

1 Microbial Systematics

2

3 **The Subdivision of the Genus *Kazachstania* Zubkova (1971) (Saccharomycetaceae)**

4

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31 **Keywords:** *Kazachstania viticola*; *Vanderwaltomyces africanus*; *Arxiozyma telluris*;
32 *Pachytichospora transvaarensis*; *Neovaughania barnettii*.

33

34

35 **Abstract**

36 The genus *Kazachstania* Zubkova was introduced as a monotypic genus including
37 *Kazachstania viticola* Zubkova (1971). Later, a large number of species were accom-
38 modated to the genus. However, the phylogenetic analyses indicated that the genus was
39 extremely diverse, since the species concerned, including the type species, *Kazachstania*
40 *viticola*, showed extremely long branches in the phylogenetic trees. The calculated pair-
41 wise sequence similarities of *K. viticola* were quite low (89.4 - 94.6%) to the remaining 17

42 representative species. The phylogenetic data obtained indicated that the genus formed the
43 so-called *Kazachstania* complex, which was subdivided into 10 taxa at the generic level.

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45

46 The genus *Kazachstania* Zubkova was introduced as a monotypic genus with
47 *Kazachstania viticola* Zubkova (1971). Later, a large number of species were
48 accommodated to the genus (Kurtzman 2003). However, many species of the genus had
49 quite long phylogenetic branches, as well as the type species, *K. viticola*, in the resulting
50 phylogenetic trees (Kurtzman 2003; Vaughan-Martini et al. 2011; Goulianova and
51 Dimitrov 2020). Such a phenomenon suggests that the genus is quite diverse phylogenetic-
52 cally and taxonomic heterogeneous-natured.

53 In a previous paper, the present authors newly constructed the phylogenetic tree based
54 on the 26S rRNA gene D1/D2 domain sequences for the representative species and
55 showed that one of the several long-branched species constituted the new genus
56 *Vanderwaltomyces* with *Vanderwaltomyces africanus*, the new combination (Malimas et
57 al. 2023c).

58

59 This paper deals with the presently available sequence data and gives the different
60 conclusion that the genus *Kazachstania* was subdivided into 10 groups at the generic
61 level.

62

63 The phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences was
64 constructed by the neighbour-joining method for 32 species of the genus *Kazachstania*
65 (Fig. 1). The type species, *K. viticola* produced a long branch with *K. kunashirensis*. The
66 calculated pair-wise sequence similarity was 95.2% between the two species (Table 1), the
67 value of which was almost similar to that (94.1%) between *Vanderwaltozyma polyspora*
68 and *Saccharomyces cerevisiae* used as reference standards (Malimas et al. 2023c). In the
69 additional phylogenetic tree derived from the maximum parsimony method (Fig. 2), *K.*
70 *kunashirensis* was located at a different position independently apart from the type
71 species, *K. viticola*, as found previously in the relationship between *Dipodascopsis*
72 *uninucleata* and *Neoaidaea tothii* (= *Dipodascopsis tothii*) (Lipomycetaceae) (Yamada et
73 al. 2022). Namely, the two species were not tightly coupled but loosely phylogenetically.
74 As a result, *K. viticola* constituted the monotypic genus *Kazachstania*. Incidentally, the
75 calculated pair-wise sequence similarities of *K. viticola* were very low (89.4 - 94.6%) to
76 the remaining 17 representative species of the 31 in the genus (Malimas et al. 2023c).

77

78 Genus I *Kazachstania* Zubkova

79 The type species is *Kazachstania viticola*.

80 1. *Kazachstania viticola* Zubkova (1971)

81 The type strain is NRRL Y-27206.

82 2. *Kazachstania kunashirensis* (James, Cai, Roberts et Collins) Kurtzman (2003)

83 Basionym: *Saccharomyces kunashirensis* James, Cai, Roberts et Collins, Int. J. Syst.
84 Bacteriol. 47: 458, 1997.

85 The type strain is NRRL Y-27209

86

87 As mentioned above, *K. kunashirensis* was not able to be classified phylogenetically in
88 the genus *Kazachstania*. However, the species was temporarily accommodated to the
89 genus (Malimaset al. 2023b, c).

90

91 Genus II *Vanderwaltomyces* Malimas, Vu, Yukphan, Tanasupawat et Yamada

92 The type species is *Vanderwaltomyces africanus*.

93 1. *Vanderwaltomyces africanus* (van der Walt) Malimas, Vu, Yukphan, Tanasupawat et
94 Yamada (2023)

95 The type strain is NRRL Y-8276.

96 Synonym: *Kazachstania africana* (van der Walt) Kurtzman (2003).

97 2. *Vanderwaltoyces martiniae* (James, Cai, Roberts et Collins) Malimas, Vu, Yukphan,
98 Tanasupawat et Yamada (2023) comb. nov.

99 Basionym: *Sacchromyces martiniae* James, Cai, Roberts et Collins. Int. J. Syst.
100 Bacteriol. 47: 458, 1997.

101 The type strain is NRRL Y-409.

102 Synonym: *Kazachstania martiniae* (James, Cai, Roberts et Collins) Kurtzman (2003).

103

104 The calculated pair-wise sequence similarity was 94.1% between *Vanderwaltomyces*
105 *africanus* and *K. viticola* (Table 1). In the phylogenetic tree derived from the neighbour-
106 joining method, *Vandervaltomyces africanus* and *K. martiniae* constituted a common
107 cluster. However, the calculated sequence similarity was 97.0% between the two species,
108 i.e., not “98% or more” nor “beyond the 98% wall” (Yamada et al. 2022; Vu et al. 2022a,
109 b; Malimas et al. 2023a, b). Therefore, the relationship between the two species was
110 similar to that (94.0%) between *Kloeckeraspora osmophilus* (subgroup *a-1*) and
111 *Kloeckeraspora occidentalis* (subgroup *a-2*) (Malimas et al. 2023b).

112

113 Genus III *Arxiozyma* van der Walt et Yarrow

114 The type species is *Arxiozyma telluris*.

115 1. *Arxiozyma telluris* (van der Walt) van der walt et Yarrow (1984)

116 The type strain is YB-4302.

117 Synonym: *Kazachstania telluris* (van der Walt) Kurtzman (2003).

118 2. *Arxiozyma bovina* (Kurtzman et Robnett) Malimas, Vu, Yukphan, Tanasupawat et
119 Yamada (2023) comb. nov.

120 Basionym: *Kazachstania bovina* Kurtzman et Robnett, J. Clin. Microbiol. 43:105.
121 2005.

122 The type strain is NRRL Y-7283.

123 3. *Arxiozyma heterogenica* (Kurtzman et Robnett) Malimas, Vu, Yukphan,
124 Tanasupawat et Yamada (2023) comb. nov.
125 Basionym: *Kazachstania heterogenica* Kurtzman et Robnett, J. Clin. Microbiol.
126 43:107. 2005.
127 The type strain is NRRL Y-27499.
128 4. *Arxiozyma pintolopesii* (Kurtzman, Robnett, Ward et Walsh) Malimas, Vu, Yukphan,
129 Tanasupawat et Yamada (2023) comb. nov.
130 Basionym: *Kazachstania pintolopesii* Kurtzman, Robnett, Ward et Walsh, J. Clin.
131 Microbiol. 43:108. 2005.
132 The type strain is NRRL Y-27500.
133 5. *Arxiozyma slooffiae* (Kurtzman et Robnett) Malimas, Vu, Yukphan, Tanasupawat et
134 Yamada (2023) comb. nov.
135 Basionym: *Kazachstania slooffiae* Kurtzman et Robnett, J. Clin. Microbiol. 43:109.
136 2005.
137 The type strain is NRRL YB-4349.
138
139 The calculated pair-wise sequence similarity was 88.8% between *Arxiozyma telluris*
140 and *K. viticola* (Table 1). The sequence similarities were 96.8 - 98.1% among the four
141 except for *Arxiozyma slooffiae*. Between *Arxiozyma slooffiae* and *Arxiozyma telluris*, the
142 calculated similarity was 95.0%.
143
144 Genus IV *Pachytichospora* van der Walt.
145 The type species is *Pachytichospora transvaarensis*.
146 1. *Pachytichospora transvaarensis* van der Walt (1978)
147 The type strain is NRRL Y-17245.
148 Synonym: *Kazachstania transvaarensis* (van der Walt) Kurtzman (2003).
149 2. *Pachytichospora humatica* (Mikata et Ueda-Nishimura) Malimas, Vu, Yukphan,
150 Tanasupawat et Yamada (2023)
151 Basionym: *Saccharomyces humatica* Mikata et Ueda-Nishimura, Int. J. Syst. Evol
152 Microbiol., 51: 2193. 2001.
153 The type strain is NBRC 10673.
154 Synonym: *Kazachstania humatica* (Mikata et Ueda-Nishimura) Kurtzman (2003).
155 3. *Pachytichospora yakushimaensis* (Mikata et Ueda-Nishimura) Malimas, Vu,
156 Yukphan, Tanasupawat et Yamada (2023)
157 Basionym: *Saccharomyces yakushimaensis* Mikata et Ueda-Nishimura, Int. J. Syst.
158 Evol Microbiol. 51: 2194. 2001.
159 The type strain is IFO 1889.
160 Synonym: *Kazachstania yakushimaensis* (Mikata et Ueda-Nishimura) Kurtzman
161 (2003).
162 4. *Pachytichospora jianica* (Lee, Liu, Young et Chang) Malimas, Vu, Yukphan,
163 Tanasupawat et Yamada (2023)

164 Basionym: *Kazachstania jiainica* Lee, Liu, Young et Chang, FEMS Yeast Res. 8: 116
165 (2008).

166 The type strain is SF1S05.

167

168 The calculated pair-wise sequence similarity was 89.5% between *Pachytichospora*
169 *transvaarensis* and *K. viticola* (Table 1). The sequence similarities were 98.7 - 99.4%
170 among the three species except for *Pachytichospora jiainica*. Between *Pachytichospora*
171 *jiainica* and *Pachytichospora transvaarensis*, the calculated similarity was 96.1%.

172 According to MycoBank Database, the genus *Grigorovia* Gouliamova et Dimitrov
173 (2020) appears to be a synonym of the genus *Pachytichospora* van der Walt (1978) (van
174 der Walt 1978; Goulianova and Dimitrov 2020).

175

176 Genus V *Mikataea* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov.

177 *Mikataea* (Mi.ka'ta.e.a. N. L. fem. n. *Mikataea*. Mikata, in honour of Kozaburo
178 Mikata, the research scientist, Institute of Fermentation Osaka, Japan, who contributed
179 greatly to the yeast systematics)

180 Cells are globose and occur singly or in pairs (Vaughan-Martini et al. 2011). Budding
181 cells are transformed to persistent asci containing two ascospores with a belt-intersporal
182 body and warty wall. Glucose, sucrose and raffinose were fermented. Growth occurs on
183 glucose, sucrose, raffinose, trehalose, inulin, glycerol, ethanol and DL-lactate (Vaughan-
184 Martini et al. 2011). Q-6 is present.

185 The type species is *Mikataea zonata*.

186 1. *Mikataea zonata* (Imanishi, Ueda-Nishimura et Mikata) Malimas. Vu, Yukphan,
187 Tanasupawat et Yamada (2023) comb. nov.

188 Basionym: *Kazachstania zonata* Imanishi, Ueda-Nishimura et Mikata, FEMS Yeast
189 Res. 7: 335, 2007.

190 The type strain is NBRC 100504.

191 2. *Mikataea gamospora* (Imanishi, Ueda-Nishimura et Mikata) Malimas. Vu, Yukphan,
192 Tanasupawat et Yamada (2023) comb. nov.

193 Basionym: *Kazachstania gamospora* Imanishi, Ueda-Nishimura et Mikata, FEMS
194 Yeast Res. 7: 336, 2007.

195 The type strain is NBRC 11056.

196

197 The calculated pair-wise sequence similarity was 93.7% between *Mikataea zonata* and
198 *K. viticola* (Table 1). The sequence similarity was 97.6% between the two species.

199

200 Genus VI *Sinensiomyces* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov.

201 *Sinensiomyces* (Si.nen'si.o.my.ces. N. L. masc. n. *Sinensiomyces*, China fungus)

202 Cells are subglobose to ellipsoidal and occur singly or in pairs (Vaughan-Martini et
203 al. 2011). Budding cells are transformed directly into asci containing two to four
204 cylindrical ascospores, which are able to be liberated from the ascus and agglutinated.

205 Glucose, galactose (+ or s), sucrose (+ or v) and raffinose were fermented. Growth occurs
206 on glucose, sucrose (+ or -), raffinose and trehalose (Vaughan-Martini et al. 2011).

207 The type species is *Sinensiomyces sinensis*.

208 1. *Sinensiomyces sinensis* (Li, Fu et Tang) Malimas, Vu, Yukphan, Tanasupawat et
209 Yamada comb. nov.

210 Basionym: *Kluyveromyces sinensis* Li, Fu et Tang, Acta Microbiol. Sin. 30: 96. 1990.

211 The type strain is NRRL Y-27222.

212 Synonym: *Kazachstania sinensis* (Li, Fu et Tang) Kurtzman (2003).

213 2. *Sinensiomyces naganishii* (Mikata, Ueda-Nishimura et Hisatomi) Malimas, Vu,
214 Yukphan, Tanasupawat et Yamada comb. nov.

215 Basionym: *Saccharomyces naganishii* Mikata, Ueda-Nishimura et Hisatomi, Int. J.
216 Syst. Evol. Microbiol. 51: 2191, 2001.

217 The type strain is NBRC 10181.

218 Synonym: *Kazachstania naganishii* (Mikata, Ueda-Nishimura et Hisatomi) Kurtzman
219 (2003).

220

221 The calculated pair-wise sequence similarity was 91.6% between *Sinensiomyces*
222 *sinensis* and *K. viticola* (Table2). The sequence similarity was 99.2% between the two
223 species.

224

225 Genus VII *Baiomyces* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov

226 *Baiomyces* (Ba.i.o.my'ces. N. L. masc. n. *Baiomyces*, Bai fungus, in honour of Dr. F.-
227 Y. Bai, the yeast taxonomist in China, who contributed greatly in the yeast systematics)

228 Cells are globose to subglobose and occur singly, in pairs or in groups (Vaughan-
229 Martini et al. 2011). Budding is multilateral on a narrow base. Pseudohyphae are not
230 formed. Budding cells are transformed into persistent asci, each of which contains one
231 ascospore. Glucose and galactose were fermented. Growth occurs on glucose and
232 galactose (Vaughan-Martini et al. 2011).

233 The type species is *Baiomyces aquatica*.

234 1. *Baiomyces aquatica* (Bai et Wu) Malimas, Vu, Yukphan, Tanasupawat et Yamada
235 (2023) comb. nov.

236 Basionym: *Kazachstania aquatica* Bai et Wu, Int. J. Syst. Evol. Microbiol. 55: 2221,
237 2005.

238 The type strain is AS 2.0706.

239 2. *Baiomyces siamensis* (Limtong, Yongmanitchai, Tun, Kawasaki et Seki) Malimas,
240 Vu, Yukphan, Tanasupawat et Yamada (2023) comb. nov.

241 Basionym: *Kazachstania siamensis* Limtong, Yongmanitchai, Tun, Kawasaki et Seki,
242 Int. J. Syst. Evol. Microbiol. 57: 421, 2007.

243 The type strain is NBRC 101968.

244

245 The calculated pair-wise sequence similarity was 94.6% between *Baiomyces aquatica*
246 and *K. viticola* (Table 1). The sequence similarity was 97.0% between the two species.

247

248 Genus VIII *Neoyamazakia* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov.

249 *Neoyamazakia* (Ne.o.ya.ma.za'ki.a. N. L. fem. n. *Neoyamazakia*, New Yamazaki, in
250 honour of Dr. Atsushi Yamazaki, NITE Biotechnology Center (NBRC), who contributed
251 greatly to the yeast systematics)

252 Cells are ovoid or elongate and occur singly or in pairs (Vaughan-Martini et al. 2011).
253 Budding is at various positions near the pole of the cells. Pseudohyphae are not produced.
254 asci are elongated, persistent and contain one occasionally two or four spherical
255 ascospores. Glucose and galactose are fermented. Growth occurs on glucose and galactose
256 (Vaughan-Martini et al. 2011).

257 The type species is *Neoyamazakia rosinii*.

258 1. *Neoyamazakia rosinii* (Vaughan-Martini, Barcaccia et Pollacci) Malimas, Vu,
259 Yukphan, Tanasupawat et Yamada (2023) comb. nov.

260 Basionym: *Saccharomyces rosinii* Vaughan-Martini, Barcaccia et Pollacci, Int. J. Syst.
261 Bacteriol. 46: 616. 1996.

262 The type strain is NRRL Y-17919.

263 Synonym: *Kazachstania rosinii* (Vaughan-Martini, Barcaccia et Pollacci) Kurtzman
264 (2003).

265 2. *Neoyamazakia piceae* (Weber, Sapaaij et van der Walt) Malimas, Vu, Yukphan,
266 Tanasupawat et Yamada (2023) comb. nov.

267 Basionym: *Kluyveromyces piceae* Weber, Sapaaij et van der Walt, Antonie van
268 Leeuwenhoek 62: 240. 1992.

269 The type strain is NRRL Y-17977.

270 Synonym: *Kazachstania piceae* (Weber, Sapaaij et van der Walt) Kurtzman (2003).

271 3. *Neoyamazakia lodderae* (van der Walt et Tscheuschner) Malimas, Vu, Yukphan,
272 Tanasupawat et Yamada (2023) comb. nov.

273 Basionym: *Saccharomyces lodderae* van der Walt et Tscheuschner, Antonie van
274 Leeuwenhoek 23: 188. 1957.

275 The type strain is NRRL Y-8280.

276 Synonym: *Kazachstania lodderae* (van der Walt et Tscheuschner) Kurtzman (2003).

277 4. *Neoyamazakia spencerorum* (van der Walt et Tscheuschner) Malimas, Vu, Yukphan,
278 Tanasupawat et Yamada (2023) comb. nov.

279 Basionym: *Saccharomyces spencerorum* Vaughan-Martini, Antonie van Leeuwenhoek
280 68: 116. 1995.

281 The type strain is NRRL Y-17920.

282 Synonym: *Kazachstania spencerorum* (van der Walt et Tscheuschner) Kurtzman
283 (2003).

284

285 The calculated pair-wise sequence similarity was 92.8% between *Neoyamazakia*
286 *rosinii* and *K. viticola* (Table 1). The sequence similarities were 95.2 - 97.7% among the
287 four species within the genus, as already shown in the genus *Kloeckeraspora* (subgroups
288 *a-1* and *a-2*) and the genus *Hanseniaspora* (subgroups *b-1* and *b-2*) (Malimas et al.
289 2023b).

290

291 Genus IX *Capriottia* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov.
292 *Capriottia* (Cap.ri.ot'ti.a N. L. fem. n. *Capriottia*, in honour of Dr. A. Capriotti, Italian
293 yeast taxonomist, who contributed to the yeast systematics)

294 Cells are subglobose to elliposoidal and occur singly or in pairs (Vaughan-Martini et
295 al. 2011). Budding cells are transformed directly into asci usually containing one, but
296 occasionally up to four, globose to short ellipsoidal ascospores. Glucose and galactose are
297 fermented. Growth occurs on glucose, galactose and trehalose (Vaughan-Martini et al.
298 2011).

299 The type species is *Capriottia servazzii*.

300 1. *Capriottia servazzii* (Capriotti) Malimas, Vu, Yukphan, Tanasupawat et Yamada
301 comb. nov.

302 Basionym: *Saccharomyces servazzii* Capriotti, Ann. Microbiol. Enzimol. 17: 83. 1967.

303 The type strain is NRRL Y-12661.

304 Synonym: *Kazachstania servazzii* (Capriotti) Kurtzman (2003).

305 2. *Capriottia unispora* (Jørgensen) Malimas, Vu, Yukphan, Tanasupawat et Yamada
306 (2023) comb. nov.

307 Basionym: *Saccharomyces unisporus* Jørgensen, Die Mikroorganismen der Gärungs-
308 industrie, 5te Aufl., p. 371, 1909. P. Parey, Berlin.

309 The type strain is NRRL Y-1556.

310 Synonym: *Kazachstania unispora* (Jørgensen) Kurtzman (2003).

311 3. *Capriottia solicola* (Bai et Wu) Malimas, Vu, Yukphan, Tanasupawat et Yamada
312 (2023) comb. nov.

313 Basionym: *Kazachstania solicola* Bai et Wu, Int. J. Syst. Evol. Microbiol. 55: 2222.
314 2005.

315 The type strain is CBS 6904.

316 4. *Capriottia aerobia* (Lu, Cai, Wu et Bai) Malimas, Vu, Yukphan, Tanasupawat et
317 Yamada (2023) comb. nov.

318 Basionym: *Kazachstania aerobia* Lu, Cai, Wu et Bai, Int. J. Syst. Evol. Microbiol. 54:
319 2434. 2004.

320 The type strain is AS 2.2384.

321

322 The calculated pair-wise sequence similarity was 94.1% between *Capriottia servazzii*
323 and *K. viticola* (Table 1). The sequence similarities were 98.8 - 100% among the four
324 species.

325

326 Genus X *Neovaughania* Malimas, Vu, Yukphan, Tanasupawat et Yamada gen. nov.
327 *Neovaughania* (Ne.o.va.ug'ha.ni.a. N. L. fem. n. *Neovaughania*, New Vaughan, in
328 honour of Dr. Vaughan-Martini, the yeast taxonomist in Italy, who contributed greatly to
329 the yeast systematics)
330 Cells are globose or slightly ovoid and occur singly or in pairs (Vaughan-Martini et al.
331 2011). Pseudohyphae are absent. Asci formed one or two globose ascospores. Glucose,
332 galactose and sucrose are fermented. Growth occurs on glucose, sucrose, raffinose,
333 galactose and trehalose (Vaughan-Martini et al. 2011).
334 The type species is *Neovaughania barnettii*.
335 1. *Neovaughania barnettii* (Vaughan-Martini) Malimas, Vu, Yukphan, Tanasupawat et
336 Yamada (2023) comb. nov.
337 Basionym: *Saccharomyces barnettii* Vaughan-Martini, Antonie van Leeuwenhoek 68:
338 116. 1995.
339 The type strain is NRRL Y-27223.
340 Synonym: *Kazachstania barnettii* (Vaughan-Martini) Kurtzman (2003).
341 2. *Neovaughania bulderi* (Middelhoven, Kurtzman et Vaughan-Martini) Malimas, Vu,
342 Yukphan, Tanasupawat et Yamada (2023) comb. nov.
343 Basionym: *Sacchromyces bulderi* Middelhoven, Kurtzman et Vaughan-Martini,
344 Antonie van Leeuwenhoek 77: 224. 2000.
345 The type strain is NRRL Y-27203.
346 Synonym: *Kazachstania bulderi* (Middelhoven, Kurtzman et Vaughan-Martini)
347 Kurtzman (2003).
348 3. *Neovaughania exigua* (Reess ex Hansen) Malimas, Vu, Yukphan, Tanasupawat et
349 Yamada (2023) comb. nov.
350 Basionym: *Saccharomyces exiguus* Reess ex Hansen, Trav. Lab. Carlsberg 2: 146.
351 1888.
352 The type strain is NRRL Y-12640.
353 Synonym: *Kazachstania exigua* (Reess ex Hansen) Kurtzman (2003).
354 4. *Neovaughania turicensis* (Wyder, Meile et Teuber) Malimas, Vu, Yukphan,
355 Tanasupawat et Yamada (2023) comb. nov.
356 Basionym: *Saccharomyces turicensis* Wyder, Meile et Teuber, Syst. Appl. Microbiol.
357 22: 423. 1999.
358 The type strain is NRRL Y-27345.
359 Synonym: *Kazachstania turicensis* (Wyder, Meile et Teuber) Kurtzman (2003).
360
361 The calculated pair-wise sequence similarity was 91.3% between *Neovaughania*
362 *barnettii* and *K. viticola* (Table 1). The sequence similarities were 98.7 - 99.6% among the
363 four species. Accordingly, the four species constituted a taxonomic homogeneous-natured
364 genus, since the calculated similarities were “98% or more”, i.e., “beyond the 98% wall”
365 (Yamada et al. 2022; Vu et al. 2022a, b; Malimas et al. 2023a, b).
366

367 As described above, the genus *Kazachstania* was extremely diverse phylogenetically
368 and produced the so-called *Kazachstania* complex. In future, a large number of species
369 will be described within the so-called *Kazachstania* complex.

370

371

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378

379 Conflict of interest

380 The authors declare that there are no conflicts of interest.

381

382 Author contributions

383 T. M., H.T.L. V., P. Y., S. T. and Y. Y. designed the study. T. M. performed the main
384 experiments. H.T.L. V. and P. Y. instructed the experiments. Y. Y. prepared the manuscript.
385 The detailed discussions were made among the five.

386

387

388 **References**

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390 and other members of the Saccharomycetaceae and the proposal of the new genera
391 *Lachancea*, *Nakaseomyces*, *Naumovia*, *Vanderwaltozyma* and *Zygorulasporea*.
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Table 1. The pair-wise sequence similarity in the 26S rRNA gene D1/D2 domain sequences in *Kazchstania* and related species.

Sp	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31							
1	100																																					
2	95.2	100																																				
3	94.1	94.1	100																																			
4	94.8	95.0	97.0	100																																		
5	88.8	90.6	89.3	90.4	100																																	
6	89.0	90.3	89.0	90.1	97.4	100																																
7	89.7	90.4	90.3	91.0	97.6	96.8	100																															
8	88.8	89.9	89.2	90.1	97.7	97.0	98.1	100																														
9	90.3	91.4	90.4	91.5	95.0	95.7	95.5	94.6	100																													
10	89.5	90.6	91.7	91.3	88.4	88.4	88.2	88.1	87.9	100																												
11	89.7	90.8	92.4	91.7	88.7	87.7	88.2	87.5	87.9	99.2	100																											
12	89.3	90.4	92.4	92.0	87.9	87.5	88.4	87.7	87.9	98.7	99.4	100																										
13	89.7	90.8	91.5	91.1	87.7	87.7	88.6	87.2	88.4	96.1	96.1	95.9	100																									
14	93.7	94.6	94.3	95.4	89.9	89.9	90.1	89.2	90.8	91.1	91.0	90.6	90.0	100																								
15	92.6	93.0	93.0	93.3	88.8	88.4	89.2	88.2	89.3	90.9	90.8	90.8	89.8	97.6	100																							
16	91.6	92.3	91.2	91.6	89.5	89.4	89.9	90.3	90.3	87.9	87.9	87.9	88.1	91.0	90.1	100																						
17	91.9	92.7	91.6	91.9	89.9	89.9	90.3	90.6	90.6	88.3	88.3	88.3	87.7	91.0	90.5	99.2	100																					
18	94.6	95.7	95.0	96.8	91.2	90.6	92.1	90.8	91.7	90.6	90.8	90.8	90.4	95.0	93.4	92.7	93.0	100																				
19	94.1	94.5	93.9	94.8	91.4	91.6	92.3	91.0	92.1	90.6	90.6	90.3	89.7	93.9	92.8	92.3	92.7	97.0	100																			
20	92.8	93.4	94.6	95.4	89.7	89.3	90.6	89.5	90.3	90.8	91.1	91.1	90.6	95.0	94.3	91.6	91.9	94.5	93.9	100																		
21	94.1	94.3	95.7	96.3	89.5	89.2	90.8	89.7	90.4	90.6	91.3	91.3	90.6	95.7	95.0	91.2	91.6	95.0	94.3	97.7	100																	
22	93.9	94.8	95.5	96.5	90.8	90.4	91.7	90.6	91.5	90.1	90.4	90.4	90.1	95.9	94.3	92.5	92.8	95.4	94.3	96.5	97.4	100																
23	94.5	93.9	94.8	95.0	89.5	89.5	90.8	89.7	90.3	89.9	90.1	89.7	89.1	95.4	93.9	91.9	92.3	94.6	93.2	95.2	96.5	97.6	100															
24	94.1	95.9	95.7	97.2	91.9	91.0	92.1	91.5	91.9	91.5	91.5	91.9	90.6	95.6	93.5	92.8	93.2	97.0	95.4	95.2	95.7	96.1	94.8	100														
25	94.6	96.1	96.5	96.8	91.5	91.0	92.1	91.4	92.3	91.2	91.9	91.9	91.0	95.2	93.5	92.8	93.2	96.7	95.8	95.7	96.6	96.3	95.0	98.8	100													
26	94.3	96.1	96.1	96.8	91.9	91.0	92.1	91.5	91.9	91.5	91.5	91.5	91.0	95.6	93.5	93.0	93.4	97.0	95.4	95.4	95.9	96.3	95.0	99.6	99.2	100												
27	94.3	96.1	96.1	96.8	91.9	91.0	92.1	91.5	91.9	91.5	91.5	91.5	91.0	95.6	93.5	93.0	93.4	97.0	95.4	95.4	95.9	96.3	95.0	99.6	99.2	100	100											
28	91.3	93.5	91.7	92.4	91.9	91.1	92.2	92.1	91.1	89.1	88.8	88.8	90.4	91.3	91.0	91.6	91.2	93.2	92.1	92.1	91.9	92.6	92.3	93.4	92.8	93.5	93.5	100										
29	91.3	93.5	91.7	92.4	91.5	91.5	91.9	91.7	91.1	89.1	88.8	88.8	90.4	91.3	91.0	91.6	91.2	92.8	92.7	92.1	91.9	92.6	92.3	93.0	92.8	93.2	93.2	99.6	100									
30	91.5	93.5	92.3	92.8	91.1	90.8	91.5	91.3	91.1	88.8	88.8	88.8	90.1	91.0	90.4	91.0	90.6	93.2	93.0	91.7	91.9	92.3	91.7	92.6	92.8	92.8	92.8	92.8	98.7	98.7	100							
31	91.5	93.7	91.9	92.3	91.3	91.0	91.7	91.5	91.0	89.3	89.0	89.0	90.6	91.5	91.2	91.7	91.4	93.0	92.8	92.3	92.1	92.4	92.1	93.2	93.0	93.4	93.4	99.4	99.4	98.8	100							

1, *Kazachstania viticola*; 2, *Kazachstania kunashirensis*; 3, *Vanderwaltomyces africanus*; 4, *Vanderwaltoyces martiniae*; 5, *Arxiozoma telluris*; 6, *Arxiozoma bovina*; 7, *Arxiozoma heterogenica*; 8, *Arxiozoma pintolopesii*; 9, *Arxiozoma slooffiae*; 10, *Pachytichospora transvaarensis*; 11, *Pachytichospora humatica*; 12, *Pachytichospora yakushimaensis*; 13, *Pachytichospora jiainica*; 14, *Mikataea zonata*; 15, *Mikataea gamospora*, 16, *Sinensiomyces sinensis*; 17, *Sinensiomyces naganishii*; 18, *Baiomyces aquatica*; 19, *Baiomyces siamensis*; 20, *Neoyamazakia rosinii*; 21, *Neoyamazakia piceae*; 22, *Neoyamazakia lodderae*; 23, *Neoyamazakia spencerorum*; 24, *Capriottia servazzii*; 25, *Capriottia unispora*; 26, *Capriottia solicola*; 27, *Capriottia aerobia*; 28, *Neovaughania barnettii*, 29, *Neovaughania bulderi*; 30, *Neovaughania exigua*; 31, *Neovaughania turicensis*. For calculating the sequence similarity, 542-548 bases were used.

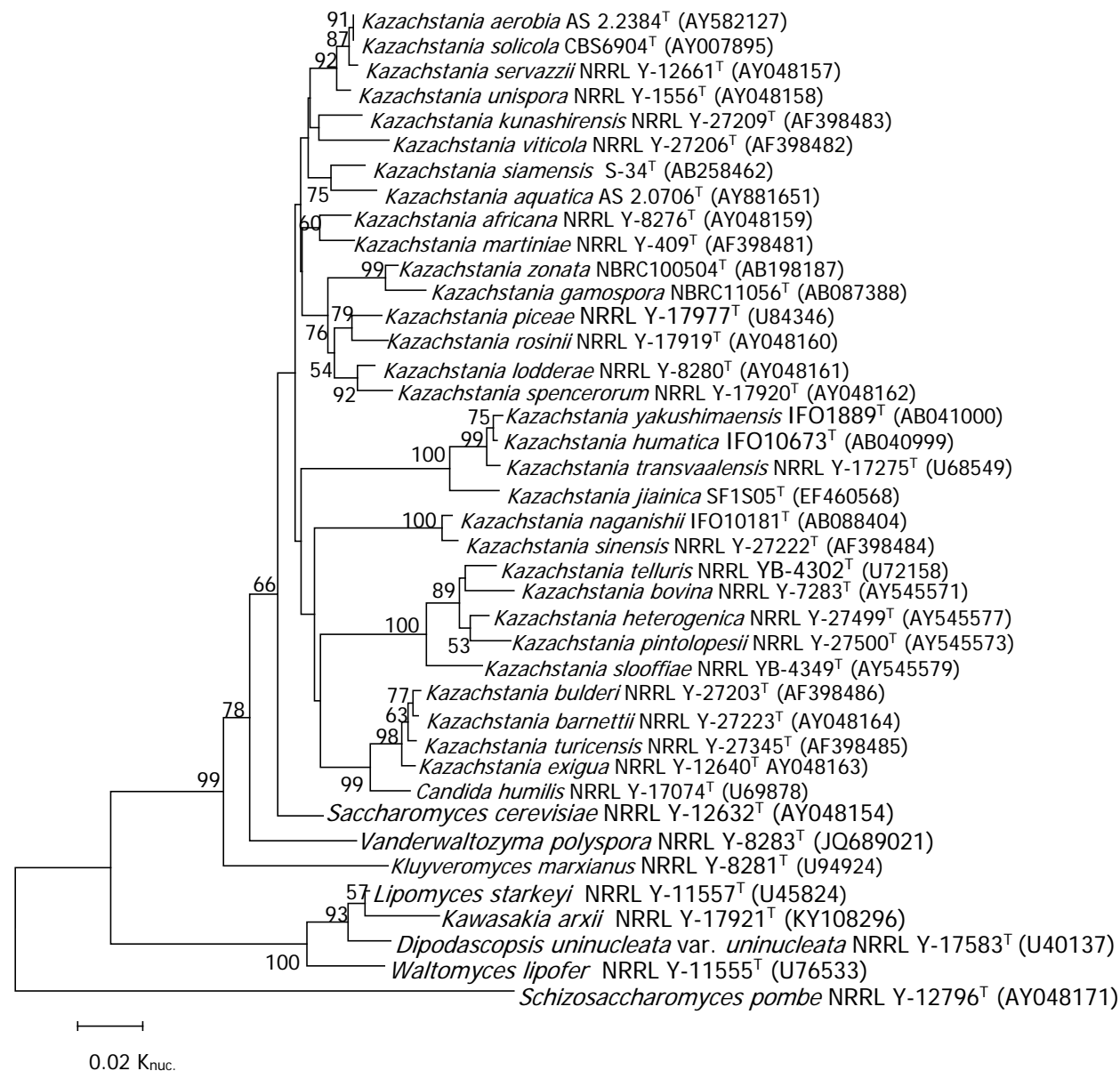


Fig. 1. The phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences with 502 bases derived from the neighbour-joining method. The numerals at the nodes of respective branches indicate bootstrap values (%) deduced from 1000 replications

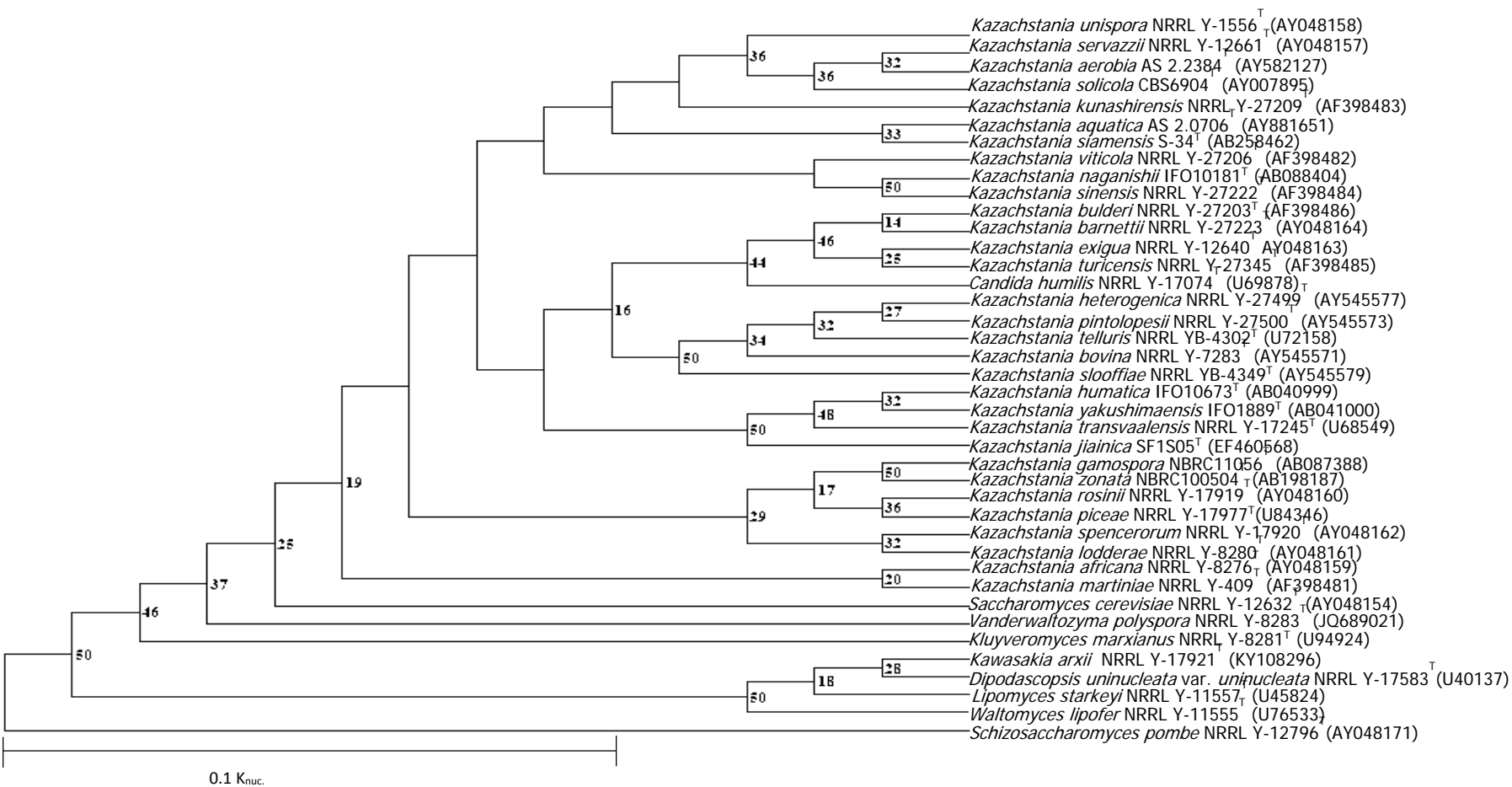


Fig. 2. The phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences with 502 bases derived from the Maximum Parsimony method. The numerals at the nodes of respective branches indicate bootstrap values (%) deduced from 1000 replications