoek. We believe that the enclosed description of these two novel, clinically relevant, species expands our knowledge of the epidemiology of the genus.

All authors have seen and approved the manuscript and I assure that it has not previously been published nor is it under consideration for publication elsewhere. There are no conflicts of interest among the authors. Thank you for your consideration of our manuscript.

Please direct correspondence to: Melissa E. Bell, Bacterial Special Pathogens Branch, Centers for Disease Control and Prevention, 1600 Clifton Road, Atlanta, GA 30333 or at jqv7@cdc.gov, phone: (404) 639-1348, facsimile (404) 639-3022.

Sincerely,

Melissa E. Bell M.Sc. Bacterial Special Pathogens Branch Division of High Consequence Pathogens and Pathology Centers for Disease Control and Prevention

1	Kroppenstedtia pulmonis sp. nov. and Kroppenstedtia sanguinis sp. nov., isolated from human patients		
2	Melissa E. Bell, Brent A. Lasker, Hans-Peter Klenk, Lesley Hoyles, Catherine Spröer, Peter Schumann, June M. Brown		
3			
4	Keywords: Kroppenstedtia species, Kroppenstedtia pulmonis, Kroppenstedtia sanguinis, polyphasic taxonomy, 16S rRNA		
5	gene, thermoactinomycetes		
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#### 28 Abstract

2

- 29 Three human clinical strains (W9323<sup>T</sup>, X0209<sup>T</sup> and X0394) isolated from lung biopsy, blood and cerebral spinal fluid,
- 30 respectively, were characterized using a polyphasic taxonomic approach. Comparative analysis of the 16S rRNA gene
- 31 sequences showed the three strains belonged to two novel branches within the genus *Kroppenstedtia*: 16S rRNA gene
- 32 sequence analysis of W9323<sup>T</sup> showed closest sequence similarity to *Kroppenstedtia eburnea* JFMB-ATE<sup>T</sup> (95.3 %),
- 33 *Kroppenstedtia guangzhouensis* GD02<sup>T</sup> (94.7 %) and strain X0209<sup>T</sup> (94.6 %); sequence analysis of strain X0209<sup>T</sup> showed
- 34 closest sequence similarity to *K. eburnea* JFMB-ATE<sup>T</sup> (96.4 %) and *K. guangzhouensis* GD02<sup>T</sup> (96.0 %). Strains X0209<sup>T</sup> and
- 35 X0394 were 99.9 % similar to each other by 16S rRNA gene sequence analysis. The DNA-DNA relatedness was 94.6 %,
- 36 confirming that X0209<sup>T</sup> and X0394 belong to the same species. Chemotaxonomic data for strains W9323<sup>T</sup> and X0209<sup>T</sup> were
- 37 consistent with those described for the genus *Kroppenstedtia*: whole-cell peptidoglycan contained LL-diaminopimelic acid;
- 38 the major cellular fatty acids were *iso*-C<sub>15</sub> and *anteiso*-C<sub>15</sub>; and the major menaquinone was MK-7. Different endospore
- 39 morphology, carbon utilization profiles, and whole cell wall sugar patterns of strains W9323<sup>T</sup> and X0209<sup>T</sup> supported by
- 40 phylogenetic analysis enabled us to conclude that the strains represent two new species within the genus Kroppenstedtia, for
- 41 which the names *Kroppenstedtia pulmonis* sp. nov. (type strain W9323<sup>T</sup> = DSM 45752<sup>T</sup> = CCUG 68107<sup>T</sup>) and
- 42 *Kroppenstedtia sanguinis* sp. nov. (type strain  $X0209^{T} = DSM 45749^{T} = CCUG 38657^{T}$ ) are proposed.

#### 44 Introduction

45

46 The family *Thermoactinomycetaceae* was established to accommodate new taxa (*Mechercharimyces* spp.) and the previously 47 described genera Thermoactinomyces, Laceyella, Thermoflavimicrobium, Seinonella and Planifilum (Matsuo et al. 2006). 48 The family now encompasses 19 genera, with representatives isolated from clinical specimens (e.g. Desmospora, 49 Marinithermofilum, Hazenella) or environmental sources (e.g. Mechercharimvces, Lihuaxuella, Geothermomicrobium and 50 *Risungbinella*) (Matsuo et al. 2006; Yassin et al. 2009; Yu et al. 2012; Buss et al. 2013; Zhou et al. 2014; Kim et al. 2015; 51 Zhang et al. 2015). Salient properties of the family Thermoactinomycetaceae include thermotolerant growth up to 60 °C, 52 Gram positivity, non-acid fastness and formation of single spores that may be sessile or on simple sporophores with the 53 structure and properties of endospores. The description of the family was emended by establishing the 16S rRNA gene 54 sequence signature nucleotides as required for suprageneric assignment (Yassin et al. 2009). Von Jan et al. (2011) further 55 emended the description of the family Thermoactinomycetaceae (to contain either LL-diaminopimelic acid or meso-56 diaminopimelic acid) when they described a new genus and species, Kroppenstedtia eburnea, that contained the isomer LL-57 diaminopimelic acid in its whole-cell peptidoglycan. 58 59 Kroppenstedtia species have been isolated from environmental and clinical sources. Although the type strain of K. eburnea 60 was isolated from an environmental source (plastic surface in a contract manufacturing company in Germany), Barker et al. 61 (2012) identified 14 strains of this species in clinical (blood, skin, peritoneal fluid, cerebral spinal fluid) samples in the 62 United States. Another species, Kroppenstedtia guangzhouensis, was isolated from soil in south China (Yang et al. 2013).

63 Three human clinical strains of *Kroppenstedtia* species were identified in a retrospective evaluation of 16S rRNA gene

64 sequences at the Special Bacteriology Reference Laboratory (SBRL). Genotypic and phenotypic data suggest that strain

65 W9323<sup>T</sup> represents a new species, for which we propose the name *Kroppenstedtia pulmonis* sp. nov., and that strains X0209<sup>T</sup>

and X0394 both represent another new species, for which we propose the name *Kroppenstedtia sanguinis* sp. nov.

67

69	Materials and methods	
70		
71	Bacterial strains	
72		
73	Strain X0209 <sup>T</sup> was obtained from the Culture Collection University of Göteborg (CCUG), Göteborg, Sweden as	
74	'Thermoactinomyces sanguinis' (CCUG 38657 <sup>T</sup> ). Two strains, W9323 <sup>T</sup> and X0394, were sent to the SBRL, Centers for	
75	Disease Control and Prevention for identification. K. eburnea DSM 45196 <sup>T</sup> was used as a phenotypic, chemotaxonomic and	
76	genetic control and K. guangzhouensis GD02 <sup>T</sup> and Melghirimyces algeriensis NariEX <sup>T</sup> were used as chemotaxonomic and	
77	genetic controls throughout this study. The GenBank accession numbers for the 16S rRNA gene sequences and the patients'	
78	data associated with these strains are given in Table 1.	
79		
80	Phenotypic analyses	
81		
82	Morphological, cultural, physiological and biochemical analyses	
83		
84	To examine morphologic features, strains were grown aerobically using brain heart infusion (BHI) broth, heart infusion agar	
85	(HIA; Becton, Dickinson and Co, Sparks, MD) supplemented with 5 % rabbit blood, HIA slants, trypticase soy agar (TSA)	
86	supplemented with 5 % sheep blood (Becton, Dickinson and Co) at 35 and 45 °C for 3 to 7 days and then examined for	
87	microscopic and macroscopic features. Gram and modified Kinyoun acid-fast staining were conducted as described	
88	previously (Berd 1973). Cultures were examined for optimal growth at 35, 45, 50 and 60 °C for 7 days on HIA slants with 5	
89	% rabbit blood. Optimal growth was determined by comparative observation of the amount of cell mass production at each	
90	temperature. All phenotypic studies were performed under optimal growth conditions (at 45 °C in air).	
91		
92	Decomposition tests for adenine, casein, esculin, hypoxanthine, tyrosine, urea and xanthine, utilization of 22 carbohydrates as	
93	sole source of carbon, utilization of acetamide and citrate, arylsulfatase production and nitrate reduction and growth in the	
94	presence of lysozyme were performed as described previously (Conville and Witebsky 2007; Conville et al. 2008; Weyant et	
95	al. 1996; Yassin et al. 1995). The decomposition of casein was compared with casein plus 0.5 % NaCl as described by von	
96	Jan et al. (2011).	

# 99 Antimicrobial susceptibility testing

101	Since no guidelines were available for the genus Kroppenstedtia, the MICs to 10 antimicrobial agents were determined	
102	following interpretative breakpoints as recommended for aerobic actinomycetes (Clinical and Laboratory Standards Institute	
103	2011) for amikacin, amoxicillin-clavulanate, ceftriaxone, ciprofloxacin, clarithromycin, imipenem, linezolid, minocycline,	
104	moxifloxacin and trimethoprim-sulfamethoxazole; the interpretive breakpoint for ampicillin used was that recommended for	
105	Enterobacteriaceae (Clinical and Laboratory Standards Institute 2015). The preparation of the inoculum suspension	
106	followed the guidelines as described previously (Clinical and Laboratory Standards Institute 2003).	
107		
108	Chemotaxonomic analyses	
109		
110	Assays of diaminopimelic acid stereoisomers and whole-cell sugars were performed by thin-layer chromatography using	
111	methods described previously (Lechevalier and Lechevalier 1970; Rhuland et al. 1955; Staneck and Roberts 1974).	
112	Isoprenoid quinones and polar lipids were extracted and purified and analyzed by the method of Minnikin et al. (1984).	
113	Analysis of isoprenoid quinones by HPLC was performed as described by Kroppenstedt (1982, 1985). Cellular fatty acids	
114	were prepared by the method of Klatte et al. (1994) and the fatty acid methyl esters were then separated as described by	
115	Sasser (1990) using the Microbial Identification System (MIDI, Inc., Sherlock version 6.1). Standardization of the	
116	physiological age of the study and reference strains was obtained by choosing the sector from a quadrant streak of culture	
117	plates.	
118		
119	Genetic analyses	
120		
121	16S rRNA gene sequence analysis	
122		
123	Purification of whole-cell DNA, amplification of near full-length 16S rRNA gene fragments, primers and nucleotides for	
124	PCR, purification of amplicons and DNA cycle sequencing were described previously (Lasker et al. 2011). Consensus 16S	
125	rRNA gene sequences were assembled and edited using Sequencher version 4.10.1 software. To identify related gene	
126	sequences in the GenBank database, consensus sequences were submitted to GenBank using BLASTn software	

127	(https://www.ncbi.nlm.nih.gov/blast/). A multiple sequence alignment was created using Clustal W (within Geneious 8.1.4),	
128	from which gaps and 5' and 3' ends were trimmed. The distance matrix was calculated using DNADIST (Kimura 2-correction	
129	parameter) (Felsenstein 1989). Phylogenetic trees were constructed using the neighbor-joining (NEIGHBOR), maximum	
130	likelihood (DNAML with global rearrangements) and maximum parsimony (DNAPARS with global rearrangements)	
131	methods available in the PHYLIP package (Felsenstein, 1989). Stability of groupings within the neighbor-joining tree was	
132	estimated by bootstrap analysis (1000 replications) using the programs SEQBOOT, DNADIST, NEIGHBOR and	
133	CONSENSE (Felsenstein 1989). The phenograms were visualized by using the program DRAWGRAM (Felsenstein 1989).	
134	The sequence of the type strain of Bacillus subtilis was used as the outgroup.	
135		
136	DNA-DNA hybridization and DNA G + C content	
137		
138	Cells for hybridization and G+C mol% determination were disrupted using a Constant Systems TS 0.75 KW (IUL	
139	Instruments, Germany). DNA in the crude lysates was purified on hydroxyapatite by chromatography as described by	
140	Cashion et al. (1977). DNA-DNA hybridization was carried out in duplicate as described by De Ley et al. (1970) as modified	
141	by Huss et al. (1983) using a model Cary 100 Bio UV/VIS-spectrophotometer equipped with a Peltier-thermostat 6 x 6	
142	multicell changer and a temperature controller with <i>in-situ</i> temperature control. DNA-DNA hybridization studies were	
143	between the clinical strains X0209 <sup>T</sup> and X0394 to confirm they were the same genomic species. DNA-DNA hybridization	
144	studies between X0209 <sup>T</sup> and W9323 <sup>T</sup> and their respective closest phylogenetically related neighbors were not conducted	
145	because of the low probability inferred from 16S rRNA gene similarities observed (Meier-Kolthoff et al. 2013).	
146		
147	DNA G+C content	
148		
149	The method of Mesbah et al. (1989) was performed to determine the G+C content of the novel type strains.	
150		
151	Results and discussion	
152	Strains W9323 <sup>T</sup> , X0209 <sup>T</sup> and X0394 were aerobic, mesophilic to thermophilic, Gram-positive bacteria but were not acid fast.	
153	Substrate hyphae were filamentous and branched, and could be seen as fringes around the colony margins; no aerial hyphae	
154	were observed. Rare elongated (paddle shaped) endospores on long, unbranched sporophores were seen on Gram-stained	
155	smears of W9323 <sup>T</sup> ; single globose endospores on unbranched sporophores were seen on Gram-stained smears of X0209 <sup>T</sup> and	

156	X0394. Colonies of all three strains showed beta hemolysis on TSA with 5 % sheep blood at 45 °C. Growth occurred at 35	
157	and 45 °C but not at 50 °C with optimum growth at 45 °C; the optimal growth at 45 °C was consistent with the type strain of	
158	K. eburnea but differed from the optimal growth at 50 °C of the K. guangzhouensis type strain. Pale yellow colonies on 7-day	
159	TSA with 5 % sheep blood had irregular margins with random surface ridges at 35 and 45 °C. Except for the production of	
160	paddle-shaped endospores on long, unbranched sporophores of strain W9323 <sup>T</sup> , the microscopic morphology was consistent	
161	with the type strains of <i>K. eburnea</i> and <i>K. guangzhouensis</i> . The macroscopic morphology of W9323 <sup>T</sup> , X0209 <sup>T</sup> and X0394	
162	was consistent with that described for the type strains of K. eburnea and K. guangzhouensis, however, none of the three	
163	strains produced aerial hyphae as reported by von Jan et al. (2011) and Yang et al (2013).	
164		
165	Table 2 gives the differential phenotypic, chemotaxonomic and genetic characteristics of the study strains and their closest	
166	phylogenetic relatives. Strains X0209 <sup>T</sup> and X0394 both were able to utilize D-mannitol; W9323 <sup>T</sup> was the only strain able to	
167	utilize D-glucose and sucrose. Strain X0209 <sup>T</sup> utilized cellobiose and salicin but the related strain X0394 did not.	
168		
169	Results of antimicrobial susceptibility testing showed strains W9323 <sup>T</sup> , X0209 <sup>T</sup> and X0394 were resistant (MIC, $\ge 8 \ \mu g/ml$ ) to	
170	clarithromycin but were susceptible to amikacin (MIC, $\leq 8$ ), amoxicillin-clavulanate (MIC, $\leq 8/4 \ \mu g/ml$ ), ampicillin (MIC, $\leq 8/4 \ $	
171	4 $\mu$ g/ml), ceftriaxone (MIC, $\leq$ 8 $\mu$ g/ml), ciprofloxacin (MIC, $\leq$ 1 $\mu$ g/ml), imipenem (MIC, $\leq$ 4 $\mu$ g/ml), linezolid (MIC, $\leq$ 8	
172	$\mu$ g/ml), minocycline (MIC, $\leq 1 \mu$ g/ml), moxifloxacin (MIC, $\leq 1 \mu$ g/ml) and trimethoprim-sulfamethoxazole (MIC, $\leq 2/38$ )	
173	$\mu$ g/ml). Except for resistance to ampicillin of <i>K. eburnea</i> JFMB-ATE <sup>T</sup> , susceptibility results of the three study strains were	
174	consistent with K. eburnea JFMB-ATE <sup>T</sup> . The antimicrobial susceptibility test results in our study were consistent with the	
175	MIC results of 14 clinical strains of K. eburnea reported by Barker et al. (2012); for example, all their strains were	
176	susceptible to all antimicrobial agents tested except for clarithromycin.	
177		

- 178 Whole-cell wall hydrolysates contained LL-diaminopimelic acid, ribose and traces of galactose (X0209<sup>T</sup> and X0394); *K*.
- 179 *eburnea* JFMB-ATE<sup>T</sup> contained ribose and glucose. The predominant menaquinones of W9323<sup>T</sup> were MK-7 (95 %) and MK-
- 180 8 (5 %); the predominant menaquinones of X0209<sup>T</sup> and X0394 were identified as MK-7 (97 %) and MK8 (3 %). Polar lipids
- 181 were phosphatidylglycerol (PG), diphosphatidylglycerol (DPG), phosphatidylethanolamine (PE),
- 182 phosphatidylmethylethanolamine (PME), two unknown phospholipids (PL) (4PL, W9323<sup>T</sup> and X0394; 5PL, X0209<sup>T</sup>) and
- 183 one unknown glycolipid (GL) (1GL, W9323<sup>T</sup>, X0209<sup>T</sup> and X0394). The fatty acid profile of the three strains consisted
- 184 predominantly of *iso*- $C_{15:0}$  (74-75 %) and *anteiso*- $C_{15:0}$  (11-13 %).

186	DNA-DNA hybridization studies between the clinical strains X0209 <sup>T</sup> and X0394 confirmed they were the same genomic	
187	species with relatedness value of 93.6% $\pm$ 6.5%. The genomic DNA G+C content of strains W9323 <sup>T</sup> and X0209 <sup>T</sup> was 45.9	
188	and 50.6 mol%, respectively. These values fall within the range of genomic DNA G+C content reported for the genus	
189	Kroppenstedtia as it exists presently (46 to 56 mol%) (von Jan et al. 2011, Yang et al 2013).	
190		
191	Based on 16S rRNA gene sequence analysis, the clinical strains were assigned within the subclade for the genus	
192	Kroppenstedtia, within the family Thermoactinomycetaceae (Fig. 1). The highest sequence similarity with strain W9323 <sup>T</sup>	
193	was to K. eburnea JFMB-ATE <sup>T</sup> (95.3 % sequence similarity), K. guangzhouensis GD02 <sup>T</sup> (94.7 % sequence similarity),	
194	Melghirimyces thermohalophilus Nari11A <sup>T</sup> (94.5 % sequence similarity), M. algeriensis DSM 45474 <sup>T</sup> (94.3 % similarity),	
195	and Desmospora activa DSM 45169 <sup>T</sup> (94.2 % similarity). The 16S rRNA gene sequences for strains X0209 <sup>T</sup> and X0394 we	
196	99.9 % identical to each other. Strain X0209 <sup>T</sup> showed the highest sequence similarity to K. eburnea JFMB-ATE <sup>T</sup> (96.4 %	
197	similarity), K. guangzhouensis GD02 <sup>T</sup> (96.0 % similarity), M. algeriensis DSM 45474 <sup>T</sup> (94.3 % similarity), M.	
198	thermohalophilus Nari11A <sup>T</sup> and D. activa DSM 45169 <sup>T</sup> (93.7 % similarity). The 16S rRNA gene sequences of strain X0209	
199	and <i>K. eburnea</i> JFMB-ATE <sup>T</sup> differed by 59 bp; the sequences of strain W9323 <sup>T</sup> and <i>K. eburnea</i> JFMB-ATE <sup>T</sup> differed by 66	
200	bp.	
201		
202	From the results of our phenotypic, chemotaxonomic and genetic studies, it is proposed that strains W9323 <sup>T</sup> and X0209 <sup>T</sup> be	
203	classified in the genus Kroppenstedtia as Kroppenstedtia pulmonis sp. nov. and Kroppenstedtia sanguinis sp. nov.,	
204	respectively.	
205		
206	Description of Kroppenstedtia pulmonis sp. nov.	
207	K. pulmonis (N. L. fem. adj. pul.mo'nis. L. n. pulmo-onis, lung; L. gen. n. pulmonis of a lung, isolated from a lung).	
208	Cells are Gram-positive, non-acid-fast and filamentous. Elongated spores (paddle shaped) are formed singly on sessile	
209	sporophores on substrate hyphae. Pale yellowish-gray colonies are wrinkled with random ridges in 3 to 7 days at 35 and 45	
210	°C on trypticase soy agar with 5 % sheep blood. Colonies are beta hemolytic but do not form aerial hyphae. Growth occurs at	

211 35 and 45 °C but not at 50 °C. D-Glucose and sucrose are utilized but i-adonitol, L-arabinose, cellobiose, citrate, dulcitol, i-

- 212 erythritol, D-fructose, D-galactose, glycerol, i-myo-inositol, lactose, maltose, D-mannitol, mannose, melibiose, raffinose, L-
- 213 rhamnose, salicin, D-sorbitol, trehalose and xylose are not utilized. Grows in the presence of lysozyme but has no

- 214arylsulfatase activity and does not reduce nitrate. Hydrolyses casein with 0.5 % NaCl but does not hydrolyse acetamide,215adenine, casein, hypoxanthine, tyrosine, urea or xanthine. Esculin hydrolysis is positive by browning but negative by UV216light absorption. Major fatty acids (>10 %) are *iso*-C<sub>15:0</sub> (75 %) and *anteiso*-C<sub>15:0</sub> (11 %). Whole-cell hydrolysates contain217LL-diaminopimelic acid and the sugar ribose. The predominant menaquinones of strain W9323<sup>T</sup> are MK-7 (95 %) and MK-8218(5 %). Polar lipids are phosphatidylglycerol (PG), diphosphatidylglycerol (DPG), phosphatidylethanolamine (PE),219phosphatidylmethylethanolamine (PME), one unknown phospholipid (4PL) and one unknown glycolipid (1GL). The type220strain (W9323<sup>T</sup> = DSM 45752<sup>T</sup> = CCUG 68107<sup>T</sup>) was isolated from lung biopsy of a 78-year-old male patient from New
- 221 York, USA. The G+C content of its genomic DNA is 45.9 mol%.
- 222

#### 223 Description of *Kroppenstedtia sanguinis* sp. nov.

224 K. sanguinis (san'gui.nis L. n. sanguis-inis, blood; L. gen. n. sanguinis, of blood).

225 Cells are Gram-positive, non-acid-fast and filamentous. Globose spores are formed singly on sessile sporophores on substrate 226 hyphae. Pale yellowish gray colonies are wrinkled with random ridges in 3 to 5 days at 35 and 45 °C on trypticase soy agar 227 with 5 % sheep blood. Colonies are beta hemolytic but do not form aerial hyphae. Growth occurs at 35 and 45 °C but not at 228 50 °C. Cellobiose, D-mannitol, mannose and salicin are utilized but i-adonitol, L-arabinose, citrate, dulcitol, i-erythritol, D-229 fructose, D-galactose, D-glucose, glycerol, i-myo-inositol, lactose, maltose, melibiose, raffinose, L-rhamnose, D-sorbitol, 230 sucrose, trehalose and D-xylose are not utilized. Grows in the presence of lysozyme but has no arylsulfatase activity and does 231 not reduce nitrate. Hydrolyses casein with 0.5 % NaCl but does not hydrolyse acetamide, adenine, casein, hypoxanthine, 232 tyrosine, urea or xanthine. Esculin hydrolysis is positive by browning but negative by UV light absorption. The predominant 233 menaquinone is MK-7 (95 %). Whole-cell hydrolysates contain LL-diaminopimelic acid and traces of the sugar galactose. 234 Major fatty acids (>10 %) are iso-C<sub>15:0</sub> (74 %) and anteiso-C<sub>15:0</sub> (12 %). Whole-cell hydrolysates contain LL-diaminopimelic 235 acid and traces of galactose and ribose. Polar lipids are phosphatidylglycerol (PG), diphosphatidylglycerol (DPG), 236 phosphatidylethanolamine (PE), phosphatidylmethylethanolamine (PME), one unknown phospholipid (5PL), and one 237 unknown glycolipid (1GL). The type strain ( $X0209^{T} = DSM 45749^{T} = CCUG 38657^{T}$ ) was isolated from the blood of a 59-238 year-old male from Gävle, Sweden. The G+C content of its genomic DNA is 50.6 mol%.

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355	Figure	Legend

- Figure 1. Neighbor joining tree showing the positions of *Kroppenstedtia sanguinis* sp. nov. and *Kroppenstedtia pulmonis* sp.
- 357 nov. within the family *Thermoactinomycetaceae*. The tree was constructed based on an analysis of ~1480 nt. Bootstrap values
- 358 shown at the nodes are expressed as a percentage of 1000 replications (neighbor joining); only values >70 % are shown. ML,
- 359 nodes common to the neighbor joining and maximum likelihood analyses; MP, nodes common to the neighbor joining and
- 360 maximum parsimony analyses (Felsenstein 1989). The sequence of the type strain *Bacillus subtilis* was used as the outgroup.
- 361

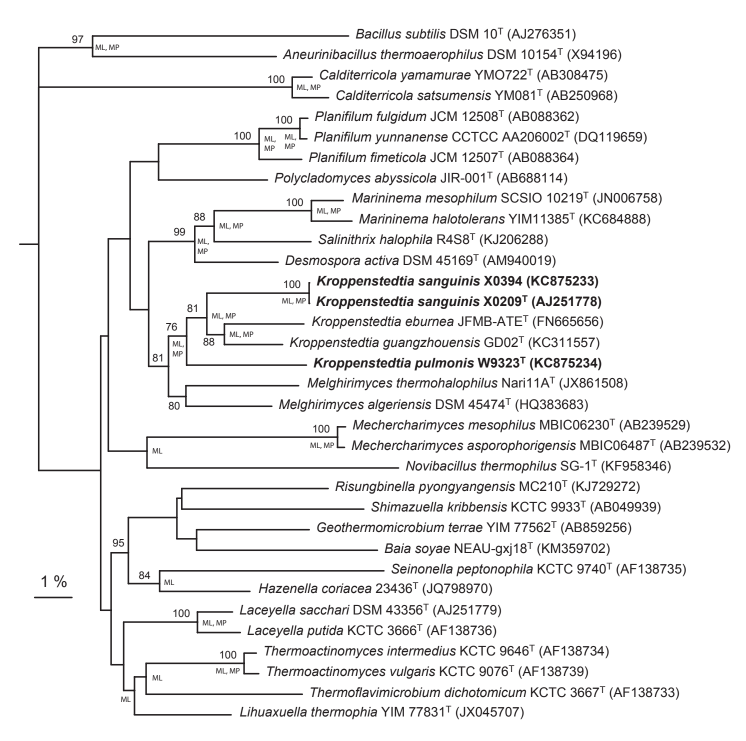


Table 1	Strains	used in	this	study
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Strain	Date received or reference	Source	Geographic source	GenBank accession number of 16S rRNA gene
Kroppenstedtia eburnea	von Jan et al.			
$(DSM 45196^{T})$	(2011)	Plastic surface	Germany	FN665656
K. guangzhouensis			-	
$(\text{GD02}^{\mathrm{T}})$	Yang et al. (2013)	Soil	China	KC311557
Melghirimyces algeriensis	Addou et al.			
$(NariEX^T)$	(2012)	Soil from salt lake	Algeria	HQ383683
		Lung, 78-year-old		
-		male		
W9323 <sup>T</sup>	2008		New York, USA	KC875234
		Blood, 59-year-old		
		male		
X0209 <sup>T</sup>	1997		Gävle, Sweden	AJ251778
		CSF, 16-year-old		
		female		
X0394	2010		Quebec, Canada	KC875233

Characteristic	K. pulmonis W9323 <sup>T</sup>	K. sanguinis X0209 <sup>T</sup>	K. sanguinis X0394	<i>K. eburnea</i> DSM 45196 <sup>T</sup>	K. guangzhouensis GD02 <sup>T a</sup>
Endospore morphology	Elongated paddle shaped	Globose	Globose	Globose	Globose
Aerial hyphae	-	-	-	- (+ <sup>b</sup> )	$NT (+^{c})$
Optimal growth on TSA	45 °C	45 °C	45 °C	45 °C	50 °C
sheep blood at 7 days					
Beta hemolysis on TSA sheep blood Utilization of:	+	+	+	+	NT
Cellobiose	-	+	-	+	-
Dulcitol	-	-	-	+	-
D-Fructose	-	-	-	+	-
D-Glucose	+	-	-	-	-
D-Mannitol	-	+	+	-	-
Mannose	-	+	-	-	NT
Salicin	-	+	-	+	NT
Sucrose	+	-	-	-	-
Whole cell-wall sugars	Ribose	Ribose	Ribose	Ribose	NT
	Glucose	Trace of galactose	Trace of galactose		
Cellular fatty acids	<i>iso</i> -C <sub>15</sub> (75.0 %)	<i>iso</i> -C <sub>15</sub> (74.0 %)	<i>iso</i> -C <sub>15</sub> (75.0 %)	<i>iso</i> -C <sub>15</sub> (73.0%)	<i>iso</i> -C <sub>15</sub> (63.5 %)
(>5%)	<i>anteiso</i> -C <sub>15</sub> (11.0 %) <i>iso</i> -C <sub>17</sub> (8.0 %)	<i>anteiso</i> -C <sub>15</sub> (12.0 %)	anteiso-C <sub>15</sub> (13.0 %)	<i>anteiso</i> -C <sub>15</sub> (13.0 %)	anteiso-C <sub>15</sub> (7.0 %) iso-C <sub>17</sub> (8 %)
	<i>iso</i> -C <sub>16</sub> (2.5 %)	<i>iso</i> -C <sub>17</sub> (7.0 %) <i>iso</i> -C <sub>16</sub> (1.4 %)	<i>iso</i> -C <sub>17</sub> (8.0 %) <i>iso</i> -C <sub>16</sub> (-)	<i>iso</i> -C <sub>17</sub> (-) <i>iso</i> -C <sub>16</sub> (4.5 %)	<i>iso</i> -C <sub>16</sub> (12.6 %)
Menaquinones	MK-7 (95.0 %) MK-8 (5.0 %)	MK-7 (97.0 %) MK-8 (3.0 %)	MK-7 (97.0 %) MK-8 (3.0 %)	MK-7 (97.0%) MK-8 (3.0%)	MK-7 (98.6 %) MK-8 (1.4 %)
Polar lipids	PG, DPG, PE, PME, 4 unknown PL,	PG, DPG, PE, PME,	PG, DPG, PE, PME, 4 unknown	PG, DPG, PE, PME, 2 unknown	PG, DPG, PE, PME, 2 unknown L
	1 unknown GL	5 unknown PL, 1 unknown GL	PL, 1 unknown GL	PL	
DNA $G + C$ mol%	45.9	50.5	50.6	54.6	56.3

All strains tested were negative for utilization of acetamide, adonitol, L-arabinose, citrate, i-erythritol, D-galactose, glycerol, i-*myo*inositol, lactose, maltose, melibiose, raffinose, L-rhamnose, D-sorbitol, trehalose and D-xylose; production of arylsulfatase and urease, hydrolysis of adenine, aesculin by fluorescence, hypoxanthine, tyrosine, xanthine and reduction of nitrate; all strains tested were positive for browning of aesculin, growth in lysozyme and hydrolysis of casein with 0.5 % NaCl. No growth on casein. PG, phosphatidylglycerol; DPG, diphosphatidylglycerol; PE, phosphatidylethanolamine; PME, phosphatidylmethylethanolamine; PL, phospholipid, GL, glycolipid; L, lipid; -, negative; +, positive; NT, not tested; TSA, trypticase soy agar. <sup>a</sup> All phenotypic data for *K. guangzhouensis* from Yang et al. (2013). All other data including cellular fatty acids, menaquinones, polar

lipids and DNA G+C content were generated from present study.

<sup>b</sup> Aerial hyphae reported by von Jan et al. (2011).

<sup>c</sup> Aerial hyphae reported by Yang et al. (2013).

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E-mail: jqv7@cdc.gov Date: January 27, 2016

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In accordance with the International Code of Nomenclature (the Bacteriological Code), the strain has been assigned a collection accession number; this number may not be subsequently altered.

The strain will be made available to the international scientific community upon the date of valid publication of the species/sub-species name(s) in the International Journal of Systematic and Evolutionary Microbiology. The depositor may request an earlier release.

Upon valid publication, the information on the strain will be transferred to the CCUG open database, on internet: http://www.ccug.se

# Edward Moore

Edward Moore, Curator - CCUG

CCUG 68107 T		Kroppenstedtia pulmonis sp. nov.		
HIS: OCC: AUTH: RESTR:	W9323 = D 16S rRNA	<- Bell M, CDC, Atlanta GA, USA SM 45752-T gene sequence (Accession Number: KC875234) ted distribution until validly published or 3 years		

Dr. Edward Moore, PhD; Professor of Bacteriology Tel: +46-31-342-4696; Fax: +46-31-82 96 17; E-mail: curator@ccug.se; URL: http://www.ccug.se CCUG: CULTURE COLLECTION UNIVERSITY OF GOTHENBURG Department of Clinical Bacteriology Sahlgrenska University Hospital Sahlgrenska Academy of the University of Gothenburg Box 7193, SE-402 34 Göteborg, Sweden

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# Edward Moore

Edward Moore, Curator - CCUG

CCUG	38657 T	Kroppenstedtia sanguinis sp. nov.
HIS: OCC: AUTH: RESTR:	X0209 = D 16S rRNA	<- PHL, Gävle SE SM 45749-T gene sequence (Accession Number: AJ251778) ted distribution until validly published or 3 years

Dr. Edward Moore, PhD; Professor of Bacteriology Tel: +46-31-342-4696; Fax: +46-31-82 96 17; E-mail: curator@ccug.se; URL: http://www.ccug.se Leibniz-Institut DSMZ-Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH



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Afasayar

Dr. E. Atasayar, Curator Actinomycetales

Geschäftsführer/ Managing Director: Prof. Dr. Jörg Overmann Aufsichtsratsvorsitzender/Head of Supervisory Board: RD Dr. David Schnieders Braunschweigische Landessparkasse (NORD/LB) Kto.-Nr./Account: 2 039 220 BLZ/Bank Code: 250 500 00 IBAN DE22 2505 0000 0002 0392 20 SWIFT (BIC) NOLADE 2 H Handelsregister/ Commercial Register: Amtsgericht Braunschweig HRB 2570 Steuer-Nr. 13/200/24030



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