Two New and Closely Related Heterothallic Species, *Pichia amylophila* and *Pichia mississippiensis*: Characterization by Hybridization and Deoxyribonucleic Acid Reassociation

C. P. KURTZMAN,† M. J. SMILEY,† C. J. JOHNSON,† L. J. WICKERHAM,† AND G. B. FUSON‡

Northern Regional Research Center, Agricultural Research, Science and Education Administration, U.S. Department of Agriculture, Peoria, Illinois 61604,† and Department of Food Science and Technology, University of California, Davis, California 95616‡

Two new species of the yeast genus *Pichia* were isolated from frass obtained from loblolly pines growing in Mississippi. The new taxa, designated *Pichia amylophila* and *Pichia mississippiensis*, are heterothallic, and *Candida obtusa* var. *arabinosa* proved to be a haploid member of the latter species. The species show interspecific mating, but ascospores from the crosses are infertile. The nuclear deoxyribonucleic acids from the species differ by 2 mol% in guanine plus cytosine content and exhibit low (25%) base sequence complementarity. Deoxyribonucleic acid relatedness was determined spectrophotometrically, and the methodology is given in detail.

During the isolation of yeasts from insect frass collected from pine trees in Mississippi, we recovered strains that were phenotypically similar to *Pichia wickerhamii* (van der Walt) Kregervan Rij and *Pichia rhodanensis* (Ramirez et Boidin) Phaff. These strains were heterothallic but failed to conjugate with mating types from either *P. wickerhamii* or *P. rhodanensis*. One of the newly isolated cultures differed from the others by its ability to assimilate starch, and genetic crosses between it and the other isolates gave only infertile and generally poorly formed ascospores. In an effort to clarify speciation, we examined the nuclear deoxyribonucleic acid (DNA) base sequence complementarity of the strains. As a result, the isolates proved to be of particular interest because they allowed an examination of species parameters through both genetic crosses and DNA reassociation studies.

**MATERIALS AND METHODS**

**Culture isolation.** Frass samples from shortleaf (*Pinus echinata* Mill.) and loblolly (*Pinus taeda* L.) pines were collected at the Piney Woods Country Life School of Piney Woods, Miss., in late October 1949, and sent by school personnel to the Northern Regional Research Center. Samples were directly streaked onto YM agar in petri plates and were enriched by incubation in liquid yeast nitrogen base supplemented with either glucose or rhamnose (21). Cultures were incubated at 25°C. Resultant yeast colonies were isolated and further purified by streaking on YM agar.

**Physiological and morphological characterization.** The methods for the morphological examination and characterization by fermentation and assimilation tests were previously given (21). Single-ascospore isolates were obtained by micromanipulation. The criteria used for speciation within the genus *Pichia* were those proposed by Kregervan Rij (7).

**DNA purification and determination of base composition.** Extraction and purification of DNA were done by a combination of the procedures of Marmur (9) and Bernardi et al. (2) as described by Price et al. (15). The ratios of absorbance at 260 nm (*A*~260~/*A*~230~ and *A*~260~/*A*~280~) were used to assess DNA purity. Purification was repeated if the preparation deviated more than 0.05 from the ratios *A*~260~/*A*~280~ = 1.86 and *A*~230~/*A*~280~ = 0.5 (11). The quality of the DNA was further assessed from analytical ultracentrifuge scans and from thermal-melt profiles. Hyperchromicity of the preparations ranged from 34.4 to 36.8%.

The guanine plus cytosine (G+C) content of the nuclear DNA was calculated from buoyant density values in cesium chloride (16, 19) and was based on three or four separate determinations made with a Spinco model E analytical ultracentrifuge equipped with an electronic scanner. *Micrococcus lysodeikticus* Fleming DNA was used as a reference; this DNA had a buoyant density of 1.7311 g/ml when compared to DNA from *Escherichia coli* K-12, whose density was taken to be 1.7100 g/ml (16).

**DNA reannealing reactions.** The extent of DNA reassociation was determined spectrophotometrically, essentially by the method reported by Seidler and Mandel (18) and Seidler et al. (17), as described below. DNA was sheared by two passages through a French press at 10,000 lb/in² and was recovered as intact double-stranded fragments. To increase the reaction rate, the reassociations were done in 5X SSC (SSC = standard saline citrate, 150 mM NaCl-15 mM sodium citrate, pH 7.0), but with the addition of 20% dimethylsulfoxide to depress the melting temperature (ca. 12°C). Incubation times were approximately 2.5 to 3 h when the strains were highly related and 5 to 6 h when they were unrelated. The temperature for reas-
associations was $T_m - 25^\circ C$, and this was determined experimentally in 5x SSC-2% dimethylsulfoxide. Incubation temperatures ranged from 54 to 58$^\circ C$ depending upon the G+C content of the DNA. The concentration of DNA in the reaction mixture was adjusted to 75 $\mu$g/ml ($A_{260} = 1.5$); e.g., in a four-place cuvette holder, one cuvette contained only the incubation medium (blank), the second had 75 $\mu$g of DNA from strain 1 per ml, the third had 75 $\mu$g of DNA from strain 2 per ml, and the fourth had a mixture of 37.5 $\mu$g of DNA from each of the two strains per ml (total of 75 $\mu$g/ml).

Reassociation experiments were carried out with a model 250 recording spectrophotometer with a model 2527 thermoprogrammer (Gilford Instruments, Inc., Oberlin, Ohio). In the initial experiments, an early model thermoprogrammer cuvette block was used that consisted of four stainless steel cuvettes with removable quartz windows. To prevent the formation of large bubbles in the cuvettes during heating and cooling, the temperature was initially raised rapidly (ca. 12$^\circ C$/min) to 75$^\circ C$, and the rate was then decreased to 3$^\circ C$/min until the temperature reached 90$^\circ C$. This was at least 5$^\circ C$ above the conclusion of the hyperchromic shift for the strains studied. The temperature was held at 90$^\circ C$ for 10 min to ensure complete denaturation of the DNA. Cooling to $T_m - 25^\circ C$ was at 3$^\circ C$/min. Small bubbles, which usually formed at the periphery of the windows, were excluded from the light path by a sheet metal mask.

Later experiments used the newly designed thermoprogrammer block with four removable fused quartz cuvettes. Although bubbling seemed not to occur in these cuvettes, the same heating and cooling protocol was used as with the older cuvette block. Leakage around the Teflon stoppers of the quartz cuvettes was prevented by compressing a double layer of Teflon tape (0.09 mm thick) into the filling port as the stopper was inserted.

Values for DNA relatedness obtained spectrophotometrically by using the quartz cuvettes were comparable to those from radioisotope studies as reported under Results. Data from the two types of cuvette assembly were comparable in the mid to upper range of DNA relatedness, but in the lower range (ca. 0 to 30%) the early model cuvettes gave readings 10 to 20% higher. In addition, variation between replicates was much greater. Consequently, only data obtained with the fused quartz cuvettes are presented.

The extent of DNA reassociation was calculated as percent relatedness by the equation of Seidler and Mandel (18): $\{1 - [\text{obs. } C_{60.5} - \text{calc. } C_{60.5}] / \text{calc. } C_{60.5}] \times 100\}$ where obs. $C_{60.5}$ is the observed $C_{60.5}$ of a renatured mixture, calc. $C_{60.5}$ is the $C_{60.5}$ of the mixture expected if the two DNA molecules are identical in sequence, and calc. $C_{60.5}$ is the $C_{60.5}$ of the mixture expected for no sequence similarity (complete additivity of the independently measured $C_{60.5}$ values).

RESULTS

Spectrophotometric determination of DNA relatedness. Seidler and Mandel (18) reported that their method for spectrophotometric measurement of reassociation of bacterial DNA gave results comparable to those obtained with radioisotope techniques. The Seidler-Mandel method appears not to have been previously used to measure reassociation of yeast DNA. In view of this, we selected for comparison yeast strains that were studied earlier with radioisotope-labeled DNA. Price and collaborators (15) employed $^{32}$P-labeled DNA and hydroxylapatite fractionation in an extensive study of speciation in four genera. For comparison, we chose four pairs from that study and used the same strains. Despite procedural differences, our results with the spectrophotometric method were quite similar to those presented by Price et al. (Table 1).

The renaturation kinetics of DNA from Debaryomyces hansenii determined spectrophotometrically are depicted in Fig. 1A along with a theoretical second-order plot calculated from the equation of Britten and Kohne (3). It is apparent that reassociation departs from theoretical second-order kinetics, and this has been noted in other studies (15, 18). By using a Wetmur-Davidson (20) plot, we estimated the DNA preparation to be composed of 94% unique and 6% repetitive, rapidly renaturing sequences (Fig. 1B).

DNA from other strains used in this study contained no more than 5 to 8% repetitive sequences, and this was comparable to preparations obtained by Price et al. (15) with hydroxyapatite fractionation of DNA renatured to a $C_{60}$ value of 0.1 to 0.2. In our spectrophotometric system, approximately 12 min elapsed between the conclusion of the DNA melt at 90$^\circ C$ and the start of measurement of reassociation at $T_m - 25^\circ C$. It appears that a major portion of the repetitive sequences renature during the cool down and are excluded from the measurement of relatedness.

Description of new taxa. Latin diagnosis of Pichia amylophila sp. nov. Species heterothallica. Ascii dehiscentes, 2-4 ascosporos piliformes habentes (1.5-2.2 $\times$ 3.0-3.5 $\mu$m). Asci spheroidales-ellipsoidales (4.0-5.0 $\times$ 4.5-7.0 $\mu$m), alii liberi, alii adnexi, sessiles seu brevibus stigmatibus retenti.

In agar maltos-cellulae singulare, aut aliique, quando dispositae in racemis, plumque spheroidales-ellipsoidales seu elongatae (2.0-5.5 $\times$ 3-14 $\mu$m). Pseudohyphae copiosa; hyphae verae (1.8-2.5 $\mu$m diametro) rarae. In agar morphologico incrementum eburneum, nubilum, umidum; margo serrata, centrum paulum sublatum.

Glucosum acriter fermentatur; galactosum, maltosum, sucrosum, lactosum, raffinosum, et trehalosum non fermentantur. In agar assimilantur glucosum, sucrosum, maltosum, cellobiosum, trehalosum, melizitosum, amylum soluble, d-xylsosum, L-arabinosum, ethanolum,
TABLE 1. Comparison of the spectrophotometric method with a radioisotope technique for measuring DNA reassociation between species of Debaryomyces and Pichia

<table>
<thead>
<tr>
<th>Test pair and strain no.</th>
<th>% DNA relatedness assessed spectro-photometrically</th>
<th>% DNA relatedness assessed using °P-labeled DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>D. hansenii and D. tyrocola</strong></td>
<td>75 ± 3.0</td>
<td>77 ± 1.0</td>
</tr>
<tr>
<td>NRRL Y-7426, NRRL Y-1458 CBS 767, CBS 766 UCD 74-86, UCD 72-47</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>D. vanriji and D. yarrowii</strong></td>
<td>71 ± 2.3</td>
<td>68 ± 1.2</td>
</tr>
<tr>
<td>NRRL Y-7430, NRRL Y-7535 CBS 3024, CBS 6246 UCD 61-24, UCD 72-48</td>
<td></td>
<td></td>
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<tr>
<td><strong>D. hansenii and D. subglobosus</strong></td>
<td>44 ± 1.4</td>
<td>40 ± 2.6</td>
</tr>
<tr>
<td>NRRL Y-7426, NRRL Y-6666 CBS 767, CBS 792 UCD 74-86, UCD 75-17</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>D. hansenii and P. vini var. vini</strong></td>
<td>2 ± 1.9</td>
<td>2.7 ± 0.7</td>
</tr>
<tr>
<td>NRRL Y-7426, NRRL Y-1459 CBS 767, CBS 810 UCD 74-86, UCD 66-20</td>
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*Standard deviation calculated from three determinations.

_Data from Price et al. (15).

**Fig. 1.** Reassociation kinetics of DNA from Debaryomyces hansenii NRRL Y-7426 determined spectrophotometrically. (A) Comparison of experimental data (solid line) with theoretical second-order plot (dashed line) calculated by the method of Britten and Kohne (3); (B) early data points presented in second-order rate plot after Wetmur and Davidson (20).


_Description of P. amylophila._ Growth on malt extract agar: After 3 days at 25°C, the cells are single or rarely clustered, speroidal, ellipsoidal, or elongate (2.0 to 5.5 by 3 to 14 μm). Pseudohyphae are abundant and well branched. True hyphae (diameter: 1.8 to 2.5 μm) are rarely seen and show no evidence of a dolipore septum when viewed under the light microscope with an oil immersion objective (Fig. 2).

Growth on morphology agar: After 7 days at 25°C, growth is cream colored, dull, and moist

FIG. 3. *P. amylophila* NRRL YB-1287. Dehisced ascospores and ascus fragments. Diploid ascospores are noticeably larger than the haploid ascospores which are also present.

FIG. 4. *P. mississippiensis* NRRL YB-1294. True hypha with septum.


FIG. 6. *P. amylophila* NRRL YB-1287-82-3 × *P. mississippiensis* NRRL YB-1294-6. Conjugated cells (right, center) and asci with aborted spores (arrows).
with a serrate margin and a slightly raised center.

Sexual reproduction: Diploid cultures formed ascospores quite abundantly on 5% malt extract agar at 25°C after 7 to 10 days. Sporulation was nearly as good on YM agar. Two hat-shaped ascospores (1.5 to 2.2 by 3.0 to 3.5 μm) were formed in each dehiscent ascus (Fig. 3). Asci are spheroidal to ellipsoidal (4.0 to 5.0 by 4.5 to 7.0 μm) and generally are free, but they may be attached and sessile or attached by short sternig mata to pseudozystes.

Isolation of 83 single ascospores from NRRL YB-1287 (viability, 76/83) gave 70 that produced sporogenous colonies and 6 that formed asporogenous colonies. There was no sporulation in mixtures of asporogenous colonies. Two of the sporogenous colonies (strains 44 and 82) infrequently produced three- to four-spored asci. Single-spore isolates from strain 82 (viability 25/37, sporogenous 14, asporogenous 11) yielded asporogenous colonies that sporulated when mixed, thus showing P. amylophila to be heterothallic despite the tendency of the parent strain to produce predominantly diploid spores. NRRL YB-1287-82-2 (a) and NRRL YB-1287-82-3 (a) were selected as representative complementary mating types. Asci from this pair predominantly formed three to four spores.

Fermentation: Glucose was strongly fermented. A bubble formed in maltose and sucrose tubes after 24 days. There was no fermentation of galactose, lactose, raffinose, or trehalose.

Assimilation of carbon compounds: Glucose, sucrose, maltose, cellobiose, trehalose, melezitose, soluble starch, d-xyllose, l-arabinose, ethanol, glycerol, d-mannitol, d-glucitol, α-methyl-d-glucoside, potassium-d-gluconate, DL-lactic acid, succinic acid, and citric acid were utilized as sole carbon sources.

Carbon compounds not assimilated: Galactose, l-sorbose, lactose, melibiose, raffinose, inulin, d-arabinose, d-ribose, l-rhamnose, d-glucoamine-hydrochloride, i-erythritol, ribitol, galactitol, salicin, calcium-2-ketogluconate, potassium-5-ketogluconate, potassium sodium saccharate, ethylacetocetate, and i-inositol were not utilized as sole carbon sources.

Assimilation of potassium nitrate: negative.

Growth in vitamin-free medium: negative.

Growth in 10% sodium chloride plus 5% glucose in yeast nitrogen base: negative.

Growth at 37°C: positive.

Starch formation: negative.

Liquefaction of gelatin: negative.

G+C content of the nuclear DNA: 45.4 mol%.

Type: The type strain, NRRL YB-1287, the only known strain of this species, was isolated from frass collected from loblolly pine, Pinus taeda L., in late October 1949, from Piney Woods, Miss. (Table 2).

Latin diagnosis of Pichia mississippiensis sp. nov. Species heterothallica. Asci dehiscentes, granulosae, elliptico-spheroidales, multo discrepantes magnitudine (1.5-2.0 × 2.5 - 3.5 μm) habent. Asci spheroidales-ellipsoidales (3.0-5.0 × 4.5-6.5 μm), liberi seu adnexit ad pseudohyphales; aliquando a cellulae pseudohyphales seu hyphales fit ascus.

In agare maltoso cellulae singulae, aliquando binae aut in racemis, plerumque spheroidales-elongatae, rara spheroidales, multo discrepantes diametro (1.1-5.0 × 2.0-12.0 μm). Pseudo hyphales copiosae, hyphae verae (1.7-3.0 μm diam) rarae. In agare morphiologico incrementum fuscon palmillum, nitens, butyrosum; cen-

<table>
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<tr>
<th>Table 2. Sources and ploidy of strains of P. amylophila and P. mississippiensis</th>
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<tbody>
<tr>
<td>Species and NRRL no.</td>
</tr>
<tr>
<td>--------------------</td>
</tr>
<tr>
<td>P. amylophila</td>
</tr>
<tr>
<td>YB-1287</td>
</tr>
<tr>
<td>P. mississippiensis</td>
</tr>
<tr>
<td>YB-1269</td>
</tr>
<tr>
<td>YB-1270</td>
</tr>
<tr>
<td>YB-1276</td>
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<tr>
<td>YB-1285</td>
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<tr>
<td>YB-1286</td>
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<tr>
<td>YB-1290</td>
</tr>
<tr>
<td>YB-1291</td>
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<tr>
<td>YB-1292</td>
</tr>
<tr>
<td>YB-1294</td>
</tr>
<tr>
<td>YB-1317</td>
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</tbody>
</table>

* The a and α designations are used for convenience and have not been correlated with the a and α mating types of Saccharomyces cerevisiae.
trum coloniae sublatum, margo non fractum.
Cultura olet subacidum sed suave.

Glucosum vehementer fermentatur. Galactosum,
maltosum, sucrosum, lactosum, raffinosum,
trehalosum non fermentantur.

Assimilantur glucosum, sucrosum, maltosum,
cellobiosum, trehalosum, melezitosum, D-xylosum,
L-arabinosum, D-arabinosum (imbecillum, negativum),
L-rhamnosum (positivum, negativum), ethanolum,
glycerolum, D-mannitolum, D-glucitolum,
α-methyl-D-glucosidum, salicinum, (positivum, imbecillum, negativum),
potassii-D-glucosan, DL-acidum lacticum, acidum succinicum,
acidum citricum (positivum, negativum). Non assimilantur galactosum,
L-sorbosum, lactosum, melibiosum, raffinosum, inulinum,
amylum solubile, D-ribosum, D-glucosaminum-HCl,
1-erythritolium, ribitolum, galactitolum, calci2-2-ketogluconas,
potassii 5-ketogluconas, potassii-sodii saccharas, ethylactoacetates,
i-inositolum, potassii nitras. Nullum augmentum
si medium caret vitaminis. Augmentum co-
oposum in temperatura 37°C. Amylum non fit;
esteres non fiant; gelatum non liqueatur.

Typus NRRL YB-1294 (CBS 7023) designat
stirpem typicam. Isolatur a fimo scarabaeorum
a Pinus taeda L. apud Piney Woods, Miss.,
United States. Typi conjungentes complementarii:
NRRL YB-1294-6 (CBS 7024) et NRRL
1294-7 (CBS 7025). Culturae servatur in Collec-
tione Culturarum, Officina Investigationum
Tractus Borealis, Peoria, Ill., United States.

**Description of P. mississippiensis.** Growth
on malt extract agar: After 3 days at 25°C,
the cells are single or infrequently in pairs or
clusters. Cells are ellipsoidal to elongate, or occa-
sionally spheroidal, and quite variable in size
(1.1 to 5.0 by 2.0 to 12.0 μm). Pseudohyphae are
abundant and well developed. True hyphae (di-
meter: 1.7 to 3.0 μm) are uncommon and show
no evidence of a dolipore septum when viewed
under the light microscope with an oil immersion
objective (Fig. 4).

Growth on morphology agar: After 7 days at
25°C, growth is light tan, glistening, and butyr-
aceous; the colony center is raised; and the margin
entire. The culture has a faint, pleasant acidic
odor.

Sexual reproduction: Cultures formed abun-
dant ascospores on YM agar and on 5% malt
extract agar at 25°C after 5 to 8 days. Two to
four hat-shaped ascospores (1.5 to 2.0 by 2.5 to
3.5 μm) were formed in each dehiscent ascus
(Fig. 5). Asci are spheroidal to ellipsoidal (3.0 to
5.0 by 4.5 to 6.5 μm) and are free or attached to
pseudohyphae. Infrequently, a pseudohyphal or
hyphal cell became an ascus.

Single ascospore isolates (viability, 26/31) pro-
duced asporogenous colonies. The species was
demonstrated to be heterothallic through pair-
ing of appropriate single-spore isolates. NRRL
YB-1294-6 (a) and NRRL YB-1294-7 (a) were
selected as representative complementary mat-
ting types.

Fermentation: Glucose was strongly fer-
mented, and occasionally a bubble formed in
maltose and sucrose tubes. There was no fer-
mentation of galactose, lactose, raffinose, or tre-
halose.

Assimilation of carbon compounds: Glucose,
sucrose, maltose, cellobiose, trehalose, melezit-
ose, D-xylose, L-arabinose, D-arabinose (weak or
negative), L-rhamnose (positive or negative),
ethanol, glycerol, D-mannitol, D-glucitol, α-
methyl-D-glucoside, salicin (positive, weak, or
negative), potassium-D-glucuronate, D-lactic acid,
succinic acid, and citric acid (positive or nega-
tive) were utilized as sole carbon sources.

Carbon compounds not assimilated: Galacto-
lose, L-sorbosum, lactose, melibiosum, raffinosum,
inulin, soluble starch, D-ribosum, D-glucosamine-
hydrochloride, 1-erythritol, ribitol, galactitol,
calcium-2-ketogluconate, potassium-5-ketoglu-
conate, potassium sodium saccharate, ethylacet-
oacetate, and i-inositol were not utilized as sole
carbon sources.

Assimilation of potassium nitrate: negative.
Growth in vitamin-free medium: negative.
Growth in 10% sodium chloride plus 5% glu-
cose in yeast nitrogen base: negative.
Growth at 37°C: positive.
Starch formation: negative.
Production of esters: negative.
Liquefaction of gelatin: negative.
G+C content of the nuclear DNA: 47.2 to 48.0
mol% (range of three strains, Table 3).

Type: NRRL YB-1294, the type strain, was
isolated from frass collected from a loblolly pine,
*Pinus taeda L.*, in late October 1949, from Piney
Woods, Miss. Other strains of this species are
listed in Table 2.

*Candida obtusa var. arabinosa* Montrocher
(14) was isolated from *Clitopilus prunulus*
(Scop. ex Fr.) Kummer in France and appeared
identical to haploid forms of *P. mississippiensis*.
Abundant conjugation and sporulation resulted
when the type strain (NRRL Y-11,748 = CBS
5837) of this asexual yeast was mixed with YB-
1294-7. Single-spore isolates from this pairing
exhibited good viability (30/34 viable), and ap-
propriate sibling pairs formed mature ascos-
spores.

By means of standard fermentation and assim-
ilation tests, *P. amylophila* is separated from *P.
mississippiensis* only by its ability to assimilate
soluble starch. Nine single-spore isolates from
NRRL YB-1287 or from crosses of its progeny were tested for starch utilization and all were found positive, so the character appeared reliable for separation of these two species.

**Interspecific mating tests.** Initially, the two new species were considered to represent the same taxon, but their difference in starch assimilation prompted us to undertake genetic crosses between them. Data in Table 4 show that the frequency of interspecific conjugation is comparable to that for intraspecific pairings; however, most of the ascospores resulting from interspecific crosses were poorly formed (Fig. 6). The results were similar when the other haploid strains of *P. mississippiensis*, including *C. obtusa* var. *arabinosa*, were paired with *P. amylophila*. Because of the rarity of apparently mature spores from interspecific crosses, single-spore isolations proved difficult, but the few spores isolated were not viable (Table 4). With the exception of NRRL YB-1291, which failed to sporulate or to show mating competence, all haploid strains of *P. mississippiensis* gave at least 50% ascospore viability when paired with the appropriate mating type from NRRL YB-1294.

In further comparisons, mating types of *P. amylophila* and *P. mississippiensis* were paired with mating types from the phenotypically similar heterothallic species *P. wickerhamii* (NRRL Y-2435-9, NRRL Y-2435-10), *P. rhodanensis* (NRRL YB-651 sm-14, NRRL Y-7854 Re-1), and *P. veronneae* Kodama (NRRL Y-7818-1, NRRL Y-7818-10). Mating responses were not detected in these mixtures or in interspecific mixtures of mating types from the latter three species.

**DNA reassociation studies.** Consistent with results from genetic crosses, DNA reassociation experiments showed high (96 to 99%) base sequence complementarity between strains of *P. mississippiensis* but low (20 to 27%) complementarity between *P. mississippiensis* and *P. amylophila* (Table 5). Nonetheless, the limited extent of reassociation shown between these two species was still noticeably higher than that of the ascospores resulting from interspecific crosses were poorly formed (Fig. 6). The results were similar when the other haploid strains of *P. mississippiensis*, including *C. obtusa* var. *arabinosa*, were paired with *P. amylophila*. Because of the rarity of apparently mature spores from interspecific crosses, single-spore isolations proved difficult, but the few spores isolated were not viable (Table 4). With the exception of NRRL YB-1291, which failed to sporulate or to show mating competence, all haploid strains of *P. mississippiensis* gave at least 50% ascospore viability when paired with the appropriate mating type from NRRL YB-1294.

Table 3. Nuclear DNA base composition of *P. amylophila*, *P. mississippiensis*, and similar species

<table>
<thead>
<tr>
<th>Species and NRRL no.</th>
<th>G+C ± SD* (mol%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. amylophila</em> YB-1287</td>
<td>45.4 ± 0.00</td>
</tr>
<tr>
<td><em>P. mississippiensis</em> YB-1269</td>
<td>47.7 ± 0.08</td>
</tr>
<tr>
<td>YB-1294</td>
<td>47.2 ± 0.10</td>
</tr>
<tr>
<td>YB-1317</td>
<td>48.0 ± 0.15</td>
</tr>
<tr>
<td><em>P. rhodanensis</em> Y-7854</td>
<td>51.9 ± 0.12</td>
</tr>
<tr>
<td>YB-651 sm-14</td>
<td>51.8 ± 0.09</td>
</tr>
<tr>
<td><em>P. wickerhamii</em> Y-2435</td>
<td>46.5 ± 0.14</td>
</tr>
<tr>
<td>C. obtusa var. <em>arabinosa</em> Y-11,748</td>
<td>47.4 ± 0.11</td>
</tr>
</tbody>
</table>

* Standard deviation (SD) calculated from three to four determinations.
* Type strain.

Table 4. Reaction between mating types from *P. amylophila* NRRL YB-1287 and *P. mississippiensis* NRRL YB-1294

<table>
<thead>
<tr>
<th>Mating types*</th>
<th>Reaction</th>
<th>Ascospore viability* (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>YB-1287-82-2 + YB-1287-82-3</td>
<td>35% conjugation; 40% of conjugants formed ascospores</td>
<td>100 (46/46)</td>
</tr>
<tr>
<td>YB-1294-6 + YB-1294-7</td>
<td>15% conjugation; 65% of conjugants formed ascospores</td>
<td>98 (44/45)</td>
</tr>
<tr>
<td>YB-1287-82-2 + YB-1294-6</td>
<td>No conjugation; no ascospores</td>
<td>Not determined</td>
</tr>
<tr>
<td>YB-1287-82-2 + YB-1294-7</td>
<td>10% conjugation; 5% of conjugants formed ascospores. Dehisced, apparently mature ascospores were no more frequent than 1 per 2 x 10^4 cells</td>
<td>0 (0/4)</td>
</tr>
<tr>
<td>YB-1287-82-3 + YB-1294-6</td>
<td>20% conjugation; 5% of conjugants formed ascospores. Dehisced, apparently mature ascospores were no more frequent than 1 per 2 x 10^4 cells</td>
<td></td>
</tr>
<tr>
<td>YB-1287-82-3 + YB-1294-7</td>
<td>No conjugation; no ascospores</td>
<td></td>
</tr>
</tbody>
</table>

* When grown alone, mating types showed neither conjugations nor ascospores.
* Determined by single ascospore isolation of dehisced ascospores.
* Dehisced, apparently mature ascospores from interspecific crosses were approximately 1.5 times larger than ascospores from intraspecific crosses, and only two were formed per ascus.
obtained when they were compared with such unrelated species as \textit{P. rhodanensis}, \textit{P. wickerhamii}, or \textit{Hansenula petersonii}.

Results from the DNA studies parallel those from genetic crosses and confirm \textit{C. obtusa} var. \textit{arabinosa} to be the imperfect form of \textit{P. mississippiensis}. It is also evident from these data that \textit{P. wickerhamii} and \textit{P. rhodanensis} represent distinct species.

**DISCUSSION**

Determination of relatedness between microorganisms by comparison of their nuclear DNA can be accomplished by several methods. One of the commonest techniques involves incubating unsheared single-stranded DNA immobilized on nitrocellulose filters in the presence of sheared single-stranded DNA labeled with radioisotopes. At the conclusion of the reaction, the amount of reassociation is measured by the amount of radioactivity retained by the filter. Price and co-workers (15) used radioactive DNA in their study of relatedness among yeasts, but the protocol differed in that the double-stranded DNA was bound to hydroxylapatite after completion of the renaturation reaction. Measurement of radioactivity in bound and unbound fractions allowed assessment of relatedness.

The spectrophotometric method for measuring reassociation is particularly attractive when difficulties are encountered with in vivo radioisotope labeling of DNA or where numerous pairwise comparisons are to be made and the time required for labeling becomes excessive. The spectrophotometric method has been used in studies of bacteria (e.g., 4, 17, 18), and results were similar to those obtained with radioisotopes. Martini and Phaff (10) used the spectrophotometric method of De Ley et al. (4) with some success to compare relatedness among yeasts, but the procedure of Seidler and Mandel (18), which monitors the reaction over a much longer period, has not been applied to yeast systematics. Our results showed good correlation with data obtained by radioisotope studies (15) and suggest that the spectrophotometric technique of Seidler and Mandel may be reliably applied to yeasts and probably other eucaryotes.

In recent years, yeast taxonomists have turned increasingly to the use of DNA base sequence comparisons for clarification of species relationships that defy solution by traditional means (1, 11-13, 15). Relatedness values of 20 to 25% or less were taken to mean that the strains in question represented different species, whereas values of 80 to 100%, as suggested by Price et al. (15), indicated that strains belong to the same species. Values in the 20 to 80% range seldom have been found, and too little data are presently available for an understanding of their significance.

Results from mating tests between \textit{P. amylolpha} and \textit{P. mississippiensis} suggested considerable genetic homology between these two species despite the lack of viable ascospores. Consequently, it was quite surprising to find that the strains in question represented different species, whereas values of 80 to 100%, as suggested by Price et al. (15), indicated that strains belong to the same species. Values in the 20 to 80% range seldom have been found, and too little data are presently available for an understanding of their significance.

Winge and Laustsen (23) and Wickerman and Burton (22) have proposed that interfertility among yeasts be considered indicative of con-specificity. However, failure to mate cannot exclude con-specificity because progeny derived from the same diploid parent have been noted to lack mating competence (8). Consequently, the decision to consider \textit{P. amylolpha} and \textit{P. mississippiensis} as separate species was made on the basis of both their limited DNA related-
ness and their inability to form viable ascospores.

The use of fertility as the sole criterion for speciation must be carefully considered, for this would require that the varieties of P. scutulata be assigned to the same species even though the extent of their DNA relatedness is the same as that of the new Pichia species described here. Conversely, defining species strictly on the basis of a certain percentage of DNA relatedness may conflict with the biological reality of genetic exchange. Almost certainly, considerable latitude will be found in the definition of species even at the genetic and molecular levels. For the present, our data suggest that DNA base sequence complementarity as low as 25% may not preclude conspecificity since sufficient genetic compatibility can exist between some strains to allow sexual reproduction.

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We are greatly indebted to H. J. Phaff, Department of Food Science and Technology, University of California, Davis, for allowing one of us (C.P.K.) use of laboratory facilities where the preliminary isolation and characterization of DNA used in this study was accomplished. We thank C. W. Price for helpful suggestions on experimental protocol and for critical review of the manuscript. H. C. Nielsen and K. R. Sexson generously gave suggestions on the operation of the Spinco analytical ultracentrifuge used for determination of G+C values. H. M. Howe kindly provided the Latin diagnoses of the new species.

REPRINT REQUESTS

Address reprint requests to: Dr. C. P. Kurtzman, Northern Regional Research Center, 1816 N. University St., Peoria, IL 61604.

LITERATURE CITED