Mushrooms are filamentous macrofungi which have huge edible, medicinal, and economical value [1]. Nowadays, more and more genome sequences of mushrooms have been completed [2]. Gene functional analysis to facilitate mushroom biotechnology is an important task in the post-genomic era. Various reverse genetics approaches including gene silencing, gene targeting, and T-DNA insertional mutagenesis are available to investigate the gene function [3–6]. Transformation is an important tool of reverse genetics research. However, screening many transformants by conventional PCR analysis is laborious and time-consuming because of the labor intensive process of DNA isolation [7]. Because of this limitation, it is necessary to develop a rapid and effective colony PCR for screening transformants in mushroom transformation experiments.

Although direct colony PCR method was already established for fungi, several fungal species could not be successfully amplified [8], pre-treatment of fungal colonies is still the most promising strategy [9]. Novozyme 234 was used for pre-treatment in lower fungi for colony PCR, but it needed additional incubation step for inactivation of enzyme [10]. Commercial kits, like Ampdirect plus (Shimadzu, Kyoto, Japan) and Y-PER (Cat# 78990; Thermo Scientific, Waltham, MA) have been applied to colonies, but mushrooms were not tested extensively for successful colony PCR. With the assistance of microwave treatment, rapid and simple template preparation from colonies and fruit bodies in several mushrooms was successful for PCR amplification [13]. However, thermal effects of microwave reduced the ability for amplification of large PCR products. Combining several kinds of pre-treatment together, a microwave, plMAN5C, proteinase K, and boiling (MMPB) method for colony PCR in red yeasts was developed [14]. This method has several drawbacks including the time-consuming manipulation and the use of expensive reagents such as proteinase K.

In this study, a rapid and simple colony PCR was developed to screen transformants in several common mushrooms including *Tremella fuciformis*, *Pleurotus ostreatus*, and *Pleurotus tuber-regium*. These strains were preserved in the Laboratory of Food Microbiology, Huazhong Agricultural University. *T. fuciformis* and *P. ostreatus* were sub-cultured on potato dextrose agar (PDA) at 25 °C, *P. tuber-regium* at 32 °C. The plasmids carrying *lac*, *hph*, and *egfp* had been used and *Agrobacterium tumefaciens* mediated transformation of these mushrooms was carried out as described before [15–17].

For screening the transformants, a small amount of the cells (approximately, 1–10 µg, 0.5–1 mm³) or...
the actively growing edge of mycelia (1–10 μg and 1–2 mm in diameter) were picked up from the selective plates PDSA (50 μg/ml hygromycin B and 200 μg/ml cefotaxime) by using a sterilized toothpick and resuspended in 20 μl Lywallzyme solution (an effective cell wall lyticase, Guangdong Institute of Microbiology, Guangdong, China) in a 200 μl PCR tube. To facilitate the disruption of the tissue, the mixture was vortexed for 10 s and an additional incubation under 34 °C for 15 min. Vortexing can be omitted in some cases, especially for mycelia because of their characteristic of dispersion. A 2 μl suspension was used as template in PCR reactions. PCR reactions were performed in a 20 μl volume containing 2 μl 10 × EasyTaq buffer, 200 μM dNTPs, 0.2 μM of each primer, and 1 U EasyTaq DNA polymerase (TransGen Biotech, Beijing, China). The primers used in this study were listed in Table 1. The PCR procedures were: 95 °C for 3 min, followed by 34 cycles of denaturation at 95 °C for 30 s, annealing at 55–58 °C (according to different primers) for 30 s, extension at 72 °C and a final elongation step at 72 °C for 5 min. Ultrapure water was used instead of DNA templates in negative controls, while vectors were used as positive controls. The PCR products were examined by 1.0% (w/v) agarose gel electrophoresis.

To make the procedure more cost-effective and convenient, we optimized it using transformants of T. fuciformis yeast-like cells (YLCs). Several important parameters such as concentration of enzyme, incubation time, quantity of template, and inactivation of enzyme were tested. The results showed lower concentration Lywallzyme (≤0.3125%) were capable of amplifying the desired fragment, higher concentration Lywallzyme (>0.3125%) failed to obtain the expected products (Figure 1(A)). A reasonable explanation is that PCR amplification was inhibited by higher concentration of Lywallzyme, while lower concentration of Lywallzyme has less inhibitory effect to PCR amplification and was enough to wound the cell wall of YLCs. We also explored whether the incubation time had influence on colony PCR in YLCs of T. fuciformis. The incubation time was varied from 15 to 120 min under the optimized enzyme concentration. The results revealed that it was not the most critical factor for PCR reactions, although longer time was beneficial for releasing DNA from cells (Figure 1(B)). In addition, the quantities of templates in supernatant (after centrifuging) was less than those in suspension, it had little influence in our protocols (Figure 1(C)). But inactivation of enzyme by heating treatment after incubation led to difficult PCR amplification. Distinctly weaker bands were obtained after 100 °C heating treatment compared to the untreated samples (Figure 1(C)).

The optimal procedure was established: picking up a suitable amount of fungal tissue (approximately 1–10 μg) to 20 μl 0.25% Lywallzyme solution, vortexing for 10 s, and then incubation at 34 °C for 15 min, and 2 μl suspension was used as templates to perform PCR. It was used to screen transformants of T. fuciformis, P. ostreatus, and P. tuber-regium. Amplification of the lac3, hph, egfp, or egfp-hph was achieved successfully in transformants of these mushrooms after enzymolysis treatment (Figure 2).

Compared with the conventional PCR analysis, the presented method requires neither the use of expensive and specialized equipment or toxic reagent like phenol used for DNA extraction. It significantly reduced the time and costs involved. In addition, in contrast to other published colony PCR protocols [10,11,14], no additional boiling step was in general required for lysing the microorganisms or inactivating the enzyme. Its simplicity makes it convenient to prepare a large amount of samples at the same time. Compared to the current protocol [13], our new method requires less harsh treatment of the sample and takes less time. A 5-minute incubation of cells and mycelia is sufficient to produce successful amplification using our colony PCR approach (data not shown). The operation steps of this method are simple, without the need for a cooling step after heat treatment. Using this procedure, more than 100 samples of mushroom DNA can be prepared in 30 min, whereas the microwave treatment method needs approximately 1 h. Noteworthy is that PCR products as large as ~3 kb were obtained using this method (data not shown). Moreover, compared to a commercially available kit (2 × T5 Direct PCR kit) (Tsingke, Beijing, China), the electrophoresis results of this colony PCR was also satisfactory with specific amplification (Figure 3). The addition of T5 DNA polymerase to the PCR system increased the rate of amplification compared to the EasyTaq DNA polymerase. However, potential primer bias existed in fungi and primer choice was critical for PCR success [18,19]. Multiple direct colony PCRs with different primer combinations or

### Table 1. Primers used in this study.

| Primers | Sequences (5’ to 3’) | Description |
|---------|----------------------|-------------|
| lac3-1-F | CGCTACGCGTCGTTGGCACTAC | lac3-1 amplification |
| lac3-1-R | AATTGGCGCCCTTCGCTTTC | |
| lac3-2-F | TCCGGCTCAATGTCGCAAC | lac3-2 amplification |
| lac3-2-R | ACAATGCCGCTTCTGCTTC | |
| hph-F | CTGTAGGGAGACTGGGTGTG | hph amplification |
| hph-R | CGGTTCCACTACGCGGAG | |
| eh-F | TGCAGTCGCTCGAGCCTACC | egfp-hph amplification |
| eh-R | TGCTGTCCTCATACAAGGAAAAC | |
| egfp-1-F | ACCCTGGTACACACCCCTGAC | egfp-1 amplification |
| egfp-1-R | CACCTTGATGCGCTTCCTTGC | |
| egfp-2-F | GGCCAAGTGACTGTCGGTCCG | egfp-2 amplification |
| egfp-2-R | GCCCTCGTGTTGGGCTTTTC | |
specific primers could be carried out to solve this problem [20]. A suitable Lywallzyme concentration is also important for successful PCR amplification, in order to provide enough for cell breakage but not so much that it causes inhibition of the PCR reaction.
In conclusion, a rapid and effective colony PCR procedure in several mushrooms was established. This method was really a useful tool to screen desired transformants from a large number of colonies at the same time.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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