Kurtzmaniella gen. nov. and description of the heterothallic, haplontic yeast species Kurtzmaniella cleridarum sp. nov., the teleomorph of Candida cleridarum

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The teleomorph of Candida cleridarum was discovered through the detection of conjugation between isolates of a large collection from the nitidulid beetles of the genus Carphophilus found in the flowers of various cacti in Arizona, USA. The previous oversight of the sexual cycle of this yeast is attributed to the inequality (ca. 5 : 1) of the two mating types. Extensive conjugation between compatible mating types is observed after overnight incubation on 5 % malt agar, followed after 3–5 days by the formation of mature ascii. The hat-shaped ascospores are reminiscent of those seen in Kodamaea species, which are members of the same guild. However, published analyses of D1/D2 large subunit rDNA sequences indicate an affinity with the genus Debaryomyces. As the latter is polyphyletic and morphologically heterogeneous, and in view of the distinct life cycle of the new teleomorph, the new genus Kurtzmaniella is described with a novel species, Kurtzmaniella cleridarum sp. nov. Given the close relatedness of Kurtzmaniella cleridarum sp. nov. to Candida quercitrusa, Candida oleophila and Candida railenensis, for which several natural isolates were available, strains of these species were mixed in pairs under the conditions found favourable for the former. Conjugation was not detected in those species. The type strain of Kurtzmaniella cleridarum sp. nov. is UWOPS 99-101.1T (=CBS 8793T=NRRL Y-48386T, h⁺), type of Candida cleridarum. The allotype is UWOPS 07-123.1 (=CBS 10688=NRRL Y-48387, h⁺).

Candida cleridarum was described by Lachance et al. (2001) from a collection of strains recovered from flowers of cacti of the deserts of Arizona and California, USA, or from beetles that occur in the flowers. The species was named after the clerid beetles found in some of the flowers, but it was later found (Lachance & Bowles, 2002) that the major vector of yeasts in these flowers is the nitidulid genus Carphophilus. Carphophilus pallipennis is known to occur consistently in the flowers of various cacti in Arizona, USA. The previous oversight of the sexual cycle of this yeast is attributed to the inequality (ca. 5 : 1) of the two mating types. Extensive conjugation between compatible mating types is observed after overnight incubation on 5 % malt agar, followed after 3–5 days by the formation of mature ascii. The hat-shaped ascospores are reminiscent of those seen in Kodamaea species, which are members of the same guild. However, published analyses of D1/D2 large subunit rDNA sequences indicate an affinity with the genus Debaryomyces. As the latter is polyphyletic and morphologically heterogeneous, and in view of the distinct life cycle of the new teleomorph, the new genus Kurtzmaniella is described with a novel species, Kurtzmaniella cleridarum sp. nov. Given the close relatedness of Kurtzmaniella cleridarum sp. nov. to Candida quercitrusa, Candida oleophila and Candida railenensis, for which several natural isolates were available, strains of these species were mixed in pairs under the conditions found favourable for the former. Conjugation was not detected in those species. The type strain of Kurtzmaniella cleridarum sp. nov. is UWOPS 99-101.1T (=CBS 8793T=NRRL Y-48386T, h⁺), type of Candida cleridarum. The allotype is UWOPS 07-123.1 (=CBS 10688=NRRL Y-48387, h⁺).

The nature of the association is not known, but one of the reasons for the specificity may well be the volatile substances released by yeasts as they ferment plant sugars. Yeast fermentation products can act as strong attractants for Carphophilus species that are pests of certain crop plants (Nout & Bartelt, 1998) and may be important in preserving beetle–yeast associations.

In three previous collections of yeasts found in cactus flowers (Lachance et al., 2001; Lachance & Bowles, 2002; and unpublished data), Candida cleridarum was clearly dominant across the entire collection area, which included sites in southern Arizona and southern California. The exception was a few samples collected in two localities near Phoenix, Arizona, where a small number of isolates of the endemic species Metschnikowia arizonensis were also found. With the objective of gaining new insights into the basis of these distributions, we recently made an intense collection of the yeasts associated with Carphophilus spp. in the flowers of cacti in five Arizona localities. Of the 107 yeast strains recovered, 102 were identified as Candida spp. in the flowers of cacti in five Arizona localities. Of the 107 yeast strains recovered, 102 were identified as Candida spp. in the flowers of cacti in five Arizona localities. Of the 107 yeast strains recovered, 102 were identified as Candida spp. in the flowers of cacti in five Arizona localities. Of the 107 yeast strains recovered, 102 were identified as Candida spp. in the flowers of cacti in five Arizona localities.
among strains of *Candida cleridarum*, which led us to show, through mating experiments, that they belong to a single ascosporic species.

**Isolation of conjugating strains**

Table 1 details the origin of recent isolates plus ten isolates that were retained from a collection conducted in 1999 (Lachance et al., 2001). The sites of the recent collection were visited in April 2007, at which time flowering was extensive in flat and cylindrical *Opuntia* species as well as *Echinocereus* species. Insects were removed aseptically from flowers and were allowed to walk for 10–20 min on the surface of agar plates. The yeasts deposited on the agar surface were then spread with an inoculation loop. To maximize the probability of detecting mating types of *M. arizonensis*, insects were allowed to walk on two different agar media. As a general isolation medium, we used YM agar (1.0% glucose, 0.5% peptone, 0.3% malt extract, 0.3% yeast extract and 2.0% agar) supplemented with 100 mg chloramphenicol l⁻¹. The second medium additionally contained (1⁻¹), 10 mg cetyltrimethylammonium bromide (CTAB) and 5 mg phloxine B. CTAB, a toxic detergent, has been used in this laboratory as a selective agent for species of the genus *Metschnikowia* (Lachance et al., 2003a). It happens that *Candida cleridarum* is also resistant to CTAB (Lachance et al., 2001). Phloxine B is a vital stain that has been used, mostly by geneticists, for the detection of colonies formed by physiological variants (Nagai, 1963). It would seem that the dye preferentially penetrates cells that have entered G₁ arrest. We have found it useful, in previous studies, for the on-site detection of interacting mating types of heterothallic *Metschnikowia* species on plates inoculated with floricolous beetle yeasts. When two *Metschnikowia* sp. colonies of complementary mating types grow in contact with each other, one of the colonies often picks up and concentrates the red colour. In the present study, we noticed that some confluent colonies of *Candida cleridarum* formed an intensely coloured zone at the point of contact. We later confirmed that the dye was concentrated at the point where cells were conjugating, although microscopic examination showed that the conjugants themselves did not absorb significant quantities of the dye, in contrast to dead cells, which appeared bright red.

**Mating type distribution**

The mating type *h*⁺, arbitrarily assigned to the type strain of *Candida cleridarum* UWOPS 99-101.1, predominated in all localities with a fivefold excess over mating type *h*⁻. Of the strains listed in the original description of *Candida cleridarum* (Lachance et al., 2001), ten were preserved in the UWOPS culture collection. Of these, only one (UWOPS 99-117.1) had the mating type *h*⁻ (Table 1). This explains in part why conjugation had been overlooked in previous collections. Another reason was the choice of mating medium. When examining mixtures of strains for possible conjugation and ascus formation, we normally use, by default, YCBAS agar (yeast carbon base, Difco, with 0.01% ammonium sulphate). This medium has generally given excellent results with heterothallic, haplontic species of the genera *Clavispora*, *Metschnikowia*, *Kodamaea*, *Starmerella* and *Wickerhamiella*, although there have been exceptions (Lachance et al., 2000). In the present case, given that the interactions were detected from the accumulation of phloxine B occurred on YM agar, a rich medium, we compared the efficacy of several media, including YCBS, YM, 20-fold diluted V8 and 5% and 10% malt agar. The last medium gave markedly better results and YCBAS agar was particularly poor.

**Characterization of new isolates**

The growth characteristics of the yeasts were determined following standard methods (Yarrow, 1998). The isolates identified as *Candida cleridarum* matched the original description well, with some minor variations that may represent artefacts of media preparation. Specifically, most strains gave a slow and weak reaction for gelatin hydrolysis, which was previously reported as negative. Two strains that were picked on the basis of the formation of smaller colonies on the medium containing CTAB had slightly different growth patterns. Strain UWOPS 07-101a1 (*h*⁺) exhibited no growth at all on sucrose, maltose or melezitose, although other growth responses were normal. Strain UWOPS 07-131.2 (*h*⁻) grew weakly on most media. Growth on citrate varied markedly across all strains.

**Kurtzmaniella gen. nov.**

The discovery of ascus formation in *Candida cleridarum* requires assignment to a teleomorphic genus. The current

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**Table 1.** Origin and mating types of isolates of *Kurtzmaniella cleridarum* sp. nov.

The 1999 collection has been described in detail by Lachance et al. (2001).

<table>
<thead>
<tr>
<th>Locality</th>
<th>Year</th>
<th>Mating type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lost Dutchman State Park</td>
<td>1999</td>
<td>2</td>
</tr>
<tr>
<td>Lost Dutchman State Park</td>
<td>2007</td>
<td>19  7</td>
</tr>
<tr>
<td>Pinahighway</td>
<td>1999</td>
<td>2</td>
</tr>
<tr>
<td>Boyce Thompson Arboretum</td>
<td>2007</td>
<td>4  2</td>
</tr>
<tr>
<td>Saguaro National Park East</td>
<td>2007</td>
<td>19  2</td>
</tr>
<tr>
<td>Saguaro National Park West</td>
<td>2007</td>
<td>20  2</td>
</tr>
<tr>
<td>Gilbert Ray Campground</td>
<td>1999</td>
<td>2</td>
</tr>
<tr>
<td>Brown Canyon, Coronado National Forest</td>
<td>2007</td>
<td>11  3</td>
</tr>
<tr>
<td>Organ Pipe Cactus National Monument</td>
<td>1999</td>
<td>1</td>
</tr>
<tr>
<td>Organ Pipe Cactus National Monument</td>
<td>2007</td>
<td>11  2</td>
</tr>
<tr>
<td>Joshua Tree National Park, California</td>
<td>1999</td>
<td>1  1</td>
</tr>
<tr>
<td>Amboy, California</td>
<td>1999</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>93</td>
<td>19</td>
</tr>
</tbody>
</table>
practice is to attempt the maintenance of genera that are monophyletic as inferred by DNA sequence analyses, as well as morphologically consistent. The present case is difficult in this respect. The closest relative of Candida fragi, as shown in Fig. 1. This and other analyses (Kurtzman & Robnett, 1998; Lachance et al., 2001; Kurtzman et al., 2001) showed that these species form a clade with a few other Candida species whose closest known ascosporic relatives are in the genus Debaryomyces. However, the latter genus has representatives in three clades. The clade nearest to Candida cleridarum and close relatives comprises, among others, Debaryomyces hansenii, which forms automictic asci with usually a single rough-walled ascospore, and Debaryomyces (Wingea) robertsi, with characteristic lenticular ascospores that come singly or in pairs. The second nearest clade contains automatic species that include Debaryomyces (Schwanniomyces) occidentalis, with mostly a single rugose, ledged ascospore, and Debaryomyces polymorphus, with two or more smooth, spheroidal ascospores. A third, more distant clade does not form a monophyletic set with the others and contains a mixture of Debaryomyces species and certain Picha species that form spheroidal ascospores without a ledge. A multi-gene study in progress (C. P. Kurtzman, personal communication) confirms this phylogenetic structure. None of the known species of the genus Debaryomyces produce pairs of hat-shaped ascospores by conjugation of heterothallic strains. In view of all the above considerations, we think it justified to erect a separate genus to accommodate the teleomorph of Candida cleridarum and eventually those of neighbouring species. We propose to name this genus Kurtzmaniella gen. nov., in honour of Cletus P. Kurtzman, in recognition of his innumerable contributions to yeast systematics. Most noteworthy are Kurtzman’s leadership in the work that led to the first and still only operational molecular barcoding system for yeast identification and his relentless efforts to provide phylogenetic circumscription of yeast genera through multi-gene sequence analysis.

Search for other Kurtzmaniella species

In view of the close relatedness of Kurtzmaniella cleridarum sp. nov. to Candida quercitrusa, Candida oleophila and Candida railenensis (Fig. 1), for which several natural isolates were available in our collection, strains of these species, including the type cultures, were mixed in every possible pair under the conditions found favourable for K. cleridarum. Conjugation was not detected for 13 strains of Candida oleophila, 9 strains of Candida quercitrusa or 12 strains of Candida railenensis. Although Candida fragi is known from only a single strain, it differs from K. cleridarum by only five substitutions, each in the ITS/5.8S (GenBank accession nos AY344066 and AY344065) and the D1/D2 LSU (U71071 and AF251552) rDNA sequences, respectively, leaving open the possibility that the two taxa might represent variants of a single polymorphic species (Lachance et al., 2003b). The type strain of Candida fragi (CBS 7702T) was therefore mixed with the two mating types of K. cleridarum and examined for signs of a mating reaction. No interactions of any kind were detected. Furthermore, profound differences between the two species at the cellular and colony level were noted. K. cleridarum forms larger, mostly ovoidal cells and smooth colonies in contrast to Candida fragi where the cells are smaller, elongate, and angular, with dull and rugose colonies.

The Vienna edition of the International Code of Botanical Nomenclature (McNeill, 2006) now allows the epitypification of species in an anamorphic genus if a teleomorph is discovered for one species in the genus. However, it is not clear how the relevant Article (59.7) would apply in the present circumstance, as the species most closely related to K. cleridarum are part of a large polyphyletic genus (Candida) in which most species are not closely related. In the absence of objective evidence such as interspecific mating, the transfer of neighbouring species to Kurtzmaniella would require making a typological decision as to which additional species to include: only Candida fragi or other species in neighbouring subclades. This would run the risk that new information would cause the later transfer of some of the species to yet another genus, an option that we do not view favourably.

Ecology and biogeography

Lachance et al. (2001) commented on the extraordinary habitat specificity of Candida cleridarum. The isolation of over a hundred more representatives confirms this
remarkable distribution. In addition, the fact that each of the five sites sampled yielded both mating types in approximately similar (5:1) ratios and the absence of any local peculiarities in growth responses (i.e. the aforementioned variation in citrate utilization is randomly distributed) suggest that the species may constitute a panmictic population across its known range. Further studies will be required to test this hypothesis. We also hope to study mating type inheritance to see if the evolutionary fitness of $h^-$ strains might be less that that of $h^+$ strains.

**Physiological convergence in the floricolous beetle yeast guild**

Another striking feature emerging from our studies of yeasts associated with floricolous beetles is the similarity of the dominant species in their growth responses, irrespective of phylogenetic affinity. The nutritional profile of *Kurtzmaniella cleridarum* is remarkably similar to that of many guild members that belong to the *Metschnikowia* and *Kodamaea* clades. Specifically, these yeasts have a very strong propensity for assimilating sucrose, galactose, maltose, melezitose, sorbose, xylose, mannitol, glucitol, succinic and citric acids, 2-keto-D-gluconic acid and N-acetyl-D-glucosamine, but not L-rhamnose or D-ribose. Suitable nitrogen sources include ethylamine, cadaverine and usually lysine, but not nitrate or nitrite. Tween 80 is often hydrolysed. All species are fermentative and sensitive to cycloheximide (100 mg l$^{-1}$) and most are resistant to CTAB (10 mg l$^{-1}$ or more). Whereas the possession of any one of these features is far from exceptional, the convergence of so many characteristics is improbable and suggests that these traits might jointly define the fundamental niche of floricolous beetle-associated yeasts.

**Latin diagnosis of *Kurtzmaniella cleridarum* sp. nov.**


**Description of *Kurtzmaniella cleridarum* sp. nov.**

*Kurtzmaniella cleridarum* (cle.ri.da’rum. N.L. gen. pl. fem. n. *cleridarum* of Cleridae, referring to the family of beetles that was originally thought to act as vectors for this yeast species).

The description is identical to that given for *Candida cleridarum* by Lachance et al. (2001) except for the following variations. Conjugation between haploid strains of compatible mating types is observed after 1 day at 25 °C. Mature asci containing usually two hat shaped ascospores, 2–2.5 × 3–4 μm occur after 3–5 days (Fig. 2). The lobe containing the ascospores gradually deliquesces over several more days. Ascospore formation is abundant on 5 % malt extract agar, weak on 1 % malt or 20-fold dilute V8 agar, and rare or absent on YCBS. After 3 weeks on malt gelatin, liquefaction is weak to moderate. The habitat...
is in flowers of cacti and associated beetles of the genus *Carpophilus* in the south-west United States.

The type strain of *Kurtzmaniella cleridarum* is strain UWOPS 99-101.1^T^, the type of *Candida cleridarum*. It was isolated from a beetle collected from a flower of *Opuntia phaeacantha* in Lost Dutchman State Park, Arizona, USA. The designated allotype is strain UWOPS 07-123.1 recovered from a specimen of *Carpophilus* sp. found in a flower of *Opuntia engelmannii*, in Brown Canyon, Coronado National Forest, Arizona, USA. The strains have been deposited in the collection of the Yeast Division of the Centraalbureau voor Schimmelcultures, Utrecht, the Netherlands, as strains CBS 8793^T^ (=NRRL Y-48386^T^) and CBS 10688 (=NRRL Y-48387). The binomial is registered under Mycobank number MB511088.

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**References**


