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The Revision of Lipomycetaceae

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Abstract: In the family Lipomycetaceae, the seven genera Waltomyces, Zygozyma, Babjevia, Smithiozyma, Kawasakia, Limtongia and Kockiozyma were introduced. However, all of them were neither accepted nor recognized. This paper described the historical surveys in the systematics of the Lipomycetaceous yeasts and confirmed the total of ten genera within the family Lipomycetaceae, i.e., the genus Lipomyces sensu stricto (the type genus), the above-mentioned seven genera, the monotypic genus Dipodascopsis and the new genus Neoaidaea from the phylogenetic and the phenotypic points of view.

Keywords: Lipomyces sensu stricto; Waltomyces; Zygozyma; Babjevia; Smithiozyma; Limtongia; Kockiozyma; Dipodascopsis; Neoaidaea gen. nov.

Supplementary Abstract

The family Lipomycetaceae Novak et Zolt					
Genus	Species				
 Lipomyces Lodder et Kreger-van Rij (1952) sensu stricto 	L. starkeyi (the type species), L.				
	kononenkoae, L. tetrasporus, L.				
	mesenbrius, L. spencermartinsiae, L.				
	dorenjongii, L. kockii, L. yamadae,				
	L. yarrowii, L. orientalis, L.				
	chichibuensis, L. kalimanthanensis,				
	L. tropicalis, L. okinawaensis				
2. Dipodascopsis Batra et Millner (1978)	D. uninucleata				
Waltomyces Yamada et Nakase (1985)	W. lipofer				
4. Zygozyma van der Walt et von Arx (1987)	Z. oligophaga				
5. Babjevia van der Walt et Smith (1995)	B. anomala (the type species), B.				
	hyphoforaminiformans, B. hyphasca				
 Smithiozyma Kock, van der Walt et Yamada (1995) 	S. japonica				
7. Kawasakia Yamada et Nogawa (1995)	K. arxii				
8. Limtongia Jindamorakot, Am-in, Yukphan et Yamada	L. smithiae				
(2011)					
9. Kockiozyma Jindamorakot, Yukphan et Yamada (2012)	K. suomiensis; K. geophila f.a., K.				
	sirexii f.a., K. yamanashiensis f.a.				
 Neoaidaea Vu, Yukphan, Tanasupawat et Yamada gen. 	N. tothii				
nov.					
The family Lipomycetaceae Novak et Zolt					
Genus	Species				
1. Lipomyces Lodder et Kreger-van Rij (1952) sensu	L. starkeyi (the type species), L.				
Kurtzman, Albertyn et Basehoar-Powers (2007)	kononenkoae, L. tetrasporus, L.				
	mesenbrius, L. spencermartinsiae, L.				
	dorenjongii, L. kockii, L. yamadae,				
	L. varrowii, L. orientalis, L.				
	chichibuensis, L. kalimanthanensis,				
	L. tropicalis, L. okinawaensis (the				
	genus <i>Lipomyces</i> sensu stricto)				
	L. lipofer, L. oligophagus, L.				
	iaponicus, L. arxii, L. smithiae, L.				
	suomiensis				
	D. uninucleata (the type species), D.				
Dipodascopsis Batra et Millner (1978) emend. Kurtzman,	Di unimercuta (the type species), Di				



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In the family Lipomycetaceae Novak et Zsolt, the seven genera *Waltomyces*, *Zygozyma*, *Babjevia*, *Smithiozyma*, *Kawasakia*, *Limtongia* and *Kockiozyma* were once introduced (Yamada and Nakase 1985; van der Walt et al. 1987; Smith et al. 1995; Kock et al. 1995; Yamada and Nogawa 1995b; Jindamorakot et al. 2011, 2012). All of them were respectively taxonomic-homogeneous natured, i.e., morphologically, physiologically, chemotaxonomically, genetically and phylogenetically. However, Kurtzman et al. (2007) divided the family into two, i.e., the monophyletic genus *Lipomyces* sensu and the paraphyletic genus *Dipodascopsis* emend. instead (Smith and de Hoog 2011; Smith and Kurtzman 2011).

This paper describes the historical surveys in the systematics of the Lipomycetaceous yeasts and recovers the seven genera mentioned above as taxonomic criteria, in addition to the genus *Lipomyces* sensu stricto, the monotypic genus *Dipodascopsis* (Jindamorakot et al. 2011, 2012) and the new genus *Neoaidaea* from the phylogenetic and the phenotypic points of view. Accordingly, the family is comprised of ten genera in total as follows.

The family Lipomycetaceae Novak et Zsolt

The type genus is Lipomyces Lodder et Kreger van-Rij.

Ten genera are included.

Genus I. Lipomyces Lodder et Kreger-van Rij sensu stricto

The type species is Lipomyces starkeyi Lodder et Kreger-van Rij.

Fourteen species are included.

1. Lipomyces starkeyi Lodder et Kreger-van Rij (1952)

2. Lipomyces kononenkoae Nieuwdorp, Bos et Slooff (1974)

3. Lipomyces tetrasporus Nieuwdorp, Bos et Slooff (1974)

4. *Lipomyces mesembrius* Botha, van der Walt et Smith (1997)

5. Lipomyces spencermartinsiae (van der Walt et Smith) van der Walt et Smith (1997)

Basionym: Lipomyces kononenkoae Nieuwdorp, Bos et Slooff subsp. spencermartinsiae van der Walt et Smith (1995)

- 6. Lipomyces doorenjongii van der Walt et Smith (1999)
- 7. *Lipomyces kockii* Smith et van der Walt (1999)
- 8. Lipomyces yamadae van der Walt et Smith (1999)
- 9. Lipomyces yarrowii Smith et van der Walt (1999)
- 10. Lipomyces orientalis Thanh (2006)
- 11. Lipomyces chichibuensis Yamazaki et Kawasaki (2014)
- 12. Lipomyces kalimantanensis Kanti, Yamazaki et Kawasaki (2017)
- 13. Lipomyces tropicalis Kanti, Yamazaki et Kawasaki (2017)
- 14. Lipomyces okinawensis Yamazaki, Yanagiba et Naganuma (2017)

The species classified in the genus *Lipomyces* sensu stricto were tightly coupled with one another in all the three phylogenetic trees based on the concatenated four-gene sequences of 18S rRNA, 26S rRNA, mitochondrial small subunit rRNA and EF1-alpha derived from the neighbour-joining, the maximum parsimony and the maximum likelihood methods (Jindamorakot et al. 2012). On the other hand, the phylogenetic positions of the *Lipomyces* species newly designated by Kurtzman et al. (2007) were quite changeable in the monophyletic genus *Lipomyces* sensu (Jindamorakot et al. 2012). In fact, the resulting genus has an extremely wide range of DNA G+C contents (14.2 mol%, from 41.5 - 55.7 mol% G+C; Jindamorakot et al. 2012) and three kinds of ubiquinone isoprenologues (Q-8, Q-9 and Q-10: Yamada 1986; Yamada et al. 1986). According to Nakase and Komagata (1970) and Yamada et al. (1973), such a genus was a taxonomic heterogeneous-natured taxon, just as found in the genus *Pichia* Hansen once designated.

Genus II. Dipodascopsis Batra et Millner

Dipodascopsis uninucleata (Biggs) Batra et Millner (1978)

Basionym: Dipodascus uninucleatus Biggs (1937).

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Two varieties are described.

- 1. Dipodascopsis uninucleata var. uninucleata
- 2. Dipodascopsis uninucleata var. wickerhamii Kreger-van Rij (1974)

When the phylogenetic tree based on the 26S rRNA gene D1/D2 domain sequences was constructed by the neighbourjoining method, the paraphyletic genus *Dipodascopsis* emend. Kurtzman et al. (2007) was split into two and produced a monotypic genus (Fig. 1), as reported previously (Jindamorakot et al. 2011, 2012), indicating that *D. uninucleata*, *D. tothii* and *D. anomala* were not tightly coupled within the genus *Dipodascopsis* emend.

Such a phylogenetic data was also given by Kurtzman (2011), i.e., the phylogenetic tree based on the concatenated threegene sequences of 26S rRNA, 18S rRNA and EF1-alpha derived from the neighbour-joining method represented that D. *uninucleata* was connected first to L. *starkeyi* but not to D. *anomala* (= B. *anomala*) and then the resulting cluster was done to D. *anomala*. In fact, the calculated sequence similarity (95.7%) between D. *uninucleata* and L. *starkeyi* was much higher than that (90.4%) between D. *uninucleara* and B. *anomala* (Table 1).

Genus III. Waltomyces Yamada et Nakase (MB 25779)

Waltomyces lipofer (Lodder et Kreger-van Rij ex Slooff) Yamada et Nakase (1985) (MB 103899)

Basionym: Lipomyces lipofer Lodder et Kreger-van Rij (1952) ex Slooff (1970).

The three partial base sequencings in positions 468-622, 155 bases (designated as region a) and in positions 1611-1835, 225 bases (designated as region b) of 26S rRNA and in positions 1451-1618, 168 bases (designated as region c) of 18S rRNA showed that the Q10-equipped species, *W. lipofer* had 88% maximum homology in region a, along with *B. anomala* (= *D. anomala*), to *L. starkeyi*, *L. tetrasporus* and *L. kononenkoae* in the genus *Lipomyces* sensu stricto (Yamada and Nogawa 1995a). In contrast, the above-mentioned three *Lipomyces* species constituted a cluster with 97% or more maximum homologies within the genus, indicating that the species is accommodated to a different genus from the genus *Lipomyces* sensu stricto (Table 1). Such similar experimental data were shown, since the species was used as outgroup in the phylogenetic trees concerned (Kurtzman et al. 2007; Yamazaki and Kawasaki 2014).

Genus IV. Zygozyma van der Walt et von Arx (MB 25149)

Zygozyma oligophaga van der Walt et von Arx (1987) (MB 131108)

Synonym: Lipomyces oligophagus (van der Walt et von Arx) Kurtzman, Albertyn et Basehoar-Powers (2007).

Four *Zygozyma* species were reported (van der Walt et al. 1987, 1989b, 1990; Smith et al. 1989). However, the four were not tightly coupled with one another in the phylogenetic trees based on the concatenated four-gene sequences derived from the three methods, i.e., the neighbour-joining, the maximum parsimony and the maximum likelihood methods (Jindamorakot et al. 2012) and chemotaxonomically grouped into two based on the ubiquinone isoprenologues, i.e., Q-8 and Q-9 (Fig. 1) (Yamada and Nogawa 1995a; Jindamorakot et al. 2011, 2012). Especially, the Q8-equipped species, *Z. oligophaga* had extremely long branches (Fig. 1) (Jindamorakot et al. 2012) and quite low sequence similarities (81.6 - 86.6%) (Table 1).

Genus V. Babjevia van der Walt et Smith (MB 27435)

The type species is Babjevia anomala (Babjeva et Gorin) van der Walt et Smith.

Three species are included.

1. Babjevia anomala (Babjeva et Gorin) van der Walt et Smith (1995) (MB 363215)

Basionym: Lipomyces anomalus Babjeva et Gorin (1975).

Synonym: Dipodascopsis anomala (Babjeva et Gorin) Kurtzman, Albertyn et Basehoar-Powers (2007).

2. Babjevia hyphoforaminiformans Yamazaki, Lorliam et Kawasaki (2020) (MB 829051)

3. Babjevia hyphasca Yamazaki, Lorliam et Kawasaki (2020) (MB 829053)

Kurtzman et al. (2007) accommodated *B. anomala* to the paraphyletic genus *Dipodascopsis* emend. as *D. anomala*. However, Jindamorakot et al. (2012) insisted that the species, *D. anomala* should be transferred again to the genus *Babjevia* as *B. anomala*, since the species was not tightly coupled with the type species, *D. uninucleata* (Fig. 1). In fact, Yamazaki et

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al. (2020) did not classified two new species to the genus *Dipodascopsis* emend. but the genus *Babjevia* as already suggested by Jindamorakot et al. (2012), indicating that the generic concept of Kurtzman et al. (2007) completely lost its validity.

In the phylognetic tree based on the concatenated four-gene sequences derived from the maximum likelihood method (Yamazaki et al. 2020), *L. lipofer*, *L. smithiae*, *Zygozyma arxii* (= *L. arxii*), *L. japonicus*, *L. oligophaga* [sic] and *Zygozyma suomiensis* (= *L. suomiensis*) were preferable to be improved to *Waltomyces lipofer*, *Limtongia smithiae*, *Kawasakia arxii*, *Smithiozyma japonica*, *Zigozyma oligophaga* and *Kockiozyma suomiensis* (Jindamorakot et al. 2011, 2012).

Genus VI. Smithiozyma Kock, van der Walt et Yamada (MB 27638)

Smithiozyma japonica (van der Walt, Smith, Yamada et Nakase) Kock, van der Walt et Yamada (1995) (MB 414161)

Basionym: Lipomyces japonicus van der Walt, Smith, Yamada et Nakase (1989).

In the three partial base sequencings mentioned above, the Q9-equipped species, *S. japonica* (= *L. japonicus*; van der Walt et al. 1989a) represented 89% maximum homology in region a, two base differences in region b and five base differences in region c to *L. starkeyi*, *L. tetrasporus* and *L. kononenkoae* in the genus *Lipomyces* sensu stricto. In contrast, the abovementioned three *Lipomyces* species constituted clusters with 97% or more maximum homologies (region a) and no base differences (regions b and c) within the genus (Yamada and Nogawa 1995a), indicating that the species is accommodated taxonomically to a different genus from the genus *Lipomyces* sensu stricto (Table 1).

In the phylogenetic trees based on the concatenated four-gene sequences, *S. japonica* was not tightly coupled, i.e., the species was connected to the cluster comprised of the Q9-equipped species including *L. starkeyi*, *D. uninucleata* and so on, when constructed by the neighbour-joining method, but to the cluster of Q8-equipped species including *Z. oligophaga* and *Kockiozyma suomiensis*, when constructed by the maximum parsimony method (Jindamorakot et al. 2012).

Genus VII. Kawasakia Yamada et Nogawa (MB 27831)

Kawasakia arxii (van der Walt, Smith et Yamada) Yamada et Nogawa (1995) (MB 442988)

Basionym: Zygozyma arxii van der Walt, Smith et Yamada (1989).

Synonym: Lipomyces arxii (van der Walt, Smith et Yamada) Kurtzman, Albertyn et Basehoar-Powers (2007).

The Q9-equipped species, *Kawasakia arxii* was very unique in representing considerably large base differences (six) from the above-mentioned three *Lipomyces* species in the partial base sequencing of 18S rRNA (region c), in contrast to *W. lipofer*, which had only one base difference in the same region (Yamada and Nogawa 1995a, b). In the remaining regions a and b, the maximum homology and the base differences of the species were 93% and only one. The calculated 26S rRNA gene D1/D2 domain sequence similarities of *Kawasakia arxii* were 85.7 - 97.5% (Table 1). In the phylogenetic trees based on the concatenated four-gene sequences, the phylogenetic positions of *Kawasakia arxii* were considerably changeable, indicating that the species was not tightly coupled to any of other species (Jindamorakot et al. 2012).

Genus VIII. Limtongia Jindamorakot, Am-in, Yukphan et Yamada (MB 582716)

Limtongia smithiae (van der Walt, Wingfield et Yamada) Jindamorakot, Am-in, Yukphan et Yamada (2011) (MB 582717)

Basionym: Zygozyma smithiae van der Walt, Wingfield et Yamada (1990).

Synonym: Lipomyces smithiae (van der Walt, Wingfield et Yamada) Kurtzman, Albertyn et Basehoar-Powers (2007).

In the three partial base sequencings mentioned above, the Q9-equipped species, *Limtongia smithiae* represented 89% maximum homology, four base differences and three base differences respectively to *L. starkeyi*, *L. tetrasporus* and *L. kononenkoae* in the genus *Lipomyces* sensu stricto (Yamada and Nogawa 1995a). The calculated 26S rRNA gene D1/D2 domain sequence similarities of the species were 82.9 - 92.4%, indicating that the species is accommodated to an independent genus (Table 1).

Genus IX. Kockiozyma Jindamorakot, Yukphan et Yamada (MB 587737)

1. Kockiozyma suomiensis (Smith, van der Walt et Yamada) Jindamorakot, Yukphan et Yamada (2012) (MB 587754)

Basionym: Zygozyma suomiensis Smith, van der Walt et Yamada (1990).

Synonym: Lipomyces suomiensis (Smith, van der Walt et Yamada) Kurtzman, Albertyn et Basehoar-Powers (2007).

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2. Kockiozyma geophila f.a. (van der Walt, Yamada et Nakase) comb. nov.

Basionym: Myxozyma geophila van der Walt, Yamada et Nakase, Syst. Appl. Microbiol. 9: 122, 1987.

The type strain is CBS 7219 = NRRL Y-17252.

3. Kockiozyma sirexii f.a. (Spaaij et Weber) comb. nov.

Basionym: Myxozyma sirexii Spaaij et Weber, Syst. Appl. Microbiol. 15: 429, 1992.

The type strain is UOFS Y-2054 = NRRL Y-27626

4. Kockiozyma yamanashiensis f.a. (Yamazaki, Yanagiba et Naganuma) comb. nov.

Basionym: Lipomyces yamanashiensis f.a. Yamazaki, Yanagiba et Naganuma, Int. J. Syst. Evol. Microbiol. 67: 2945, 2017.

The type strain is NBRC 110621 = CBS 14748.

In the three partial base sequencings mentioned above, the Q8-equipped species, *Kockiozyma suomiensis* represented 83% maximum homology in region a, 12 base differences in region b, and four base differences in region c respectively to *L. starkeyi*, *L. tetrasporus*, and *L. kononenkoae* in the genus *Lipomyces* sensu stricto (Yamada and Nogawa, 1995a). The calculated 26S rRNA gene D1/D2 domain sequence similarities of the species were 85.9 - 90.6%, indicating that the species is accommodated to an independent genus (Table 1). On the other hand, *Myxozyma geophila*, the Q8-equipped anamorphic species showed 92% maximum homology, and one and one base difference to *Kockiozyma suomiensis* in the three partial base sequencings (Yamada and Nogawa, 1995a).

In addition, the sequence similarity between *Kockiozyma suomiensis* and *Myxozyma geophila* was 98.0%, the value of which was almost identical with that (98.1%) between *Octosporomyces octosporus* and *Octosporomyces osmophilus* (Malimas et al. 2023). From the results obtained above, the generic name of *Kockiozyma* was able to be given to the anamorphic species as *Kockiozyma geophila* f.a. (Lachance 2012) as well as *Kockiozyma melibiosi* f.a., *Kockiozyma mucilagina* f.a., *Kockiozyma neotropica* f.a., *Kockiozyma vanderwalti* f.a. and *Kockiozyma neglecta* f.a. (Vu et al. 2022).

Dipodascopsis tothii and *B. anomala* (= *D. anomala*) were tightly coupled in the three phylogenetic trees (Jindamorakot et al. 2012), so that the two species appeared to be classified in a single genus (Fig. 1). Due to the priority of *B. anomala*, the single genus was to be named *Babjevia*. However, the distance between the two species, *D. tothii* and *B. anomala* was not so short, when compared with *Kluyveromyces polysporus* (= *Vanderwaltozyma polyspora*; Kurtzman 2003) and *Saccharomyces cerevisiae* (Jindamorakot et al. 2012). In fact, the calculated pair-wise sequence similarity (95.6%) between the two species was interestingly almost the same as that (95.7%) between *D. uninucleata* and *L. starkeyi* and that (95.2%) between *W. lipofer* and *L. starkeyi*, and not so high to accommodate the two species to the single genus *Babjevia* (Table 1), indicating that an additional genus can be phylogenetically introduced for *D. tothii*. Phenotypically, the species was quite different from *B. anomala*, e.g., in the morphology of asci and ascospores and the mode of asexual reproduction (Smith and de Hoog 2011).

Yamazaki et al. (2017a, b) constructed the phylogenetic trees based on the translation elongation factor 1-alpha (EF1-alpha) gene sequences derived from the neighbour-joining method. In the phylogenetic trees, it is of interest that *D. tothii* was found inside and distantly separated from *B. anomala* as well as *D. uninucleata*. In contrast, the cluster including *L. starkeyi* and its relatives and the cluster including *Kockiozyma suomiensis* and its related *Myxozyma* species were tightly coupled without any exception. In this respect, *D. tothii* was quite different phylogenetically from the remaining two species of the paraphyletic genus *Dipodascopsis* emend. Kurtzman et al. (2007), and thus the introduction of a separate new genus was confirmed.

Genus X. Neoaidaea Vu, Yukphan, Tanasupawat et Yamada gen. nov.

Neoaidaea (Ne.o.a.i'da.e.a. N. L. fem. n. *Neoaidaea*, new Aida, in honour of Dr. Ko Aida, Professor Emeritus, The Institute of Applied Microbiology, The University of Tokyo, Tokyo, Japan, who introduced the isoprenoid quinone analyses into the microbial systematics.

The colonies are cream-coloured, moist and creamy after 10 d (Smith and de Hoog 2011). Budding cells are absent. Hyphae

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are firm with inflated cells and intensely amyloid. Arthroconidia are absent. Gametangia are formed laterally on hyphae as curved branches. Asci arise after fusion of a terminal cell with its penultimate cell. Asci are acicular, 50 - 100 \Box m long and contain 30 - 100 ascospores. Asci open by rupturing at the apex. Ascospores are subhyaline to hyaline, broadly ellipsoidal, occasionally bean-shaped, 1×1.5 - 2 \Box m and without slime. Fermentation is absent. Grows on glucose, inulin, sucrose, raffinose, galactose, lactose, trehalose, maltose, methyl- \Box -glucoside, starch, cellobiose, salicin, L-sorbose, xylose, arabinose, L-arabinose, ethanol, glycerol, ribitol, mannitol, glucitol, *myo*-inositol, succinate (w) and gluconate (v), but not on melibiose, L-rhamnose, ribose, methanol, erythritol, galactitol and DL-lactate. No growth on vitamin-free medium (Smith and de Hoog 2011). Ubiquinone-9 (Q-9) is present (Cottrell and Kock 1989).

Mycobank number is 846280.

Neoaidaea tothii (Zolt) Vu, Yukphan, Tanasupawat et Yamada comb. nov.

Basionym: Dipodascus tothii Zolt, Acta Bot. Hung. 9: 226, 1963.

Synonym: Dipodascopsis tothii (Zolt) Batra et Millner (1978).

The characteristics of the species are the same as those described in the genus (Smith and de Hoog 2011).

The holotype is CBS $759.85^{T} = NBRC \ 10813^{T}$.

MycoBank number is 846281.

For the detailed characteristics of the taxa mentioned above and the type strains concerned, refer to the related monographs and/or articles.

Kurtzman (2003) introduced 'clade' or 'phylogenetic circumscribed genus' as generic concept. However, it was not yet perfect to get the taxonomic homogeneous-natured taxon for the genus. Namely, the branch lengths should be additionally considered in the phylogenetic trees.

In the seven ranks used for the systematics of plants and animals, the homogeneous-natured taxa can be available only when the lower-ranked two taxa, i.e., species and genus are dealt with taxonomically. Therefore, the present authors naturally selected the homogeneous-natured taxon in the generic designation, since the longer the phylogenetic distances are the more taxonomic heterogeneities will be increased.

Thus, the monophyletic genus *Lipomyces* Lodder et Kreger-van Rij sensu Kurtzman et al. (2007) included the phylogeneticdistant and the phenotypic-different species, i.e., *L. lipofer*, *L. oligophagus*, *L. japonicus*, *L. arxii*, *L. smithiae* and *L. suomiensis*, all of which were re-classified as *Waltomyces lipofer*, *Zygozyma oligophaga*, *Smithiozyma japonica*, *Kawasakia arxii*, *Limtongia smithiae* and *Kockiozyma suomiensis* respectively, in addition to *L. starkeyi*, the type species and its closely related *Lipomyces* species that constituted the genus *Lipomyces* sensu stricto (Jindamorakot et al. 2011, 2012), and so corresponded taxonomically to the so-called family.

On the other hand, the paraphyletic genus *Dipodascopsis* Batra et Millner emend. Kurtzman et al. (2007) was a monotypic genus, which included only *D. uninucleata* (Jindamorakot et al. 2012). For the remaining two species, one was transferred again to the genus *Babjevia* as *B. anomala* (Jindamorakot et al. 2012; Yamazaki et al. 2020) and the other was classified in the new genus *Neoaidaea* as *Neoaidaea tothii* gen. nov., comb. nov.

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Conflicts of interest

The authors declare that there are no conflicts of interest.

Author contributions

Y.Y., H.T.L.V., P.Y. and S.T. designed the study. H.T.L.V. performed the main experiments. P.Y. instructed how to make the experiments. Y.Y. prepared the manuscript. The detailed discussion was made among Y.Y., H.T.L.V., P.Y. and S.T.

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This article is dedicated to Professor Dr. Johannes Petrus van der Walt, South Africa for his excellent taxonomic study on the Lipomycetaceous yeasts, especially in the findings of the Q8-equipped Zygozyma oligophaga and its related species.

Preliminary reports were opened [19, 27]

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	Sequence similarity (%)				_						
Species											
	1	2	3	4	5	6	7	8	9	10	11
1. Lipomyces starkeyi											
2. Waltomyces lipofer	95.2										
3. Zygozyma oligophaga	85.5	83.0									
4. Babjevia anomala	91.7	91.0	81.6								
5. Smithiozyma japonica	96.4	93.6	85.5	89.9							
6. Kawasakia arxii	97.5	94.1	85.7	92.2	95.7						
7. Limtongia smithiae	92.4	89.9	82.9	86.8	90.6	91.0					
8. Kockiozyma suomiensis	90.6	88.9	85.9	86.2	90.3	90.6	86.4				
9. Myxozyma geophila	90.4	89.0	86.6	85.9	90.4	90.3	85.7	98.0			
10. D. uninucleata var. uninucleata	95.7	94.0	85.5	90.4	95.0	95.4	91.2	90.2	90.2		
11. D. uninucleata var. wickerhamii	95.6	93.8	85.4	90.3	94.9	95.2	91.0	90.1	90.1	99.8	
12. Neoaidaea tothii	92.6	90.5	83.4	95.6	90.1	95.2	87.8	87.8	86.7	90.6	90.5

The pair-wise 26S rRNA gene D1/D2 domain sequence similarities were calculated for 555-568 bases with the program BioEdit (version 7.2.5) (Hall, *GERF Bull Biosci* 2011; **2**: 60,).

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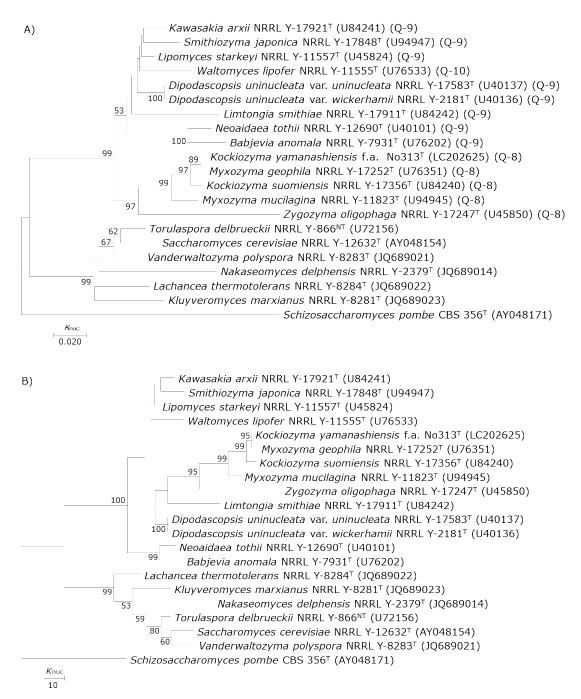


Fig. 1. The phylogenetic relationships of the Lipomycetaceous yeast species. The phylogenetic trees based on the 26S rRNA gene D1/D2 domain sequences (526 bases) were constructed for the Lipomycetaceous yeast species by the neighbourjoining method (A) and by the maximum parsimony method (B). Saccharomyces cerevisiae and its related species were utilized for reference standards. Schizosaccharomyces pombe was used as outgroup. The evolutionary distances were computed using the Kimura 2-parameter method (Kimura, J Mol Evol 1980; 16: 111). The numerals at the nodes of the respective branches indicate bootstrap values (%) deduced from 1000 replications (Felsenstein, Evolution 1985; 39: 783). The bootstrap values below 50% were deleted. For the additional phylogenetic trees based on the concatenated four-gene sequences, i.e., 18S rRNA, 26S rRNA, mitochondrial small subunit rRNA, and EF1-alpha, derived from the three methods, refer to Jindamorakot et al. (2012). According to the theory of Kurtzman et al. (2007), the Lipomycetaceous yeast species are designated as *L. starkeyi* and its relatives, *Lipomyces lipofer*, *L. oligophagus*, *L. japonicus*, *L. smithiae*, *L. arxii*, *L. suomiensis*, *Dipodascopsis uninucleata*, *D. anomala* and *D. tothii* instead.