

1 Short Communication

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3 **The Generic Circumscription of *Tetrapisispora* Ueda-Nishinura et Mikata (1999)**
4 **(Saccharomycetaceae): The Proposal of *Nishimuraea* gen. nov.**

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29 **ABSTRACT**

30 The genus *Tetrapisispora* was introduced with the following four species; *T. arboricola*, *T.*
31 *iriomotenensis*, *T. nanseiensis* and *T. phaffii* (the type species). Later, the three species
32 were additionally accommodated to the genus; *T. blattae*, *T. fleetii* and *T. namnaonensis*.
33 Of the seven species, *Tetrapisispora blattae* represented an abnormally long branch in the
34 phylogenetic trees based on the 26S rRNA gene D1/D2 domain and the 18S rRNA gene
35 sequences derived from the neighbour-joining method. The calculated pair-wise 26S
36 rRNA gene D1/D2 domain sequence similarities were extremely low (81.2 - 95.7%)
37 within the genus. Except for *T. blattae*, the similarities were 90.3 - 95.7% among the
38 remaining five species. From the data obtained, *Nishimuraea* gen. nov. was proposed
39 along with *Nishimuraea blattae* comb. nov.

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41 **Keywords:** *Tetrapisispora phaffii*; *Tetrapisispora blattae*; *Tetrapisispora namnaonensis*;
42 *Nishimuraea* gen. nov.; *Nishimuraea blattae* comb. nov.

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45 The genus *Tetrapisispora* Ueda-Nishimura et Mikata was introduced for the following
46 four species, *T. arboricola*, *T. iriomotenensis*, *T. nanseiensis* and *T. phaffii* (= *Kluyvero-*
47 *myces phaffii*) (Ueda-Nishimura and Mikata 1999). Subsequently, *T. blattae*, (= *Kluyvero-*
48 *myces blattae*), *T. fleetii* and *T. namnaonensis* were included in the genus (Kurtzman 2003;
49 Kurtzman et al. 2004; Sumpradit et al. 2005).

50 Of the seven species, *T. blattae* was characteristic of an extremely long branch in the
51 phylogenetic trees (Lachance 2011; Sumpradit et al. 2005). Especially, *T. blattae* was
52 located outside the cluster comprised of the remaining six species and then connected
53 surprisingly to that of *Lachancea thermotolerans* (= *Zygosaccharomyces thermo-*
54 *tolerans*)/*Zygosaccharomyces rouxii* in the phylogenetic tree based on the D1/D2 domain
55 of the LSU rDNA derived from the neighbour-joining method (Sumpradit et al. 2005).

56

57 This paper deals with the presently available sequence data and gives the different
58 conclusion that *T. blattae* constitutes an independent taxon at the generic level since the
59 species has an abnormally long branch and the phylogenetic position of the species is not
60 found inside the cluster of *T. phaffii*, the type species and the remaining five.

61

62 The phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences (LSU
63 D1/D2) was constructed by the neighbour-joining method (Fig. 1). *Tetrapisispora phaffii*,
64 the type species was connected to *T. namnaonensis* and *T. fleetii* first and then to the
65 remaining three species, *T. iriomotensis*, *T. nanseiensis* and *T. arboricola*. The phylo-
66 genetic branches of the six species were very long within the genus. The lengths of the
67 species almost corresponded to those of the nine species in the different genera (Fig. 1).
68 *Tetrapisispora blattae*, the remaining one was surprisingly located outside the six species
69 as well as the nine species used for reference standards.

70 The calculated pair-wise sequence similarities were 81.2 - 95.7% within the genus and
71 90.3 - 95.7% among the five species except for *T. blattae* (Table 1). The calculated values
72 were neither '98% or more' nor 'beyond the so-called 98% wall' (Yamada et al., 2022; Vu
73 et al., 2022a, b; Malimas et al., 2023a-d). In contrast, the sequence similarity between *T.*
74 *blattae* and *T. phaffii* was quite low (82.6%), the value of which was much lower than that
75 (88.8%) between *T. blattae* and *Saccharomyces cerevisiae* and to that (87.7%) between *T.*
76 *blattae* and *Vanderwaltozyma polyspora*. Incidentally, the sequence similarity between *V.*
77 *polyspora* and *S. cerevisiae* was 93.8%. From the phylogenetic data obtained above, *T.*
78 *blattae* was quite unique.

79

80 In the phylogenetic tree based on the 18S rRNA gene sequences (SSU) (Fig. 2), the
81 branches were on the whole shorter among the species examined, as shown and discussed
82 previously (Malimas et al., 2003a-d). The five species of the genus *Tetrapisispora*
83 represented quite short phylogenetic branches in contrast to those of LSU D1/D2.

84 However, it is of great interest that there was almost no change in the branch length of *T.*
85 *blattae*.

86 The calculated pair-wise sequence similarities were very high (99.3 - 99.8%) among
87 the five species except for *T. blattae* (Table 2). In contrast, the sequence similarity between
88 *T. blattae* and *T. phaffii* was quite low (94.4%), the value of which was the same as that
89 (94.4%) between *T. blattae* and *S. cerevisiae* and also to that (94.4%) between *T. blattae*
90 and *V. polyspora*. Incidentally, the sequence similarity between *V. polyspora* and *S.*
91 *cerevisiae* was 98.9%. From the phylogenetic data obtained above, *T. blattae* was quite
92 unique.

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94 *Tetrapisispora blattae* was different phylogenetically from the remaining six species of
95 the genus. Therefore, the species is adequate to be taxonomically distinguished at the
96 generic level.

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98 The family Saccharomycetaceae

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100 *Nishimuraea* Malimas, Vu, Yukpnan, Tanasupawat et Yamada gen. nov.

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102 *Nishimuraea* (Ni.shi'mu.ra.e.a. N. L. fem. n. *Nishimuraea*, Nishimura, in honour of
103 Dr. K. Ueda-Nishimura, Institute for Fermentation, Osaka, Japan, who contributed greatly
104 in the yeast systematics)

105 Growth is butyrous, glossy and white to cream coloured (Lachance 2011). Cells are
106 globose to ellipsoid and occur singly, in pairs or in chains. Pseudohyphae and true hyphae
107 are not formed. Ascus arises from diploid cells, and one to eight or more spherical to
108 ellipsoidal ascospores are produced. The spores are liberated from the ascus soon
109 (Lachance 2011). Glucose and galactose are fermented. Growth occurs on glucose,
110 galactose and glycerol. Q-6 is present (Lachance 2011).

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MycoBank number is 848291.

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113 *Nishimuraea blattae* (Henninger et Windisch) Malimas, Vu, Yukphan, Tanasupawat et
114 Yamada comb. nov.

115 Basionym: *Kluyveromyces blattae* Henninger et Windisch, Arch. Microbiol. 109: 155.
116 1976.

117 The type strain is CBS 8765. Mycobank number is 848292.

118 Synonym: *Tetrapisispora blattae* (Henninger et Windisch) Kurtzman (2003).

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120 As described above, *Nishimuraea blattae* is extremely unique phylogenetically. The
121 additional species are expected to be described in the genus *Nishimuraea*.

122

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130 Conflict of interest

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

132

133 Author contributions

134 T. M., H.T.L. V., P. Y., S. T. and Y. Y. designed the study. T. M. performed the main
135 experiments. H.T.L. V. and P. Y. instructed the experiments. Y. Y. prepared the manuscript.

136 The detailed discussions were made among the five.

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178 *Tetrapisispora* 属の限界: 新属 *Nishimuraea* の提案

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189 *Tetrapitisporea* 属には7種が存在する. その中の1種, *Tetrapisispora blattae*は異常
190 に長い系統枝を有し, the type species, *Tetrapisispora phaffii* をはじめとする他の6種
191 とは系統樹の上では、異なった挙動を示した. rRNA遺伝子塩基配列での7種の類
192 似度は, 81.2 - 95.7% と極めて低い値を示した. さらに, *T. blattae*/*T. phaffii* (the type
193 species) 間の塩基配列類似度は, 82.6%であった. 従って, 本種に対して属を別にす

194 ることが可能であり, 新属 *Nishimuraea* および *Nishimuraea blattae* comb. nov. を
 195 とともに提案した.

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Table 1. The pair-wise sequence similarity in the 26S rRNA gene D1/D2 domain sequences in *Tetrapisispora* species.

Species	1	2	3	4	5	6	7	8
1. <i>T. phaffii</i>	100							
2. <i>T. iriomotensis</i>	90.3	100						
3. <i>T. nanseiensis</i>	90.7	91.9	100					
4. <i>T. arboricola</i>	91.2	93.2	95.7	100				
5. <i>T. namnaonensis</i>	91.3	92.4	92.1	93.1	100			
6. <i>N. blattae</i>	82.6	81.6	81.8	82.3	81.2	100		
7. <i>S. cerevisiae</i>	81.9	84.2	83.9	84.8	84.1	88.8	100	
8. <i>V. polyspora</i>	81.2	83.0	82.5	83.0	82.3	87.7	93.8	100
Species	9	10	11	12				
9. <i>L. starkeyi</i>	100							
10. <i>W. lipofer</i>	95.2	100						
11. <i>D. uninucleata</i>	95.7	94.0	100					
12. <i>K. arxii</i>	97.5	94.1	95.4	100				

T, *Tetrapisispora*; *N*, *Nishimuraea*; *S*, *Saccharomyces*; *V*, *Vanderwaltozyma*; *L*, *Lipomyces*; *W*, *Waltomyces*; *D*, *Dipoascopsis*; *K*, *Kawasakia*. The total sequences were of 549 - 568 bases.

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The type strains were used.

Table 2. The pair-wise sequence similarity in the 18S rRNA gene sequences in *Tetrapisispora* species.

Species	1	2	3	4	5	6	7	8
1. <i>T. phaffii</i>	100							
2. <i>T. iriomotensis</i>	99.6	100						
3. <i>T. nanseiensis</i>	99.5	99.4	100					
4. <i>T. arboricola</i>	99.5	99.3	99.8	100				
5. <i>T. namnaonensis</i>	99.8	99.5	99.7	99.6	100			
6. <i>N. blattae</i>	94.4	94.4	94.3	94.4	94.4	100		
7. <i>S. cerevisiae</i>	96.0	96.1	96.2	96.1	96.1	94.4	100	
8. <i>V. polyspora</i>	96.0	96.1	96.0	96.0	96.1	94.4	98.9	100
Species	9	10	11	12				
9. <i>L. starkeyi</i>	100							
10. <i>W. lipofer</i>	95.5	100						
11. <i>D. uninucleata</i>	95.7	97.6	100					
12. <i>K. arxii</i>	95.9	96.7	97.5	100				

T, *Tetrapisispora*; *N*, *Nishimurea*; *S*, *Saccharomyces*; *V*, *Vanderwaltozyma*; *L*, *Lipomyces*; *W*, *Waltomyces*; *D*, *Dipoascopsis*; *K*, *Kawasaki*. The total sequences were of 1705 - 1734 bases. The type strains were used.

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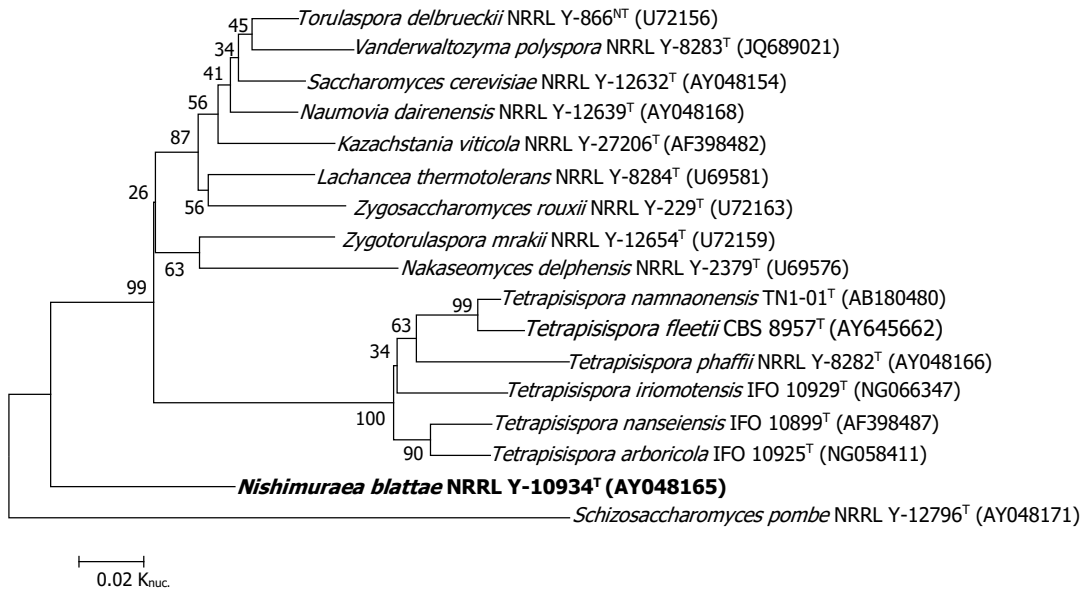


Fig. 1. The phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences with 524 bases derived from the neighbour-joining method. The numerals at the nodes of respective branches indicate bootstrap values (%) deduced from 1000 replications.

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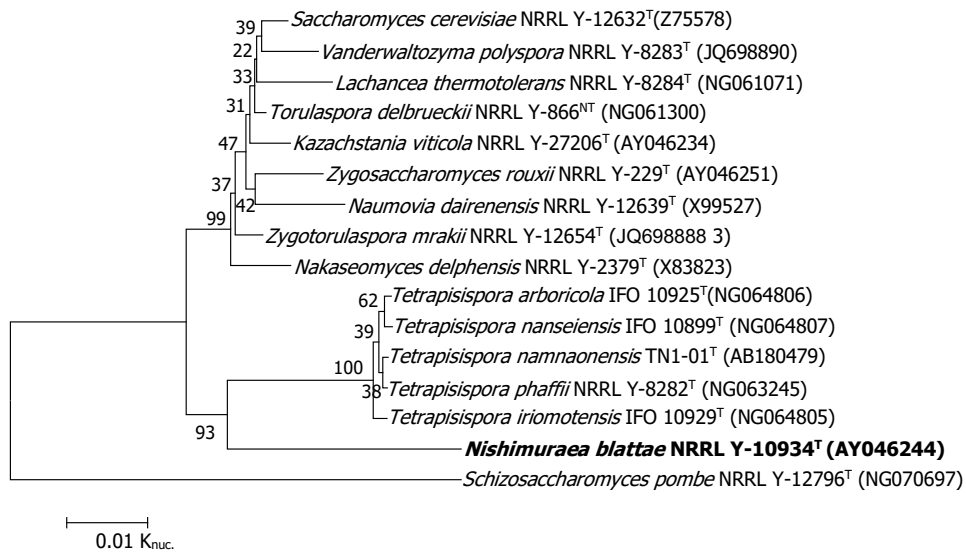


Fig. 2. The phylogenetic tree based on the 18S rRNA gene sequences with 1672 bases derived from the neighbour-joining method. The numerals at the nodes of respective branches indicate bootstrap values (%) deduced from 1000 replications.