Establishment of uracil auxotrophic dikaryotic strains of *Lentinula edodes* by crossbreeding

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The uracil auxotrophic monokaryotic strain 423-9 of *Lentinula edodes* was crossed with nine monokaryons (cro2-2-9, W66-1, xd2-3-2, QingKe 20A, 241-1-1, 9015-1, L66-2, 241-1-2, and Qing 23A) derived from wild type strains of *L. edodes*. Nine dikaryotic hybrids were established from these crosses. These hybrids were fruited and 496 single spore isolates were obtained. Among these single spore isolates, 166 were identified as monokaryons under a microscope. We screened these monokaryons on selective medium and obtained 19 uracil auxotrophic monokaryons. By using the Monokaryon-monokaryon crossing method among the uracil auxotrophic monokaryons, 56 uracil auxotrophic dikaryotic strains were established on selective medium. These dikaryotic strains were unable to grow on minimal medium without uracil and exhibited slow growth rates on PDA plates compared to the wild type strain. The uracil auxotrophic dikaryotic strains also showed more vigorous growth on sawdust cultivation medium containing uracil than that without uracil. The fruiting tests showed that they formed normal fruiting bodies on the sawdust medium containing uracil. The results show that the uracil auxotrophic dikaryotic strain of *L. edodes* could be produced by mating, and will provide a valuable resource for future genetic studies and for spawn protection and identification.

**Key Words:** *Lentinula edodes*, uracil auxotroph, dikaryon.
breeding involving isolation by a routine separation method to develop an auxotrophic strain that cannot grow well without nutrient supplementation.

Materials and Methods

Strains, media, and growth conditions

The uracil auxotrophic monokaryotic strain of *L. edodes* 423-9 was obtained by UV mutagenesis of protoplasts.

The following media were employed for mycelia growth: Potato Dextrose Agar (PDA): 200 g/L potatoes, 20 g/L glucose, 20 g/L agar; Potato Dextrose Agar containing uracil (PDAU): PDA containing 0.05 mmol/L uracil; Potato Dextrose (PD): 200 g/L potatoes, 20 g/L glucose; Minimal medium (MM): KH₂PO₄ 1.0 g/L, (NH₄)₂HPO₃ 1.5 g/L, MgSO₄·7H₂O 0.3 g/L, Thiamine HCL 500 μg/L, agar 15 g/L; Minimal medium containing uracil (MMU): MM containing 0.05 mmol/L uracil; Minimal medium containing uracil and 5′-FOA (MMUF): MM containing 0.05 mmol/L uracil, 0.5 g/L 5′-FOA. All the media were sterilized for 15 mins at 121°C. Fifteen milliliters of each media was poured into individual plastic Petri dishes with a 9 cm diameter. The mycelia were kept in a dark room at 25°C for 1–2 w.

The spores were germinated on PDA medium and kept in a dark room at 25°C for 2 w.

The dikaryotic hybrids were inoculated onto sterile sawdust cultivation bags for producing fruiting bodies. The sawdust substrate was composed of 8:2 mixtures of oak sawdust and rice bran, including 63% water content (Lee et al. 2012). The sawdust cultivation bags were incubated at 25°C to 30°C for 3 months.

Selection of parental strains for crossing experiments

The uracil auxotrophic monokaryotic strain of *L. edodes* 423-9 was crossed with nine monokaryons (cro2-2-9, W66-1, xd2-3-2, QingKe 20A, 241-1-1, 9015-1, L66-2, 241-1-2, and Qing 23A) derived from wild type strains of *L. edodes*. Nine dikaryotic hybrids (Z1, Z2, Z3, Z4, Z5, Z6, Z7, Z8, and Z9) were obtained from crossing two uracil auxotrophic monokaryons.

Basidiospore production and germination

Mature, large, thick, disease- and insect-pest-free fruiting bodies of Z1-Z9 were collected to prepare spore prints. The spores were collected in Eppendorf tubes filled with sterile water and serially diluted with sterile water until 2–3 spores were visible in a single field of view under a microscope. Spore dilution samples were spread on the PDA plate at 25°C for incubation (Zhang et al. 2002). The spores were allowed to germinate for 2 w.

Microscopic observation

Selected single colonies were transferred to PDA medium and incubated for 7 d. For each selected colony, it was further purified by transferring a small mycelia block from the edge of the colony and placing it in the center of a new PDA plate. After 10 d of incubation, mycelia were observed under a microscope, and monokaryons and dikaryons were distinguished by clamp connection.

Uracil auxotrophic mutant selection

Fresh inocula from the margins of monokaryotic colonies were transferred to MM, MMU, and MMUF plates, respectively, for selecting the uracil auxotrophic mutant. The uracil auxotrophic mutants were identified based on whether they could grow well on MMU and MMUF but could not grow on MM.

Monkaryon-monkaryon (Mon-Mon) crossing method

Small pieces of mycelia from two uracil auxotrophic monokaryons were placed side by side on MMU plates for mon-mon mating. The uracil auxotrophic dikaryotic strains were identified based on the detection of clamp connections after fusion between hyphae of different monokaryons.

Genomic DNA preparation

All strains were cultivated in 100 ml of PD medium at 25°C for 10 d with constant shaking (150 rpm). Fungal mycelia were harvested using sterile cloth and washed twice with sterile water. Preparation of genomic DNA was performed using a method described previously (Zhang et al. 2006).

PCR testing

Detection of the *pyrF* gene (GenBank: KJ934231.1) was conducted using primers *LEpyrF-F*1150 and *LEpyrF-R*1150 (*Table 2*). Amplification included an initial denaturation at 94°C for 5 min, followed by 35 cycles of denaturation at 94°C for 30 s, annealing at 55°C for 30 s, and elongation at 72°C for 75 s, and a final elongation step at 72°C for 10 min.

Mycelial growth rate measurement

Tested strains were incubated on PDA plates and PDAU plates, and the mycelial growth rate was measured 7 d later. The dikaryotic strains were also incubated in test tubes

| Table 1. | The uracil auxotrophic monokaryotic strain of *L. edodes* 423-9 crossed with nine monokaryons derived from wild type *L. edodes* |
| --- | --- |
| **F1** | **P** |
| Z1 | 423-9 × cro2-2-9 |
| Z2 | 423-9 × W66-1 |
| Z3 | 423-9 × xd2-3-2 |
| Z4 | 423-9 × QingKe 20A |
| Z5 | 423-9 × 241-1-1 |
| Z6 | 423-9 × 9015-1 |
| Z7 | 423-9 × L66-2 |
| Z8 | 423-9 × 241-1-2 |
| Z9 | 423-9 × Qing 23A |

| Table 2. | The primers to amplify the *pyrF* gene |
| --- | --- |
| **Gene** | **Primer** | **Sequence (5′-3′)** |
| *pyrF* | *LEpyrF-F*1150 | CGCTTTATCTTTTATCTCGTGCTCAT |
| | *LEpyrF-R*1150 | GAACGGTGTAGTAAGACAGGTAGTCA |
filled with sawdust cultivation medium which contained various concentrations of Uracil (0, 0.025, 0.05, 0.075 mmol/L), and the mycelial growth rate was measured 15 d later.

**Fruiting test**

Five uracil auxotrophic dikaryotic strains were selected at random for fruiting tests. The dikaryotic stains were inoculated onto sterile sawdust cultivation bags, and the media were supplemented with uracil (0.075 mmol/L).

**Results**

The collected spores from newly generated dikaryons Z2, Z3, Z5, Z6, Z7, and Z8 germinated while those of Z1, Z4, and Z9 did not. Four hundred and ninety-six single colonies were picked after the spores were germinated on PDA medium. After 10 d of incubation, mycelia from these single spores were observed under a microscope, and monokaryons and dikaryons were distinguished by the existence of clamp connections. From the 496 colonies, 166 monokaryons and 330 dikaryons were identified. These monokaryons were screened on selective medium, and 19 uracil auxotrophic monokaryotic strains (Z2-13, Z2-14, Z2-15, Z2-18, Z2-77, Z2-97, Z3-1, Z3-2, Z3-3, Z3-4, Z3-6, Z3-7, Z3-8, Z3-27, Z3-41, Z3-42, Z5-25, Z6-9, and Z7-11) were obtained (Fig. 1). The proportion of uracil auxotroph monokaryotic strains was therefore 11.4%.

Mon-Mon mating among the above-mentioned 19 uracil auxotrophic monokaryotic strains was performed. Finally, 56 new dikaryons (UH1-UH56) were established and identified based on the detection of clamp connections under a microscope after the fusion between hyphae of different monokaryons (Fig. 2). By checking the selection media (MM, MMU, and MMUF), all 56 crossing dikaryons were uracil auxotrophy (Fig. 3). As shown in Fig. 4, the *pyrF* gene in the uracil auxotrophic dikaryotic strains completely corresponded with that of the parent strain 423-9, which has an 83 bp deletion in the *pyrF* gene.

The uracil auxotrophic dikaryotic strains exhibited slower growth rates on PDA plates compared to wild type strains (Fig. 5), but exhibited vigorous growth on PDAU plates. The growth rate of wild type strains was similar on PDAU plates compared to PDA plates (Fig. 6). The uracil auxotrophic dikaryotic strains also showed more vigorous growth on sawdust cultivation medium containing uracil than that

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**Fig. 1.** The growth of the uracil auxotrophic monokaryon Z2-18 of *L. edodes* and *L. edodes* strain 135 on selective media. Contrary to the wild type *L. edodes* strain 135, the uracil auxotrophic monokaryon Z2-18 of *L. edodes* could grow well on MMU and MMUF but could not grow on MM.

**Fig. 2.** Microscopic examination of uracil auxotrophic dikaryons of *L. edodes*. The configuration of clamp connections is indicated by the arrows (400×). Bar, 10 μm.

**Fig. 3.** The growth of the uracil auxotrophic dikaryons UH25 of *L. edodes* and *L. edodes* strain SUXIANG on selective media.

**Fig. 4.** PCR amplification of *pyrF* gene. Lanes 1 and 2, DNA isolated from wide type *L. edodes* strain 135 and *L. edodes* strain SUXIANG; lane 3, DNA isolated from the uracil auxotrophic monokaryotic strain 423-9 of *L. edodes*; lanes 4 to 9, DNA isolated from the uracil auxotrophic dikaryotic strains UH1, UH9, UH18, UH20, UH26, and UH48 of *L. edodes*, respectively; lane 10, negative control with water; M, DNA molecular size markers (DL2000 marker).
without uracil. The growth rate of wild type strains was similar on cultivation medium containing various concentrations of uracil (Fig. 7).

The fruiting tests of the 5 uracil auxotrophic dikaryotic strains (UH1, UH18, UH20, UH25, UH26) showed that they all formed normal fruiting bodies (Fig. 8) on the sawdust medium containing uracil.

**Discussion**

In this study, we obtained 56 uracil auxotrophic dikaryons by using the Mon-Mon crossing method. All five strains produced fruiting bodies. The results suggested that uracil auxotrophy was stable during sexual development and could be transferred to target strains with advantageous traits by mating. On the other hand, the testing results showed these auxotrophic strains exhibited retarded growth rates on PDA medium compared to the wild type strains but exhibited a range of higher growth rates on PDAU medium. These characteristics of the auxotrophic strains could provide three potential values for spawn protection and identification. First, the uracil auxotrophy represents a new type of molecular marker to track strains and spawns, which may give breeders a powerful piece of evidence to protect their cultivars. Second, the auxotrophy phenotype provides a simply way to help growers determine whether the strain was contaminated based on its growth rates on the PDA and MM media. The contaminations by used spawns of *L. edodes* in commercial production have brought great losses to the farmers in China every year. Third, the auxotrophy will only exhibit normal growth on the sawdust cultivation medium containing the corresponding nutrition, such as uracil in this study, provided by the breeders. The potential advantages this auxotrophic marker may provide may enhance breeders’ enthusiasm to breed more new varieties of mushrooms. In addition, the establishment of this and
other auxotrophic markers should enhance genetic research on this important edible fungus.

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