

1 ***Pseudomonas daroniae* sp. nov. and *Pseudomonas dryadis* sp. nov. isolated from**
2 **pedunculate oak affected by Acute Oak Decline in the United Kingdom.**

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15
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17
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19 Plant Pathogenic Bacteria, (United Kingdom).

20
21 The GenBank/EMBL/DDBJ accession numbers are as follows: MK159357 - MK159378 (16S
22 rRNA), MK293898 - MK293919 (*gyrB*), MK293920 - MK293941 (*rpoB*), MK293876-
23 MK293897 (*rpoD*) and QJUH00000000 - QJUN00000000 (whole genome).

35

36 **ABSTRACT**

37 Twenty-two cream-coloured bacterial strains were isolated from oak trees affected by acute
38 oak decline (AOD) in Southern England. Isolates were Gram-negative, motile, slightly curved
39 rods, aerobic, non-spore-forming, and catalase and oxidase positive. 16S rRNA gene sequence
40 analysis placed the strains in two separate phylogenetic clusters in the *Pseudomonas straminea*
41 group, with *Pseudomonas flavescens* as the closest phylogenetic relative. Multilocus sequence
42 analyses (MLSA) of the genes *gyrB*, *rpoD* and *rpoB* supported the delineation of the strains
43 into two separate taxa, which could be differentiated phenotypically and chemotaxonomically
44 from each other, and their closest relatives. Average nucleotide identity (ANI) and *in silico*
45 DNA-DNA hybridisation values revealed percentages of genome similarity below the species
46 threshold (95% and 70% respectively) between the strains and the closest relatives, confirming
47 their novel species status. Therefore, on the basis of this polyphasic approach we propose two
48 novel *Pseudomonas* species, *Pseudomonas daroniae* sp. nov. (type strain FRB 228^T = LMG
49 31087^T = NCPPB 4672^T) and *Pseudomonas dryadis* sp. nov. (type strain FRB 230^T = LMG
50 31087^T = NCPPB 4673^T).

51

52 Bacteria are consistently isolated from pedunculate oak trees (*Quercus robur*) affected by acute
53 oak decline (AOD) in the United Kingdom [1]. AOD has been described as a polymicrobial
54 syndrome that causes vertical cracks in the bark, from which dark fluid emanates. The bacterial
55 communities isolated from AOD-affected oaks at different sites in the UK show a significantly
56 similar composition pattern, where *Brenneria goodwinii* and *Gibbsiella quercinecans* appear
57 to be the predominant bacterial species [1]. In addition to these species, isolates identified as
58 belonging to the genus *Pseudomonas* by 16S rRNA gene and *gyrB* sequencing have been
59 consistently isolated from AOD-affected inner bark, and also in larval galleries created in the
60 phloem by the buprestid beetle *Agrilus biguttatus* [2].

61

62 Species belonging to the genus *Pseudomonas* [3] are ubiquitous microorganisms commonly
63 isolated from practically all environmental niches (such as water, soil and air), in addition to
64 those coexisting, or causing disease in humans, plants and animals. They are aerobic, Gram-
65 negative bacilli and motile by means of polar or peritrichous flagella. The genus *Pseudomonas*
66 is a heterogeneous taxon with an increasing number of species [4]. The current classification
67 approach for novel species of the genus *Pseudomonas* is polyphasic and includes genotypic,
68 phenotypic, and chemotaxonomic characterisation of bacterial isolates. In the present study,

69 16S rRNA gene sequence analysis was performed, along with multilocus sequence analysis
70 (MLSA) of the concatenated nucleotide sequences of the *gyrB*, *rpoD* and *rpoB* genes. The
71 phylogenetic analyses were supported by whole genome relatedness values, and PCR-based
72 fingerprinting methods were used to assess the genetic diversity of the bacterial isolates.
73 Moreover, hypersensitivity reactions against a plant host were investigated.

74

75 Based on the data obtained from this polyphasic study, the AOD-associated bacterial strains
76 constitute two novel species in the genus *Pseudomonas* for which we propose the names
77 *Pseudomonas daroniae* sp. nov. and *Pseudomonas dryadis* sp. nov.

78

79 Isolation

80 Bacterial strains were isolated from the inner bark and sapwood, as well as galleries created in
81 the phloem by *Agrilus biguttatus*. Oak trees were located in three sites in the United Kingdom:
82 in two deciduous woodlands: Bisham Woods (51.54911°N, 0.77071°W) and Great Monk
83 Wood (51.89955°N, 000.64606°E), and in the Royal Botanic Gardens, Kew. A list of strains
84 used in this study are presented in Suppl. Table S1. Strains were initially isolated in potato
85 yeast glucose agar, and routinely cultured at 25°C on King's B agar (KB) (Oxoid, UK) and in
86 Luria Bertrani (LB) broth.

87

88 Genotypic characterisation

89 Genomic DNA was extracted following an alkali lysis method [5]. Amplification and
90 sequencing of the 16S rRNA, *gyrB*, *rpoD* and *rpoB* genes for each of the 22 strains was as
91 previously described [6–8]. After alignment and trimming, the final sequence lengths were as
92 follows: 16S rRNA – 1399 bp, *gyrB* – 759 bp, *rpoD* – 636 bp, and *rpoB* – 750 bp. The 16S
93 rRNA gene sequences for each of the 22 strains were queried against the EZBioCloud 16S
94 rRNA database [9]. The pairwise similarities obtained for the strains were 98.73 – 99.23% to
95 *P. flavescens*, 98.66 - 99.23% to *P. seleniipraecipitans*, 98.31 - 98.87% to *P. punonensis* 98.1
96 - 98.73% to *P. straminea* and 97.96 - 98.66% to *P. argentinensis*. Values of greater than 95%
97 were observed to other members of the genus. For phylogenetic comparisons, the 16S rRNA
98 gene sequences of the 36 closest related strains were downloaded from EzBioCloud. Sequences
99 of *gyrB*, *rpoD* and *rpoB* from the closest phylogenetic neighbours, as determined by
100 EzBioCloud and BLAST [10], were downloaded from the GenBank database (Suppl. Table
101 S2) for analysis of the concatenated gene datasets. All sequences were aligned and trimmed in
102 BioEdit 7.2.5 [11] and *gyrB*, *rpoD* and *rpoB* gene datasets were concatenated in

103 SequenceMatrix [12]. Nucleotide sequences were translated to amino acid sequences with
104 BioEdit.

105

106 Maximum likelihood (ML) and Bayesian Inference (BI) trees were constructed in PhyML [13]
107 and MrBayes [14], respectively. The best-fit nucleotide substitution model – based on the
108 Akaike information criterion (AIC) – was selected by the software Smart Model Selection
109 (SMS) [15] in ML trees, and by JModelTest [16] in BI trees. The general time reversible model
110 with invariable sites and gamma distribution (GTR+I+G) was the best substitution model for
111 the 16S rRNA gene ML and BI phylogenetic trees, as well as for the concatenated (*gyrB*, *rpoD*
112 and *rpoB*) ML tree. The best-fit nucleotide substitution model for each individual protein-
113 encoding gene alignment and the concatenated alignment for the BI tree was the transitional
114 model with invariable sites and gamma distribution (TIM2+I+G). The best substitution model
115 for the ML tree based in amino acid concatenated sequences was: LG +G+I+F. “+F” indicates
116 the use of the empirical amino acid distribution in the analysed alignment, instead of the model
117 default distribution; and being “LG” the name of the model, given after the authors [17].

118

119 Phylogenetic inference was tested 1,000 times by the bootstrap method in all ML trees, and
120 bootstrap values less than 50% were not included in the trees. The branch-support values in BI,
121 or posterior probabilities (PP) [18, 19] were determined by Markov Chain Monte Carlo
122 sampling. Four chains were run simultaneously 10,000,000 times, one in every 1,000 trees was
123 sampled, and the first 1,000 trees were discarded whilst the other 9,000 were used to calculate
124 the PP. Values of PP lower than 0.5 are not included in the BI trees. *Acinetobacter baumannii*
125 DSM 30007^T was chosen as outgroup in both 16S rRNA analyses, and *P. pertucinogena* LMG
126 1874^T was the outgroup for all concatenated nucleotide and amino acid phylogenetic analyses.

127

128 The topology of both the ML and BI 16S rRNA phylogenetic trees was mostly congruent (Fig.
129 1 and Suppl. Fig. S1). The type strains of both proposed novel species, FRB 228^T and FRB230^T,
130 formed their own cluster within the *P. straminea* group on the border of the *P. fluorescens*
131 lineage [20] with 83% bootstrap support in the ML tree, but lacking support in the BI tree. The
132 closest relative, according to the 16S rRNA phylogenetic analyses, is *P. flavescens* although
133 without significant bootstrap or PP support.

134

135 MLSA based on the protein-coding genes *gyrB*, *rpoD* and *rpoB* have proven useful for
136 examining phylogenetic relationships within the genus *Pseudomonas* [21, 22]. To further

137 clarify the taxonomic position of the *Pseudomonas* strains isolated from AOD-affected oak,
138 MLSA of the *gyrB*, *rpoD* and *rpoB* genes was performed. Both phylogenetic tree constructions
139 (ML and BI) clearly separated the 22 AOD-associated strains into two clusters (MLSA cluster
140 A and MLSA cluster B) with 100% bootstrap and PP support, constituting each a potential
141 novel species (Fig. 2 and Suppl. Fig. S2). MLSA cluster A (*P. daroniae* sp. nov.), containing
142 strains only from Bisham Woods, formed an independent branch with *P. seleniipraecipitans*
143 and *P. flavescens* as the closest phylogenetic relatives, although without significant bootstrap
144 or PP values. MLSA cluster B (*P. dryadis* sp. nov.) contained strains from both Bisham and
145 Great Monk Woods as well as Kew Gardens. The closest phylogenetic neighbours to MLSA
146 cluster B, without bootstrap support in ML trees but with good PP values in BI trees, are
147 *P. argentinensis*, *P. punonensis*, *P. flavescens*, *P. seleniipraecipitans* and *P. straminea*, as well
148 as MLSA cluster A. The ML tree based on the concatenated amino acid sequences was
149 congruent with the trees constructed from nucleotide sequences (Suppl. Fig. S3).

150

151 Three PCR-based DNA fingerprinting techniques including BOX, ERIC (enterobacterial
152 repetitive intergenic consensus) and RAPD (random amplification of polymorphic DNA) were
153 used to investigate the genetic diversity of the AOD-associated *Pseudomonas*. Amplification
154 was performed on all 22 strains, and closest related strains, according to the published protocols
155 [23–25]. Primers BOX-A1R, ERIC-1R and ERIC-2, and OPA-04 were used for the BOX,
156 ERIC, and RAPD PCR reactions, respectively.

157

158 RAPD-PCR was found to be the most discriminative fingerprinting method for the
159 *Pseudomonas* strains. The RAPD patterns for MLSA cluster A showed differences between
160 strain P18A and the remaining strains (Suppl. Fig. S4), while four different band patterns were
161 observed for strains belonging to MLSA cluster B: one for strains isolated in Great Monk
162 Wood (FRB 230^T, P26B, GM38b, GM48c and GM50b), one for Bisham Woods (P6B), and
163 two for strains isolated in Kew Gardens (one pattern for strain I151 and another pattern for
164 I160, I163 and I166). These results suggest that there are two clonal populations within MLSA
165 cluster A and four clonal populations within MLSA cluster B.

166

167 **Genome Features**

168 Representative strains from each MLSA cluster were selected for whole genome sequencing.
169 FRB 228^T, P9A, P18A, P23A, FRB 230^T, P6B and P26B were sequenced by MicrobesNG

170 (Birmingham, UK) using the Illumina HiSeq platform. The reads were trimmed using
171 Trimmomatic [26], and their quality was assessed using in-house scripts combined with the
172 following software: Samtools, BedTools and bwa-mem [27–29].

173

174 De novo assembly was performed on samples using SPAdes version 3.11.1 [30] and the
175 resulting contigs were annotated in Prokka 1.11 [31]. Assembly of strain FRB 228^T yielded 56
176 contigs > 1,000 bp with a total length of 5.43 Mbp, while FRB 230^T generated 71 contigs >
177 1,000 bp with a total length of 5.76 Mbp. The genome sequences of strains FRB 228^T, P9A,
178 P18A, P23A, FRB 230^T, P6B and P26B were submitted to GenBank and received the accession
179 numbers QJUH00000000, QJUI00000000, QJUI00000000, QJUK00000000,
180 QJUN00000000, QJUL00000000 and QJUM00000000, respectively. The DNA G + C mol%
181 content for strains FRB 228^T, P9A, P18A and P23A ranged from 61.97 – 62.02 mol%, and
182 64.94 – 64.97 mol% for strains FRB 230^T, P6B and P26B.

183

184 The average nucleotide identity (ANI) was calculated using JSpecies [32]. Whole genomes of
185 the potential novel *Pseudomonas* species and the closest related species available from
186 GenBank were uploaded and compared. ANI values of 98.3 – 100% were obtained when strains
187 FRB 228^T, P9A, P18A and P23A (MLSA cluster A) were compared to each other, while strains
188 FRB 230^T, P6B and P26B (MLSA cluster B) exhibited 98.6 – 100%. The threshold for species
189 delimitation of 95% ANI has been shown to correspond to the DNA-DNA hybridisation value
190 of 70% [33], confirming that each of the MLSA species clusters corresponds to a single taxon.
191 When the whole genomes of the closest relatives (including *P. flavescens* and *P.*
192 *selenipraecipitans*) were compared to strains belonging to each novel species, ANI values
193 ranging from 74.2 to 87.2% were obtained. In addition to ANI, *in silico* DNA-DNA
194 hybridisation (*isDDH*) using the genome to genome distance calculator (GGDC), was
195 performed on the genomes of the novel species and closest related species [34]. The values
196 generated by *isDDH* were in agreement with the ANI values, with strains in each novel species
197 sharing > 87.9% whole genome similarity, and <39.2% to their closest relatives. The
198 percentages generated by ANI and *isDDH* analysis indicate that the *Pseudomonas* isolates are
199 well below the cut-off limit when compared to the closest related species, supporting their
200 classification as two novel species: *Pseudomonas daroniae* sp. nov. (MLSA cluster A) and
201 *Pseudomonas dryadis* sp. nov. (MLSA cluster B). A detailed summary of the pairwise whole
202 genome comparisons is listed in Suppl. Table S3a and S3b.

203

204

205 Phenotypic and chemotaxonomic characterisation

206 Morphological, physiological and biochemical characteristics were determined for
207 representative strains from the two novel species (*P. daroniae* sp. nov.: FRB 228^T, P9A, P18A,
208 P23A and *P. dryadis* sp. nov.: FRB 230^T, P6B and P26B), as well as type strains from the
209 closest phylogenetic relatives. Cell size and motility were measured using light microscopy
210 and the microscopy imaging software CellSens version 1.11 (Olympus Life Science, Tokyo,
211 Japan). Strains were grown on KB agar for fluorescent pigment production analysis [35] and
212 Gram reaction was determined following growth on LB agar [36]. Oxidase tests were
213 performed according to the manufacturer's instructions (bioMérieux), while catalase activity
214 was monitored by spreading fresh bacterial colonies in 3% w/w hydrogen peroxide (H₂O₂) and
215 recording gas bubbles formation. Tolerance ranges of temperature (4-10°C, 33°C, 37°C, 39°C
216 and 41°C) were studied on KB agar and tryptone soy agar (TSA), while NaCl tolerance (3.5%,
217 4.5%, 5.5%, 6.5%, 7.5%, 8.5% and 9.5%) and pH (6.0, 7.0, 8.0 and 9.0) were studied only on
218 TSA at 25°C. Plates were incubated and monitored for bacterial growth for 6 days. To test if
219 they were able to grow anaerobically, *P. daroniae* sp. nov. and *P. dryadis* sp. nov. strains were
220 incubated in TSA for 4 days at 35°C in anaerobic conditions.

221

222 Transmission electron microscopy (TEM) was used to image cells of *P. daroniae* FRB 228^T
223 and *P. dryadis* FRB 230^T (Fig. 3). The strains were grown in tryptone soy broth in shaking
224 conditions. Cell cultures with an optical density of 1.3 were centrifuged 10 minutes at 3000 g.
225 Cell pellets were resuspended in 200 µL of Tris-saline buffer (50mM Tris pH 8.0, 130mM
226 NaCl). This washing step was performed twice. Five µL of the samples were applied to glow-
227 discharged carbon coated copper grids (Cu 300 mesh). Grids were washed and stained for a
228 minute with 2% (w/v) uranyl acetate. Digital images were acquired in a Tecnai T12 TEM
229 microscope with a CETA 16M camera (ThermoFisher Scientific) at a nominal magnification
230 of 2900.

231

232 Cells from both *P. daroniae* sp. nov., and *P. dryadis* sp. nov. were slightly curved, motile rods,
233 with average sizes of 1.7 µm x 0.4 µm and 2.3 µm x 0.4 µm, respectively. Cells from both
234 species had a single polar flagellum, observed by TEM. All strains were Gram-negative, and
235 positive for catalase and oxidase activity. All isolates grew in visible, cream-coloured, 2 mm
236 diameter colonies on KB agar after incubation at 25°C for 48 hours. Only strain P18A from
237 MLSA cluster A produced a fluorescent pigment, whereas all strains belonging to MLSA

238 cluster B produced fluorescent pigments. All strains of cluster A grew at 33°C but failed to
239 grow at or above 37°C, whereas strains of cluster B grew well at 33°C, 37°C and 39°C, but
240 failed to grow at 41°C. Only strain P18A from cluster A grew at 4–10°C, while all strains of
241 cluster B could grow at 4–10°C. However, compared to typical growth at 25°C all strains
242 produced relatively smaller colonies at these lower temperatures. All strains from both novel
243 species grew at pH 6.0, 7.0, and 8.0. However, after 6 days, only small colonies of P18A
244 appeared on pH 9.0 plates. All isolates from both species grew at a NaCl concentration of 3.5%
245 (w/v) but only FRB 228^T grew on 5.5% (w/v) NaCl plates. No growth from either species was
246 observed on plates containing a higher concentration of NaCl.

247

248 Metabolic profiles, carbohydrate fermentation and carbon utilisation data were obtained using
249 API 20 NE strips, (bioMérieux,), API 50 CH galleries (bioMérieux,) and Biolog GN2
250 Microplates (Biolog), respectively. All tests were performed according to the manufacturer's
251 instructions and strains *P. flavescens* LMG 18387^T, *P. seleniipraecipitans* LMG 25475^T,
252 *P. argentinensis* LMG 22563^T, *P. straminea* LMG 21615^T and *P. punonensis* LMG 26839^T
253 were included as reference strains using the same experimental conditions.

254

255 The phenotypic features that can distinguish *P. daroniae* sp. nov. from its closest relative, *P.*
256 *flavescens*, are the inability to ferment D-maltose, D-saccharose (sucrose) and D-trehalose (API
257 50 CH), the inability to oxidise sucrose, D-trehalose and urocanic acid, and the ability to
258 oxidise D-galacturonic acid, D-glucuronic acid, glucuronamide and D-alanine (Biolog GN2).
259 *P. dryadis* sp. nov. can be phenotypically differentiated from its closest neighbours (*P.*
260 *flavescens*, *P. argentinensis*, *P. punonensis*, *P. seleniipraecipitans*, *P. straminea* and *P.*
261 *daroniae* sp. nov.) by the inability to assimilate L-arabinose (API 20 NE), the inability to
262 produce acid from L-arabinose and D-galactose (API 50 CH), as well as the lack of oxidation
263 of D-galactose and D-gluconic acid (Biolog GN2).

264

265 Interestingly, all strains of *P. daroniae* sp. nov. and *P. dryadis* sp. nov. share an exclusive
266 feature, not present in any of the closest neighbours. This is the ability to weakly ferment D-
267 lyxose, a very rare sugar in nature, that can be found as part of bacterial glycolipids [37]. A
268 summary of the most distinguishing characteristics is presented in Table 1, while detailed
269 results of the phenotypic assays are available in Suppl. Table S4.

270

271 Fatty acid methyl ester (FAME) profiles were determined by Fera Science Ltd. (York, UK) for
272 strains of *P. daroniae* sp. nov. (FRB 228^T, P9A, P18A, P23A) and *P. dryadis* sp. nov. (FRB
273 230^T, P6B P26B) using the MIDI microbial identification system Sherlock Version 6.2. Strains
274 were grown in TSA medium at 28°C for 24 h for this analysis. The results obtained were
275 compared against the library TSBA6 6.10. Similar profiles were obtained for strains of both
276 novel species with summed features 3 (C_{16:1} ω6c and/or C_{16:1} ω7c) and 8 (C_{18:1} ω7c and/or C_{18:1}
277 ω6c) as the major fatty acids. As expected, both proposed novel species possess C_{10:0} 3-OH,
278 C_{12:0} 3-OH and C_{12:0} which are characteristic for members of the genus *Pseudomonas* [38].
279 Furthermore *P. daroniae* sp. nov. and *P. dryadis* sp. nov. display these fatty acids in similar
280 amounts to other members of the *P. straminea* group, along with C_{16:0} and summed features 3
281 and 8. The fatty acid profiles for *P. daroniae* sp. nov., *P. dryadis* sp. nov. and their closest
282 phylogenetic relatives are displayed in Table 2. Other chemotaxonomic characteristics, which
283 could be considered for differentiation of species, are the presence of polar lipids and quinones.
284 However, there is no information available for polar lipids for the five species of the *P.*
285 *straminea* group (*P. straminea*, *P. seleniipraecipitans*, *P. flavescens*, *P. argentinensis* and *P.*
286 *punonensis*) [39–43], and the presence of Q9 has only been reported for two species, namely
287 *P. straminea* and *P. punonensis* [39, 40, 41, 42, 43]. Based on the results provided here from
288 genotypic, phenotypic, genomic and chemotaxonomic data, we propose the classification of
289 two novel species, *P. daroniae* sp. nov. and *P. dryadis* sp. nov.

290

291 **Protologue**

292

293 **Description of *Pseudomonas daroniae* sp. nov.**

294 *Pseudomonas daroniae* (da.ron.i'ae. N.L. fem. adj. *daroniae* from Daron, the Celtic goddess
295 of oak).

296

297 Cells are Gram-negative, slightly curved rods (1.7 μm x 0.4 μm), motile by a polar flagellum,
298 strictly aerobic, non-spore forming, and positive for catalase and oxidase activity. Colonies on
299 KB agar are cream-coloured, circular and convex, with entire margins and measure 2 mm in
300 diameter after 48 hours of incubation at 25°C. Only strain P18A produces fluorescent pigment.
301 Growth is optimum at 28°C and pH 6.0 - 8.0, but strains can grow at 33°C and weakly at 37°C.
302 Only strain P18A is able to grow small colonies on KB at refrigeration temperatures (4-10°C)
303 and at pH 9.0. All strains grow on 3.5% (w/v) NaCl supplemented TSA plates, and only FRB
304 228^T grows on 5.5% (w/v) NaCl.

305 Strains are negative for arginine dihydrolase, indole production, urease, β -glucosidase,
306 gelatinase and β -galactosidase. There is no evidence of denitrification or nitrification activity.
307 D-glucose, L-arabinose, D- mannose, D-mannitol, potassium gluconate, caprate, malate and
308 citrate are assimilated. Acid is produced from: glycerol, L-arabinose, D, galactose, D-glucose,
309 D-fructose, D-mannose, D-mannitol, and weakly from D-lyxose and D-arabitol (API 50
310 CHB/E). L-arabinose, D-fructose, D-galactose, α -D-glucose, D-mannitol, D-mannose, pyruvic
311 acid methyl ester, succinic acid mono-methyl ester, acetic acid, *cis*-aconitic acid, citric acid,
312 D-galacturonic acid, D-gluconic acid, D-glucuronic acid, γ -hydroxybutyric acid, itaconic acid,
313 α -ketoglutaric acid, D,L-lactic acid, quinic acid, succinic acid, bromosuccinic acid, succinamic
314 acid, glucuronamide, L-alaninamide, D-alanine, L-alanine, L-asparagine, L-aspartic acid, L-
315 glutamic acid, hydroxy-L-proline, L-proline, L-pyroglutamic acid, L-serine, γ -amino butyric
316 acid, 2-aminoethanol and glycerol are oxidised (Biolog GN2). Reactions for D-arabitol,
317 dextrin, formic acid, glycogen, D-psicose, turanose α -hydroxybutyric acid, β -hydroxybutyric
318 acid, α -ketobutyric acid, α -ketovaleric acid, propionic acid, D-saccharic acid, D,L-carnitine,
319 putrescine, 2,3-butanediol , and D,L, α -glycerol phosphate are variable. Major fatty acids are
320 C_{12:0}, C_{16:0} and summed features 3 (C_{16:1} ω 6c and/or C_{16:1} ω 7c) and 8 (C_{18:1} ω 7c and/or C_{18:1}
321 ω 6c).

322

323 The type strain FRB 228^T (LMG 31087^T = NCPPB 4672^T) was isolated from *Quercus robur*
324 stem tissue displaying symptoms of acute oak decline in the United Kingdom. The DNA G+C
325 content of the type strain (QJUH00000000) is 62.01 mol%.

326

327 **Description of *Pseudomonas dryadis* sp. nov.**

328 *Pseudomonas dryadis* (dry.a'dis. L. gen. n. of a Dryad, the oak tree nymph)

329

330 Cells are Gram-negative, slightly curved rods (2.3 μ m x 0.4 μ m), motile by a polar flagellum,
331 strictly aerobic, non-spore forming, and positive for catalase and oxidase activity. Colonies on
332 KB agar are cream-coloured, circular and convex, with entire margins and measure 2 mm in
333 diameter after 48 hours of incubation at 25°C. All strains produce fluorescent pigment. Growth
334 is optimum at 28°C but is observed up to 39°C, while weak growth occurs at 4 to 10°C. Growth
335 occurs in the pH range 6.0 to 8.0, and on 3.5% (w/v) NaCl supplemented TSA plates.

336

337 Negative for urease, indole production, arginine dihydrolase, β -glucosidase, gelatinase and β -
338 galactosidase. D-glucose, D-mannose, D-mannitol, potassium gluconate, caprate, malate and
339 citrate are assimilated (API 20 NE). Acid is produced from glycerol, D-glucose, D-fructose,
340 D-mannose, D-mannitol, and weakly from D-lyxose, D-fucose and D-arabitol (API 50
341 CHB/E). L-arabinose, D-arabitol, α -D-glucose, D-mannitol, pyruvic acid methyl ester, succinic
342 acid mono-methyl ester, acetic acid, *cis*-aconitic acid, citric acid, γ -hydroxybutyric acid,
343 itaconic acid, α -ketoglutaric acid, D,L-lactic acid, quinic acid, succinic acid, bromosuccinic
344 acid, L-aspartic acid, L-glutamic acid, hydroxy-L-proline, L-proline, γ -aminobutyric acid and
345 glycerol are oxidised (Biolog GN2). Reactions for dextrin, glycogen, tween 40, tween 80, D-
346 fructose, D-galactose, D-mannose, turanose, formic acid, α -hydroxybutyric acid, β -
347 hydroxybutyric acid, α -ketobutyric acid, propionic acid, succinamic acid, L-asparagine, L-
348 pyroglutamic acid, L-serine, 2-aminoethanol, 2,3-butanediol are variable. Major fatty acids are
349 C_{12:0}, C_{16:0} and summed features 3 (C_{16:1} ω 6c and/or C_{16:1} ω 7c) and 8 (C_{18:1} ω 7c and/or C_{18:1}
350 ω 6c).

351

352 The type strain FRB 230^T (LMG 31087^T = NCPPB 4673^T) was isolated from inner bark and
353 beetle galleries in the phloem of *Quercus robur* in the United Kingdom. The G+C content of
354 the type strain (QJUN00000000) is 64.97 mol%.

355

356 AUTHOR STATEMENTS

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371 **ABBREVIATIONS**

372 AIC: Akaike information criterion

373 ANI: average nucleotide identity

374 AOD: acute oak decline

375 BI: Bayesian inference

376 ERIC-PCR: enterobacterial repetitive intergenic consensus

377 FAME: Fatty acid methyl ester

378 GGDC: genome to genome distance calculator

379 GTR+I+G: general time reversible model with invariable sites and gamma distribution

380 *IsDDH*: *in silico* DNA-DNA hybridisation

381 ML: Maximum likelihood

382 MLSA: multilocus sequence analysis

383 PP: Posterior probabilities

384 RAPD-PCR: random amplification of polymorphic DNA

385 TIM2+I+G: transitional model with invariable sites and gamma distribution

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Figure 1: Maximum likelihood phylogenetic tree of the 16S rRNA gene sequences obtained from the AOD-associated type strains of *Pseudomonas daroniae* sp. nov., *Pseudomonas dryadis* sp. nov. and the closest phylogenetic neighbours. A final dataset of 38 partial 16S rRNA gene sequences of 1399 nucleotides long was compared to infer the evolutionary relationships of the taxa, using the ML method in PhyML. Bootstrap support values higher than 50% (from 1,000 replicates) are shown next to the branches. The closest phylogenetic neighbour to both *Pseudomonas daroniae* sp. nov. and *Pseudomonas dryadis* sp. nov. (in bold) is *Pseudomonas flavescens* with low bootstrap support. The scale indicates the nucleotide substitutions per site. *Acinetobacter baumannii* DSM 30007^T (X811660) was used as an outgroup. Species names are followed by the strain number and the GenBank accession number. ^T = type strain.

Figure 2: Maximum likelihood phylogenetic tree of the concatenated *gyrB*, *rpoB* and *rpoD* gene sequences obtained from 12 AOD-associated strains of *Pseudomonas daroniae* sp. nov., 10 strains of *Pseudomonas dryadis* sp. nov. and the closest phylogenetic neighbours. A final dataset of 58 gene sequences of 2172 nucleotides long was compared to infer the evolutionary relationships of the taxa using the ML method in PhyML. Bootstrap support values higher than 50% (from 1,000 replicates) are shown next to the branches. AOD-associated *Pseudomonas* strains grouped in two well-defined clusters (100% bootstrap support): MLSA cluster A (*Pseudomonas daroniae* sp. nov.) and MLSA cluster B (*Pseudomonas dryadis* sp. nov.), in bold. The scale indicates the nucleotide substitutions per site. *Pseudomonas pertucinogena* LMG 1874^T was used as an outgroup. Species names are followed by the strain number. ^T = type strain.

Figure 3: Transmission electron microscopy images of cells from AOD-associated bacteria *Pseudomonas daroniae* FRB 228^T (left) and *Pseudomonas dryadis* FRB 230^T (right). Scale bar: 1 μ m.

Table 1. Distinguishing phenotypic features of *Pseudomonas daroniae* sp. nov., *Pseudomonas dryadis* sp. nov. and closest phylogenetic neighbours. Data were generated in this study using the commercial tests API 20 NE, API 50 CH and Biolog GN2. For complete phenotypic profile results see Suppl. Table S4. All strains tested were negative for indole production, acidification from D-glucose, esculin hydrolysis, gelatin hydrolysis, presence of arginine dihydrolase, urease and β -galactosidase; and assimilation of N-acetyl-glucosamine, D-maltose, adipate and phenyl-acetate. All strains failed to ferment erythritol, D-arabinose, L-xylose, D-adonitol, methyl- β D-xylopyranoside, L-sorbose, L-rhamnose, dulcitol, inositol, D-sorbitol, methyl- α D-mannopyranoside, methyl- α D-glucopyranoside, N-acetylglucosamine, amygdalin, arbutin, esculin ferric citrate, salicin, D-cellobiose, D-lactose, inulin, D-melezitose, D, raffinose, amidon, glycogen, xylitol, D-turanose, D-tagatose, L-fucose, L-arabitol, potassium gluconate, potassium 2-ketogluconate and potassium 5-ketogluconate. None of the strains oxidised α -cyclodextrin, N-acetyl-D-galactosamine, N-acetyl-D-glucosamine, adonitol, D-cellobiose, i-erythritol, L-fucose, gentiobiose, m-inositol, α -D-lactose, lactulose, maltose, D-melibiose, D-raffinose, L-rhamnose, D-sorbitol, xylitol, D-glucosaminic acid, p-hydroxy-phenylacetic acid, sebacic acid, glycyl-L-aspartic acid, L-histidine, L-leucine, L-ornithine, L-phenylalanine, D-serine, inosine, uridine, thymidine, phenylethylamine and α -D-glucose-1-phosphate. α -Ketovaleric acid was not oxidised by any of the *Pseudomonas dryadis* sp. nov. or the reference strains tested. All strains were able to assimilate D-glucose, D-mannose, D-mannitol, potassium gluconate, caprate, malate and citrate. All strains displayed cytochrome oxidase activity, and were able to ferment D-glucose, D-fructose, D-mannose and D-mannitol. Strains were positive for oxidation of L-arabinose, D-mannitol, pyruvic acid methyl ester, cis-aconitic acid, citric acid, itaconic acid, α -ketoglutaric acid, D,L-lactic acid, bromosuccinic acid, L-aspartic acid, L-glutamic acid, L-proline, and γ -aminobutyric acid. All strains except for *Pseudomonas dryadis* sp. nov. ferment L-arabinose and D-galactose; assimilate L-arabinose and D-galactose; and oxidise D-gluconic acid, L-asparagine, L-pyroglutamic acid.

(1) *Pseudomonas daroniae* sp. nov. (n=3) (2) *Pseudomonas dryadis* sp. nov. (n=3) (3) *Pseudomonas flavescens* LMG 18387^T (4) *Pseudomonas argentinensis* LMG 22563^T (5) *Pseudomonas punonensis* LMG 26839^T (6) *Pseudomonas seleniipraecipitans* LMG 25475^T (7) *Pseudomonas straminea* LMG 21615^T. ^T = type strain.

Characteristic	1	2	3	4	5	6	7
API 20 NE							
Nitrate reduction to nitrites (nitrification)	-	-	-	+	-	+	-
Nitrate reduction to nitrogen (denitrification)	-	-	-	-	-	+	-
Assimilation L-arabinose	+	-	+	+	+	+	+

(API 50 CH) Acid from:

L-Arabinose	+	-	+	+	+	+	+
D-Xylose	-	-	-	+	-	-	-
D-Galactose	+	-	+	+	+	+	+
D-Maltose	-	-	+	+	-	-	-
D-Melibiose	-	-	-	+	-	-	-
D-Saccharose (sucrose)	-	-	+	-	-	-	-
D-Trehalose	-	-	+	+	-	-	-
Gentiobiose	-	-	-	+	-	-	-
D-Lyxose	w+	w+	-	-	-	-	-

(Biolog GN2) Oxidation of:

D-Galactose	+	-	+	+	+	+	+
Sucrose	-	-	+	-	-	-	-
D-Trehalose	-	-	+	+	-	-	-
Succinic acid mono-methyl ester	+	+	+	+	+	+	w+
Acetic acid	+	+	+	-	w+	+	-
D-Galacturonic acid	+	-	-	+	+	+	+
D-Gluconic acid	+	-	+	+	+	+	+
D-Glucuronic acid	+	-	-	+	+	+	+
γ -hydroxybutyric acid	+	+	+	w+	+	+	w+
Malonic acid	-	-	w+	-	+	-	-
Quinic acid	+	+	+	+	+	-	+
Succinic acid	+	+	+	-	+	+	+
Bromosuccinic acid	+	+	+	+	+	+	+
Glucuronamide	+	-	-	-	+	+	+
L-alaninamide	w+	-	-	-	w+	w+	+
D-alanine	+	w+	-	-	w+	+	w+
L-alanine	+	w+	+	-	+	+	+

L-alanyl-glycine	-	-	-	-	w+	w+	w+
Hydroxy-L-proline	+	+	+	-	+	+	+
Urocanic acid	-	-	+	-	w+	-	+
Glycerol	+	+	+	w+	w+	w+	w+

+: 100 % of strains positive; -: 100 % of strains negative; w+: 100 % of strains weakly positive.

Table 2: Percentages of cell fatty acid methyl esters (FAMES) of *Pseudomonas daroniae* sp. nov, and *Pseudomonas dryadis* sp. nov. Summed features are sets of two or more fatty acids, which the Microbial Identification System (MIDI) did not manage to separate. Summed feature 3 is C_{16:1} ω6c and/or C_{16:1} ω7c and summed feature 8 is C_{18:1} ω7c and/or C_{18:1} ω6c. Data was generated in this study, except for the profiles of reference strains, which were obtained from [43]. Values are displayed as average percentage per species investigated, with the standard deviation shown in parentheses. *Pseudomonas daroniae* sp. nov. displayed traces (<1%) of C_{12:0} 2-OH, C_{10:0}, C_{14:0}, C_{17:0}, C_{18:0}, C_{17:1} ω8c and C_{19:0} cyclo ω8c. Traces (<1%) of the fatty acids C_{10:0}, C_{14:0}, C_{18:0} and C_{17:0} iso were found in *Pseudomonas dryadis* sp. nov.

(1) *Pseudomonas daroniae* sp. nov. (n=4) (2) *Pseudomonas dryadiae* sp. nov. (n=3) (3) *Pseudomonas flavescens* LMG 18387^T (4) *Pseudomonas argentinensis* LMG 22563^T (5) *Pseudomonas punonensis* LMT03^T (6) *Pseudomonas straminea* IAM 1598^T. - = not detected. ^T = type strain.

Fatty acid	1	2	3	4	5	6
C_{10:0} 3-OH	3.34 (±0.21)	2.8 (±0.20)	3.74	2.4	4.83	3.91
C_{12:0} 3-OH	3.76 (±0.20)	3.10 (±0.45)	3.55	2.58	4.54	3.57
C_{12:0}	9.88 (±0.60)	9.29 (±0.51)	9.23	7.88	8.31	9.58
C_{16:0}	20.2 (±0.63)	22 (±0.37)	19.75	19.69	15.2	17.63
C_{17:0} cyclo	2.88 (±1.11)	2.74 (±0.42)	-	-	-	-
Summed feature 3	22.15 (±2.13)	17.9 (±0.61)	22.4	21.3	23.7	22.4
Summed feature 8	34.45 (±0.90)	39.68 (±0.29)	38.51	41.52	40.82	39.73