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## Taxonomic Description template

# 1 Robertkochia solimangrovi sp. nov., isolated from

# <sup>2</sup> mangrove soil, and emended description of the genus

# **3** Robertkochia

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18			
19	Keywords: Robertkochia solimangrovi; polyphasic taxonomy; Flavobacteriaceae; mangrove		
20			
21	The full length 16S rRNA gene of strain CL23 <sup>T</sup> has been deposited at		
22	EMBL/DDBJ/GenBank with accession number MK258111.		
23	The whole genome shotgun project of strain $CL23^{T}$ and <i>R. marina</i> $CC$ -AMO-30D <sup>T</sup> are		
24	available at EMBL/DDBJ/GenBank under accession QKWN00000000 and QXMP00000000		
25	respectively.		
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# **ABSTRACT**

To date, there is sparse information for the genus Robertkochia with Robertkochia marina  $CC-AMO-30D^{T}$  as the only described member. We report here a new species isolated from mangrove soil of Malaysia Tanjung Piai National Park and perform polyphasic characterization to determine its taxonomy position. Strain CL23<sup>T</sup> is a Gram-negative, yellow-pigmented, strictly aerobic, catalase-positive and oxidase-positive bacterium. The optimal growth conditions were determined to be at pH 7.0, 30-37°C and 1-2% (w/v) NaCl. The major respiratory quinone was menaquinone-6 (MK-6) and the highly abundant polar lipids were four unidentified lipids, a phosphatidylethanolamine and two unidentified aminolipids. The 16S rRNA similarity between CL23<sup>T</sup> and *R. marina* CC-AMO-30D<sup>T</sup> is 96.67%. Strain CL23<sup>T</sup> and *R. marina* CC-AMO-30D<sup>T</sup> are clustered together and were distinguished from taxa of closely related genera in 16S rRNA phylogenetic analysis. Genome sequencing revealed the strain  $CL23^{T}$  has a genome size of 4.4 Mbp and a G+C content of 40.72 mol%. Overall genome indexes (OGRIs) including digital DNA-DNA hybridization (dDDH) value and average nucleotide identity (ANI) are 17.70% and approximately 70%, below the cut-off 70% and 95% respectively, indicated that strain CL23<sup>T</sup> is a distinct species to that of R. marina CC-AMO-30D<sup>T</sup>. Collectively, based on phenotypic, chemotaxonomic, phylogenetic and genomic evidence presented, strain CL23<sup>T</sup> is proposed as a new species with the name *Robertkochia solimangrovi* sp. nov. (=KCTC  $72252^{T}$  =LMG 31418<sup>T</sup>). An emended description of the genus *Robertkochia* is also proposed. 

Flavobacteriaceae is one of the widely spread bacterial families composed of 158 genera at 64 65 the time of writing [1]. The genus *Robertkochia* was introduced by Hameed et al. in 2014 [2] as one of the new genera in the family *Flavobacteriaceae*. Until now, the genus consisted of a 66 single species Robertkochia marina CC-AMO-30D<sup>T</sup>, which was isolated from surface 67 seawater at Taichung harbour, Taiwan [2]. The species was described as Gram negative, 68 69 strictly aerobic, orange-pigmented and with iso-C<sub>15:0</sub>, iso-C<sub>15:1</sub> G and iso-C<sub>17:0</sub> 3-OH as 70 predominant fatty acids. The report for Robertkochia is scarce as the previous study only focused on taxonomic assignment with one species reported so far [2]. Furthermore, the 71 genome of this genus and prospective application have not been studied or reported. 72

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74 Robertkochia and many other members of the Flavobacteriaceae are halophilic or 75 halotolerant bacteria that reside in diverse saline environments such as seawater, mangrove 76 forest and marine sediment [3-5]. Mangroves are inter-tidal wetlands that connect terrestrial 77 and marine ecosystems [6]. Due to periodic tidal flats, drastic changes in salinity and nutrient 78 availability of the mangrove environment make it a unique ecosystem [7]. Free living and symbiotic bacteria in such environment were found to play essential roles in maintaining 79 mangrove ecosystem such as recycling of organic matter and biotransformation of minerals 80 [8-10]. It was estimated that less than 5% of species in mangrove environment have been 81 described so far [11]. Therefore, it could be considered as one of the interesting areas to be 82 explored. In the present study, strain CL23<sup>T</sup> was isolated from soil obtained from mangrove 83 forest located at Tanjung Piai National Park, Johor, Malaysia. This strain was characterized 84 using polyphasic approach (phenotypic, chemotaxonomic and genomic aspects) following the 85 recommended guidelines [12, 13] and new criteria for classification [14] to elucidate its 86 taxonomy position. The results indicated that strain CL23<sup>T</sup> represents a new species within 87 Robertkochia genus, with the name Robertkochia solimangrovi sp. nov. is proposed. 88

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## 99 ISOLATION AND HOME HABITAT

Soil from the mangrove forest was sampled at Tanjung Piai National Park (GPS location: 100 101 1°16'06.0" N, 103°30'31.2" E) in September 2017 with permit (CJB F No. 734342) granted by Johor National Parks Corporation. The soil samples were serially diluted with sterile 102 distilled water (10<sup>-1</sup> to 10<sup>-8</sup>). A 0.1 ml of diluted sample was spread onto marine agar 2216 103 (MA; BD Difco) and incubated at 30-35°C for 1 to 14 days. A yellow-pigmented strain 104 designated as CL23<sup>T</sup> was isolated from MA and re-streaked twice to obtain a pure culture. 105 The strain was maintained in marine broth 2216 (MB; BD Difco) with 20 % (v/v) glycerol at 106 -80°C. Strain CL23<sup>T</sup> was deposited at Korean Collection for Type Cultures (KCTC) and 107 Belgian Co-ordinated Collections of Micro-organisms (BCCM) under accession of KCTC 108 72252<sup>T</sup> and LMG 31418<sup>T</sup>, respectively. For comparative polyphasic taxonomy 109 characterization, R. marina CC-AMO-30D<sup>T</sup> (=JCM 18552<sup>T</sup>) was obtained from Japan 110 Collection of Microorganisms (JCM). Both strains were routinely cultured on MA and in MB 111 112 at 30°C for 48 h, unless specified otherwise.

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# 116 **16S rRNA PHYLOGENY**

Genomic DNA was extracted using DNeasy Blood and Tissue kit (Qiagen) and was purified 117 by DNA Clean and Concentrator<sup>™</sup>-25 (Zymo Research) following manual instructions. The 118 16S rRNA gene of strain CL23<sup>T</sup> was amplified by PCR using universal primers: 27F (5'-119 AGAGTTTGATCMTGGCTCAG-'3) and 1525R (5'-AAGGAGGTGWTCCARCC-3') [15]. 120 The 16S rRNA gene was sequenced at Apical Scientific Pte. Ltd., Seri Kembangan, Malaysia. 121 After the sequencing, the raw sequences were trimmed, and the sequences were aligned using 122 ClustalW. The nearly full-length 16S rRNA gene was searched against EzBioCloud database 123 for identification. The amplified 16S rRNA gene of strain CL23<sup>T</sup> was also cross-checked 124 with the genome data to ensure the acquisition of full-length gene (1522 bp). The 16S rRNA 125 gene of strain CL23<sup>T</sup> (MK258111) shared highest similarity (96.67%) with R. marina CC-126 AMO-30D<sup>T</sup> (JX235674), which is below the accepted threshold of 98.7% for species 127 delineation [14]. The 16S rRNA gene similarity was less than 94% between strain CL23<sup>T</sup> and 128

other members of closely related genera: *Joostella marina* En5<sup>T</sup> (93.82%), *Joostella atrarenae* M1-2<sup>T</sup> (93.82%), *Zhouia spongiae* HN-Y44<sup>T</sup> (93.75%) and *Pustulibacterium marinum* E403<sup>T</sup> (93.35%).

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Phylogenetic trees of 16S rRNA were built following the Neighbor-joining (NJ) [16] and 133 Maximum Likelihood (ML) [17] algorithms using MEGA 7.0 software [18] based on 1000 134 bootstrap replications [19] and Kimura-2 parameter. Following the 16S rRNA phylogenetic 135 analysis (Fig. 1), strain CL23<sup>T</sup> and *R. marina* CC-AMO-30D<sup>T</sup> formed a clade in NJ and ML 136 trees, confirming the placement of strain CL23<sup>T</sup> within *Robertkochia* genus. The high 137 bootstrap value at the node separating the branch of strain CL23<sup>T</sup> and *R. marina* CC-AMO-138 30D<sup>T</sup> in 16S rRNA phylogenetic tree supported that these two strains are distinct between 139 each other. 140

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# 144 PHENOTYPIC AND CHEMOTAXONOMIC CHARACTERIZATION

Colony morphology was observed on MA at 30°C after 48 h of incubation. The Gram 145 staining was performed according to the protocol as described previously [20]. The malachite 146 green staining was used to assess the presence of endospore in 7-day old cultures [21]. The 147 Gram stain reaction and endospore formation were examined under light microscope (Nikon 148 ECLIPSE E200). Cell morphology was examined under scanning electron microscope (SEM; 149 150 JEOL JSM-IT300LV). The bacterial motility was investigated by hanging-drop approach [22]. The presence of flexirubin-type pigment was determined by flooding the cells with 20 % 151 152 (w/v) KOH [12].

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154 Catalase activity was detected by effervescence using 3 % (v/v) H<sub>2</sub>O<sub>2</sub> while oxidase activity 155 was determined by oxidation of tetramethyl-*p*-phenylenediamine. Hydrolysis of starch, casein, 156 L-tyrosine, hypoxanthine, xanthine, Tween 20, Tween 40, Tween 60, Tween 80, 157 carboxymethyl-cellulose (CMC) and xylan were tested according to Smibert and Krieg [21]. 158 Bile esculin hydrolysis was investigated using the method of Facklam and Moody [23]. Other 159 biochemical characteristics were revealed by API 20 E and API 20 NE kits (BioMérieux, 160 France). Carbohydrate utilization and enzyme activity profile of both strains were investigated by API 50 CHB and API ZYM kits (BioMérieux, France), respectively. All API
kits were carried out by following the manufacturer's instructions with slight modification in
which inoculation was supplemented up to 2 % (w/v) NaCl.

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Growth under anaerobic condition was tested by incubating the bacteria on MA for 14 days at 165 166 30°C using AnaeroGen (Oxoid) in an anaerobic jar (Mitsubishi Gas Chemical). Growth was tested on the following media: Reasoner's 2A agar (R2A; HiMedia), Nutrient agar (NA; 167 Merck), Tryptic soy agar (TSA; Merck), Luria-Bertani agar (LBA; Conda) and Muller Hinton 168 169 agar (MHA; Sigma) supplemented with 2 % (w/v) NaCl at 30°C for 7 days. The temperature range (4, 9, 15, 20, 25, 30, 37, 40, 42, 45 and 50°C) and the optimum temperature for growth 170 were determined using MB at pH 7. The pH range (in intervals of 1.0 pH unit) and optimum 171 pH for growth were investigated using MB at 30°C. The pH was adjusted with the following 172 buffer systems: 50 mM citrate phosphate (pH 4–5), 50 mM sodium phosphate (pH 6–8) and 173 50 mM glycine–NaOH (pH 9–10) [24]. The pH was verified after autoclaving. To test NaCl 174 tolerance and optimal concentration, the bacteria were grown in a medium containing yeast 175 extract (1.0 g l<sup>-1</sup>), peptone (5.0 g l<sup>-1</sup>), MgCl<sub>2</sub> (5.0 g l<sup>-1</sup>), MgSO<sub>4</sub>·7H<sub>2</sub>O (2.0 g l<sup>-1</sup>), CaCl<sub>2</sub> (0.5 g 176 1<sup>-1</sup>), KCl (1.0 g 1<sup>-1</sup>) and NaCl (0, 0.5, 1–11 %, w/v) [10]. 177

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Antibiotic susceptibility of bacteria against 21 antibiotics was tested using the disk diffusion method on MA at 30°C for 48 h [25]. The antibiotics disks (Oxoid) used were: ampicillin (10  $\mu$ g), bacitracin (10 IU), carbenicillin (100  $\mu$ g), chloramphenicol (100  $\mu$ g), clindamycin (2  $\mu$ g), doxycycline (30  $\mu$ g), erythromycin (60  $\mu$ g), gentamicin (10  $\mu$ g), kanamycin (50  $\mu$ g), lincomycin (2  $\mu$ g), minocycline (30  $\mu$ g), neomycin (30  $\mu$ g), novobiocin (5  $\mu$ g), oleandomycin (15  $\mu$ g), oxacillin (1  $\mu$ g), penicillin G (10 IU), piperacillin (100  $\mu$ g), polymyxin B (300 IU), rifampicin (5  $\mu$ g), streptomycin (10  $\mu$ g) and tetracycline (30  $\mu$ g).

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Strain CL23<sup>T</sup> was determined as a Gram negative, rod-shaped, non-spore forming, oxidase 187 188 positive and catalase positive bacterium with motile ability by gliding. The colony was in a circular form with 0.5–1.0 mm diameter, smooth surface, convex elevation, entire margin and 189 has translucent property on MA after 48 h incubation. Under SEM, cells of strain CL23<sup>T</sup> were 190 0.2-0.4 µm in width and 2.3-3.2 µm in length. The notable distinctive features to 191 differentiate strain CL23<sup>T</sup> and *R. marina* CC-AMO-30D<sup>T</sup> are shown in Table 1. In terms of 192 morphology, strain CL23<sup>T</sup> is yellow pigmented while *R. marina* CC-AMO-30D<sup>T</sup> was found 193 to be orange pigmented. Strain CL23<sup>T</sup> grew well in 15–42°C, pH 5–9 and 0–9 % (w/v) NaCl, 194

and in general strain CL23<sup>T</sup> demonstrated a broader growth range compared to *R. marina* 195 CC-AMO-30D<sup>T</sup> (Table 1). The optimal growth conditions of strain CL23<sup>T</sup> were observed at 196 30–37°C, pH 7 and 1–2 % (w/v) NaCl. Strain CL23<sup>T</sup> was also able to produce acetoin,  $\beta$ -197 galactosidase and weakly positive toward amygdaline according to API 20 E but not for R. 198 marina CC-AMO-30D<sup>T</sup>. Based on API ZYM, strain  $CL23^{T}$  was able to produce  $\alpha$ -199 galactosidase,  $\beta$ -galactosidase and  $\alpha$ -mannosidase, which were absent in *R. marina* CC-200 201 AMO-30D<sup>T</sup>. Both strains were further distinguished by the hydrolysis capability of gelatin, Tween 20, Tween 40, Tween 60, and exhibiting resistance towards ampicillin, penicillin G, 202 203 piperacillin and bacitracin (Table 1).

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For the chemotaxonomic analysis, cellular fatty acids were extracted following the protocol 205 of Microbial Identification System (MIDI, version 6.1) [26]. Biomass of strain CL23<sup>T</sup> and its 206 reference strain *R. marina* CC-AMO-30D<sup>T</sup> were harvested from MA after 48 h of incubation 207 at 30°C. The cells were saponified with methanolic base, then the resulting sodium salts of 208 fatty acids were methylated. In the final step, methyl esters were transferred to the organic 209 phase and washed. Fatty acid methyl esters were analyzed on an Agilent 6890 equipped with 210 Ultra-2 capillary column and subsequently identified in the RTSBA6 library. As exhibited in 211 Table 2, the predominant cellular fatty acid of strain CL23<sup>T</sup> and *R. marina* CC-AMO-30D<sup>T</sup> 212 were found to be iso- $C_{15:0}$ , iso- $C_{15:1}$  G and iso- $C_{17:0}$  3-OH (> 10%). Nonetheless, some fatty 213 acid patterns and abundance of strain CL23<sup>T</sup> varied when compared to *R. marina* CC-AMO-214  $30D^{T}$ , such as summed features 3 (3.64%) and 9 (5.24%) were constituted in strain CL23<sup>T</sup> but 215 none for *R. marina* CC-AMO-30D<sup>T</sup>. On top of that, the amount of iso-C<sub>16:0</sub>, anteiso-C<sub>15:0</sub> and 216 iso-C<sub>16:0</sub> 3-OH of strain CL23<sup>T</sup> are remarkably lower than *R. marina* CC-AMO-30D<sup>T</sup> (Table 217 218 2).

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The polar lipids and respiratory quinone analyses of strain CL23<sup>T</sup> were performed by Dr. 220 Brian Tindall at the Identification Service, DSMZ, Braunschweig, Germany. In brief, the 221 respiratory quinones were extracted by solvent methanol: hexane (2:1 v/v), separated by TLC 222 and High Performance Liquid Chromatography (HPLC) following the standard method by 223 Tindall [27]. The polar lipids were extracted using chloroform: methanol solvent and 224 separated by two-dimensional silica gel thin layer chromatography (TLC) [28]. Total lipid 225 material was identified using molybdatophosphoric acid and specific functional groups were 226 determined using spray reagents specific for defined functional groups. 227

The major respiratory quinone of strain CL23<sup>T</sup> was identified to be menaquinone-6 (MK-6), 229 which matched to R. marina [2] and other members in Flavobacteriaceae family [12]. In 230 terms of polar lipids, strain CL23<sup>T</sup> has four unidentified lipids (L1, L2, L3 and L4), a 231 phosphatidylethanolamine (PE) and two unidentified aminolipids (AL1 and AL2) as major 232 polar lipids (Fig. S1). Additionally, three unidentified glycolipids (GL1, GL2 and GL3) and 233 an unknown lipid (L5) were observed in minor amounts. The unidentified lipids (L1–L3) and 234 glycolipids (GL1-GL3) were not detected in *R. marina* CC-AMO-30D<sup>T</sup> [2]. Moreover, an 235 unidentified phospholipid (PL) was contained in R. marina CC-AMO-30D<sup>T</sup> in which this 236 lipid was not found in strain  $CL23^{T}$  [2]. 237

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# 241 GENOMIC CHARACTERIZATION

The genome of reference strain *R. marina* CC-AMO-30D<sup>T</sup> was not available at the time of study, therefore, both the genomes of strain  $CL23^{T}$  (NCBI accession: QKWN00000000) and *R. marina* CC-AMO-30D<sup>T</sup> (NCBI accession: QXMP00000000) were sequenced in this study. Whole genome sequencing of strain  $CL23^{T}$  was accomplished on an Illumina HiSeq 2500 platform (2 × 150 bp). The raw reads were filtered, and the quality data was *de novo* assembled using SOAPdenovo 2.04 [29]. The resulting genome was annotated using NCBI Prokaryotic Genome Annotation Pipeline (PGAP) [30].

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The assembled genome of strain CL23<sup>T</sup> consists of 23 contigs with 322× depth of sequencing 250 coverage (average), made up the size of genome with 4,407,290 bp in length and a GC 251 content of 40.72 mol%. The genome size of strain  $CL23^{T}$  is significantly larger than R. 252 marina CC-AMO-30D<sup>T</sup> (3,571,649 bp). The GC content of strain CL23<sup>T</sup> is slightly lower 253 than R. marina CC-AMO-30D<sup>T</sup> (43.67 mol%). Based on PGAP annotation, a total of 3669 254 protein coding genes was found in genome of strain CL23<sup>T</sup>. The genes responsible for 255 phosphatase activity were found in the genome of strain CL23<sup>T</sup> and *R. marina* CC-AMO-256  $30D^{T}$  with a total of 12 and 7 phosphatases were encoded respectively (Table S1). This 257 258 correlated to API ZYM results in which both strains were positive to acidic and alkali phosphatases. Notably, the number of phosphatases annotated is higher in strain CL23<sup>T</sup> as 259 compared to *R. marina* CC-AMO-30D<sup>T</sup>. On the other hand, strain CL23<sup>T</sup> consists of a series 260

of genes for assimilatory sulfate reduction into sulfite (sulfate adenylyltransferase subunit 261 CysN and CysD, adenylylsulfate kinase and phosphoadenylylsulfate reductase) and then 262 sulfite reduction into sulfide (FAD-binding oxidoreductase and LLM class flavin-dependent 263 oxidoreductase) (Table S1). Nevertheless, the genes responsible for reduction of sulfite to 264 sulfide are absent in *R. marina* CC-AMO-30D<sup>T</sup> (Table S1). Furthermore, strain CL23<sup>T</sup> also 265 encodes a set of genes for reduction of nitrate to ammonia (NirBD and NrfAH) in which 266 NirBD genes were not found in genome of R. marina CC-AMO-30D<sup>T</sup> (Table S1). These 267 genes suggest that strain CL23<sup>T</sup> participates in nutrient recycling in mangrove environments. 268 269

Multilocus sequence analysis (MLSA) was conducted on five housekeeping genes of strain 270 CL23<sup>T</sup>, *R. marina* CC-AMO-30D<sup>T</sup> and related genera, which the sequences were retrieved 271 from genome data. The sequences of housekeeping genes were aligned individually and then 272 concatenated in the following order: *rpoB–gyrB–recA–mutL–atpD*. The phylogenetic tree of 273 concatenated housekeeping genes was constructed using MEGA 7.0 similarly as described in 274 section "16S rRNA phylogeny". In this tree (Fig. 2), strain CL23<sup>T</sup> and *R. marina* CC-AMO-275 30D<sup>T</sup> are clustered together but well distinguished from each other with high level of support 276 (>90% bootstrap value). Likewise, the phylogenetic tree based on whole genome sequences 277 that built using REALPHY 1.12 [31] also supported that both strain CL23<sup>T</sup> and *R. marina* 278  $CC-AMO-30D^{T}$  are grouped in the same clade (Fig. S2). 279

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To further underpin the classification of strain CL23<sup>T</sup> as a new species, the overall genome related indexes (OGRIs) were determined. Average nucleotide identity based on BLAST (ANIb) was calculated using JSpeciesWS [32]. ANI based on USEARCH (OrthoANIu) was determined by ChunLab's online ANI calculator [33]. The digital DNA-DNA hybridization (dDDH) value was calculated by Genome-to-Genome Distance Calculator [34].

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The ANIb and OrthoANIu values between strain  $CL23^{T}$  and *R. marina* CC-AMO-30D<sup>T</sup> are 69.35% and 70.47% respectively. These ANI values are below the recommended 95–96% for species delineation [35]. Similarly, the dDDH value between two strains was found to be 17.70%, lower than 70%, the cut off for species boundary [34]. Combining the interpretation of ANI and dDDH values, the result revealed the identity of strain  $CL23^{T}$  as a distinct species within the same genus as *R. marina* CC-AMO-30D<sup>T</sup>.

Based on polyphasic taxonomy characterization including phenotypic, chemotaxonomic, phylogenetic and genomic aspects, the results clearly indicated that strain  $CL23^{T}$  (=KCTC 72252<sup>T</sup> =LMG 31418<sup>T</sup>) represents a new species within the genus *Robertkochia*, for which the name *Robertkochia solimangrovi* sp. nov. is proposed.

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# 301 DESCRIPTION OF ROBERTKOCHIA SOLIMANGROVI SP. NOV.

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# Robertkochia solimangrovi sp. nov. (so.li.man.gro'vi. L. neut. n. solum soil; N.L. neut. n. *mangrovum* a mangrove; N.L. gen. n. solimangrovi of soil of a mangrove, pertaining to where the type strain was isolated.)

306 The cells are Gram-negative, rod shape, approximate 0.2–0.4 µm in width and 2.3–3.2 µm in length with motile ability by gliding. Colony is yellow-pigmented, in circular form with 0.5– 307 308 1.0 mm diameter, smooth surface, convex elevation, entire margin and has translucent property after 48 hours incubation at 30°C on MA. Flexirubin-type pigment is absent. Cells 309 are positive for oxidase and catalase. Growth occurs at 15–42 °C (optimum, 30–37°C), pH 310 5–9 (optimum, pH 7) and in the presence of 0–9 % (w/v) NaCl (optimum, 1–2 % (w/v) NaCl). 311 Grows well on MA, however, no growth is observed on R2A, NA, LBA, TSA and MHA 312 media. No growth is observed on MA in anaerobic condition. The predominant fatty acids 313 are iso-C<sub>15:0</sub>, iso-C<sub>15:1</sub> G and iso-C<sub>17:0</sub> 3-OH. The major isoprenoid quinone is menaquinone-6 314 (MK-6). The major polar lipids are four unidentified lipids, a phosphatidylethanolamine and 315 two unidentified aminolipids. Xylan, esculin, Tween 20, 40 and 60 are hydrolyzed. L-316 Tyrosine is weakly hydrolyzed. Casein, starch, CMC, Tween 80, xanthine and hypoxanthine 317 are not hydrolyzed. In the API 20 E strip, positive for ONP-β-D-galactopyranoside and 318 acetoin production; weakly positive for fermentation/oxidation of amygdaline; negative for 319 320 arginine dihydrolase, lysine decarboxylase, ornithine decarboxylase, tryptophane deaminase, 321 urease and gelatinase, production of H<sub>2</sub>S and indole, utilization of citrate, fermentation/oxidation of D-glucose, D-mannitol, inositol, D-sorbitol, L-rhamnose, D-322 323 saccharose, D-melibiose and L-arabinose. In the API 20 NE strip, positive for hydrolysis of pNP-β-D-galactopyranoside and esculin ferric citrate; negative for nitrate reduction, indole 324

production, arginine dihydrolase, gelatinase and urease, fermentation of D-glucose and 325 assimilation of D-glucose, D-arabinose, D-mannose, D-mannitol, N-acetyl-glucosamine, D-326 maltose, potassium gluconate, capric acid, adipic acid, malic acid and phenylacetic acid. In 327 the API 50 CHB strip, acid is produced from D-galactose, D-glucose, D-mannose, esculin 328 ferric citrate, D-cellobiose, D-maltose, D-lactose, D-melibiose, D-saccharose, D-trehalose, D-329 melezitose, p-raffinose, amidon, glycogen and gentibiose; acid is weakly produced from 330 methyl- $\alpha$ -D-glucopyranoside, arbutin, salicin and D-turanose; acid is not produced from 331 glycerol, erythritol, D-arabinose, L-arabinose, D-ribose, D-xylose, L-xylose, D-adonitol, 332 333 methyl-β-D-xylopyranoside, D-fructose, L-sorbose, L-rhamnose, dulcitol, inositol, D-mannitol, D-sorbitol, methyl-α-D-mannopyranoside, N-acetyl-glucosamine, amygdalin, inulin, xylitol, 334 D-lyxose, D-tagatose, D-fucose, L-fucose, D-arabitol, L-arabitol, potassium gluconate, 335 potassium 2-ketogluconate and potassium 5-ketogluconate. In the API ZYM strip, alkali 336 phosphatase, esterase (C4), esterase lipase (C8), leucine arylamidase, valine arylamidase, 337 338 cvstine arylamidase, trypsin, chymotrypsin, acid phosphatase, naphtol-AS-BIphosphohydrolase,  $\alpha$ -galactosidase,  $\beta$ -galactosidase,  $\alpha$ -glucosidase,  $\beta$ -glucosidase, *N*-acetyl-339  $\beta$ -glucosaminidase and  $\alpha$ -mannosidase are present; weak positive reaction for  $\alpha$ -fucosidase 340 and negative results for lipase (C14) and  $\alpha$ -fucosidase. Cells are susceptible to carbenicillin, 341 342 clindamycin, doxycycline, lincomycin, minocycline, novobiocin, oleandomycin, rifampicin and tetracycline, but not to ampicillin, bacitracin, chloramphenicol, erythromycin, gentamicin, 343 344 kanamycin, neomycin, oxacillin, penicillin G, piperacillin, polymyxin B and streptomycin.

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The type strain is  $CL23^{T}$  (=KCTC 72252<sup>T</sup> =LMG 31418<sup>T</sup>), isolated from soil of mangrove collected from Tanjung Piai National Park, Johor, Malaysia.

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The EMBL/DDBJ/GenBank accession for 16S rRNA gene of strain CL23<sup>T</sup> is MK258111. Genome metrics are as follows: genome size, 4,407,290 bp; number of contigs, 23; G+C content, 40.72 mol%. The Whole Genome Shotgun project of strain CL23<sup>T</sup> is available at EMBL/DDBJ/GenBank under accession QKWN00000000. The version described in this paper is QKWN01000000.

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# 355 EMENDED DESCRIPTION OF GENUS ROBERTKOCHIA

The characteristics of the genus *Robertkochia* are described according to Hameed et al. 2014 [2] with following amendments and additional information. Oxidase is either positive or negative and catalase is positive. The DNA G+C content of the type strain of type species is 43.67 mol% based on genome data. The Whole Genome Shotgun project of type strain of type species is available at EMBL/DDBJ/GenBank under accession QXMP00000000. The version described in this paper is QXMP01000000.

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# 365 AUTHOR STATEMENTS

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387 Ethical statement

388 No human and animal experiments are involved.

390 Conflicts of interest

391 All authors declared that there are no conflicts of interest.

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### 393

# 394 **ABBREVIATIONS**

ANI, average nucleotide identity, dDDH, digital DNA-DNA hybridization; MA, marine agar;
MB, marine broth; MK, menaquinone; ML, maximum likelihood; NJ, neighbor-joining;
OGRI, overall genome related index; PE, phosphatidylethanolamine.

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# 522 FIGURES AND TABLES

523

**Table 1.** Differential phenotypic characteristics of strain CL23<sup>T</sup> and *Robertkochia marina* 

525 CC-AMO- $30D^{T}$ .

Strains: 1, strain CL23<sup>T</sup>; 2, *R. marina* CC-AMO-30D<sup>T</sup>. All data were obtained from this study. 526 +, Positive reaction; -, negative reaction; w, weakly positive reaction. All strains were 527 positive for catalase; hydrolysis of xylan and aesculin; production of acid from D-glucose, 528 esculin ferric citrate, D-cellobiose, D-maltose, D-saccharose, D-trehalose, D-melezitose, 529 amidon and glycogen in API 50 CHB strips; and activity of alkali phosphatase, esterase (C4), 530 esterase lipase (C8), leucine arylamidase, valine arylamidase, cystine arylamidase, trypsin, 531 chymotrypsin, acid phosphatase, naphtol-AS-BI-phosphohydrolase,  $\alpha$ -glucosidase,  $\beta$ -532 glucosidase and *N*-acetyl-β-glucosaminidase. Both strains were negative for flexirubin-type 533 pigment; growth under anaerobic condition; growth on R2A, NA, LBA, TSA and MHA 534 media; hydrolysis of casein, starch, CMC, Tween 80, xanthine and hypoxanthine; nitrate 535 reduction; indole and H<sub>2</sub>S production; urease; acid production from glycerol, erythritol, D-536 arabinose, L-arabinose, D-ribose, D-xylose, L-xylose, D-adonitol, methyl-B-D-xylopyranoside, 537 D-fructose, L-sorbose, L-rhamnose, dulcitol, inositol, D-mannitol, D-sorbitol, methyl-a-D-538 mannopyranoside, N-acetyl-glucosamine, amygdalin, inulin, xylitol, p-lyxose, p-tagatose, p-539 fucose, L-fucose, D-arabitol, L-arabitol, potassium gluconate and potassium 2-ketogluconate 540 and potassium 5-ketogluconate in API 50 CHB strips; and activity of lipase (C14) and β-541 glucuronidase (API ZYM). 542

Characteristics	1	2
Colony pigmentation	Yellow	Orange
Oxidase	+	—
Growth parameters		
pH range	5–9	6–7
Temperature range (°C)	15–42	20–40
Temperature optimum (°C)	30–37	30
NaCl range (%, w/v)	0–9	0.5–4
NaCl optimum (%, w/v)	1–2	2
Hydrolysis of		
Tween 20	+	W
Tween 40	+	_
Tween 60	+	W
Tyrosine	W	—

Gelatin	—	+
Production of		
Acetoin	+	_
Oxidation of		
Amyldaline	W	_
Utilization of		
D-Galactose	+	_
D-Mannose	+	—
Arbutin	W	_
Salicin	W	_
D-Lactose	+	_
D-Melibiose	+	_
D-Raffinose	+	_
Gentiobiose	+	_
D-Turanose	W	—
Enzyme activity (API ZYM)		
α-Galactosidase	+	_
β-Galactosidase	+	—
α-Mannosidase	+	W
α-Fucosidase	W	_
Antibiotic susceptibility (per disc)		
Ampicillin (10 µg)	_	+
Penicillin G (10 IU)	_	+
Piperacillin (100 µg)	_	+
Bacitracin (45 µg)	_	+

**Table 2.** Cellular fatty acid profiles (%) of strain CL23<sup>T</sup> and *Robertkochia marina* CC-AMO-

559 30D<sup>T</sup>.

560 Strains: 1, strain CL23<sup>T</sup>; 2, *R. marina* CC-AMO-30D<sup>T</sup>. All data presented in the table are

from this study. TR, trace ( $\leq 0.5\%$ ); -, not detected. Major components (> 10%) are

562 highlighted in bold.

Fatty acid	1	2	
Branched saturated	Branched saturated		
iso-C <sub>13:0</sub>	TR	2.4	
iso-C <sub>14:0</sub>	_	2.4	
iso-C <sub>15:0</sub>	21.8	19.9	
iso-C <sub>16:0</sub>	3.4	6.1	
anteiso-C <sub>15:0</sub>	2.3	5.8	
Unsaturated			
C <sub>15:1</sub> ω5c	0.7	_	
C <sub>17:1</sub> $\omega$ 6c	1.7	_	
C <sub>17:1</sub> $\omega$ 8c	0.8	—	
Branched unsaturated			
iso-C <sub>15:1</sub> G	10.8	23.3	
iso-C <sub>16:1</sub> G	_	1.6	
iso-C <sub>16:1</sub> H	1.0	—	
anteiso-C <sub>15:1</sub> A	TR	2.8	
Hydroxy			
C <sub>15:0</sub> 2-OH	0.9	1.5	
С15:0 З-ОН	2.0	0.6	
C <sub>16:0</sub> 3-OH	1.4	TR	
С <sub>17:0</sub> 3-ОН	1.1	TR	
iso-C <sub>16:0</sub> 3-OH	2.6	6.5	
iso-C <sub>17:0</sub> 3-OH	29.5	15.5	
Summed features *			
3 *	3.6	—	
9 #	5.2	_	

<sup>\*</sup> Summed features are groups of two or three fatty acids that cannot be separated by GLC

s64 with the MIDI system.

<sup>+</sup>Summed feature 3 consisted of iso- $C_{15:0}$  2-OH,  $C_{16:1}\omega$ 6c and/or  $C_{16:1}\omega$ 7c and annotated here as

566 iso- $C_{15:0}$  2-OH based on the equivalent chain length (ECL).

567 <sup>#</sup> Summed feature 9 consisted of iso- $C_{17:1}\omega$ 9c and/or  $C_{16:0}$  10-methyl.

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### **Figure legends**





Fig. 1. Neighbor joining 16S rRNA phylogenetic tree manifesting the relationship of strain CL23<sup>T</sup> with closely related taxa of family *Flavobacteriaceae*. Corresponding Genbank accession numbers are indicated in parentheses. Bootstrap values  $\geq 70\%$  based on 1000 resampled datasets are depicted as percentages at nodes. Bootstrap value from left to right for NJ and ML calculated with same sequence set. Filled circles indicate that corresponding nodes were also recovered in dendrograms generated using ML algorithm. The sequence of Cytophaga hutchinsonii ATCC 33406<sup>T</sup> was used as outgroup. Bar, 0.05 substitutions per nucleotide position. 





Fig. 2. Neighbor joining phylogenetic tree based on the concatenated sequences of five 590 housekeeping genes: rpoB-gyrB-recA-mutL-atpD, indicating the position of strain CL23<sup>T</sup>. 591 Bootstrap values ≥70% based on 1000 resampled datasets are depicted as percentages at 592 nodes; value <70% is indicated by a dash. Bootstrap value from left to right for NJ and ML 593 calculated with same sequence set. Filled circles indicate that corresponding nodes were also 594 recovered in dendrograms generated using ML algorithm. The sequence of Cytophaga 595 hutchinsonii ATCC 33406<sup>T</sup> was used as outgroup. Bar, 0.05 substitutions per nucleotide 596 position. 597 598

# **Supplementary Materials**

*Robertkochia solimangrovi* sp. nov., isolated from mangrove soil, and emended description of the genus *Robertkochia* 

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**Fig. S1.** Polar lipids profile of strain CL23<sup>T</sup>. Unidentified lipids; L1–L5, phosphatidylethanolamine; PE, unidentified aminolipids; AL1–AL2, unidentified glycolipids; GL1–GL3.



**Fig. S2.** Neighbor joining phylogenomic tree manifesting the relationship of strain  $CL23^{T}$  with closely related taxa of family *Flavobacteriaceae*. Corresponding Genbank accession numbers are indicated in parentheses. Bootstrap values  $\geq$ 50% based on 1000 resampled datasets are depicted as percentages at nodes. Bootstrap value from left to right for NJ and ML calculated with same sequence set. Filled circles indicate that corresponding nodes were also recovered in dendrograms generated using ML algorithm. Bar, 0.005 substitutions per nucleotide position.

# Supplementary tables

**Table S1.** List of potential genes for phosphatases, sulfur reduction and nitrate reduction encoded in the genome of strain  $CL23^{T}$  and *R. marina* CC-AMO-30D<sup>T</sup>.

Category	Bacterial strain	NCBI Annotation	Accession
Phosphatases	CL23 <sup>T</sup>	alkaline phosphatase family protein	TRZ44267
		alkaline phosphatase	TRZ44500
		alkaline phosphatase	TRZ44343
		pyrophosphatase	TRZ44378
		sodium-translocating pyrophosphatase	TRZ44400
		alkaline phosphatase family protein	TRZ43533
		alkaline phosphatase family protein	TRZ43596
		alkaline phosphatase	TRZ42861
		HAD family phosphatase	TRZ42760
		alkaline phosphatase family protein	TRZ42969
		alkaline phosphatase family protein	TRZ41972
		HAD family phosphatase	TRZ41063
	R. marina CC-	alkaline phosphatase family protein	TRZ46762
	$AMO-30D^{T}$	alkaline phosphatase family protein	TRZ45488
		alkaline phosphatase family protein	TRZ45685
		sodium-translocating pyrophosphatase	TRZ44743
		HAD family phosphatase	TRZ42656
		pyrophosphatase	TRZ41149
		HAD family phosphatase	TRZ40862
Sulfur	CL23 <sup>T</sup>	sulfate adenylyltransferase subunit CysN	TRZ46029
reduction		sulfate adenylyltransferase subunit CysD	TRZ46030
		adenylyl-sulfate kinase	TRZ46031
		phosphoadenylylsulfate reductase	TRZ44200
		phosphoadenylylsulfate reductase	TRZ42776
		FAD-binding oxidoreductase	TRZ41175
		LLM class flavin-dependent oxidoreductase	TRZ41182
	R. marina CC-	sulfate adenylyltransferase subunit CysN	TRZ40960
	$AMO-30D^{T}$	sulfate adenylyltransferase subunit CysD	TRZ40970
		adenylyl-sulfate kinase	TRZ40959
		phosphoadenylylsulfate reductase	TRZ46694
Nitrate	CL23 <sup>T</sup>	nitrite reductase (NirBD)	TRZ44395
reduction		nitrite reductase (NAD(P)H) (NirBD)	TRZ42280

	nitrite reductase (NAD(P)H) small subunit	TRZ42281
	(NirBD)	
	NAD(P)H-nitrite reductase (NirBD)	TRZ42287
	ammonia-forming cytochrome c nitrite	TRZ42033
	reductase (NrfAH)	
	cytochrome c nitrite reductase small	TRZ42034
	subunit ( <i>NrfAH</i> )	
R. marina CC-	ammonia-forming cytochrome c nitrite	TRZ44150
$AMO-30D^{T}$	reductase (NrfAH)	
	cytochrome c nitrite reductase small	TRZ44178
	subunit ( <i>NrfAH</i> )	