INTRODUCTION

Chinese dark tea, including Yunnan Pu-erh tea, Hunan Fu-zhuan tea, Hubei Qing-zhuan tea, Sichuan Bian-xiao tea, and Guangxi Liu-bao tea, is a kind of microbial fermented tea in which fermentation allows microorganisms to induce auto-oxidation and nonenzymatic auto-oxidation to complete the process (Lv et al., 2014; Xu et al., 2011). Pu-erh tea (Pu-erh shucha), a traditional Chinese dark tea, has been produced in Yunnan Province of China with a production history of more than a thousand years (Peng et al., 2013). Based on recent researches, Pu-erh tea has definite efficacy on reduction of waist fat buildup and atherosclerosis probability, antioxidation, antibiosis, antiancancer, antidiabetics (Lee & Foo, 2013; Su, Wang, Song, Bai, & Li, 2016), hypolipidemic (Deng, Lin, Shyur, & Lin, 2015; Way et al., 2009), and antirheumatic (Lyu et al., 2013; Nyambe & Williamson, 2016).

Pu-erh tea is normally made from sun-dried green tea leaves (Camellia sinensis var. assamica (JW Masters) Kitamura) by microbial solid-state fermentation (pile fermentation) at high temperature (about 50°C) and high humidity condition (Lv, Zhang, Lin, & Liang, 2013; Zhang et al., 2016). Due to the participation of microorganism, the storage of Pu-erh tea could be regarded as postfermentation.

Correlation analysis between filamentous fungi and chemical compositions in a pu-erh type tea after a long-term storage

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Abstract
Storage environment caused the difference between Jinhua Pu-erh tea (JPT) and General Pu-erh tea. In this study, fungal flora and chemical compositions were analyzed. The results showed that storage environment caused significant (p < .05) differences of theaflavins (TF), theabrownins (TB), tea polyphenols (TP), and water-soluble sugars (WSS), and a highly significant (p < .01) difference of thearubigins (TR). Aspergillus niger, Aspergillus pallidofulvus, Aspergillus sesamicola, and Aspergillus tamarii were isolated from Pu-erh teas and identified based on colony characteristics and ITS, β-tubulin, and calmodulin gene sequences, respectively. A. pallidofulvus, A. sesamicola, and P. manginii were dominant fungi in JPT and generated macroscopic yellow cleistothecia after a long-term storage. Correlation analysis showed that dominant fungi exhibited significantly (p < .05 or p < .01) positive or negative corrections with TF, TB, TP, WSS, TR, and gallic acid. This study revealed dominant fungi including A. pallidofulvus, A. sesamicola, and P. manginii and their effects on given chemical compositions.

KEYWORDS
filamentous fungi, liquid chromatography, metabolites, storage, tea

1 | INTRODUCTION

Chinese dark tea, including Yunnan Pu-erh tea, Hunan Fu-zhuan tea, Hubei Qing-zhuan tea, Sichuan Bian-xiao tea, and Guangxi Liu-bao tea, is a kind of microbial fermented tea in which fermentation allows microorganisms to induce auto-oxidation and nonenzymatic auto-oxidation to complete the process (Lv et al., 2014; Xu et al., 2011). Pu-erh tea (Pu-erh shucha), a traditional Chinese dark tea, has been produced in Yunnan Province of China with a production history of more than a thousand years (Peng et al., 2013). Based on recent researches, Pu-erh tea has definite efficacy on reduction of waist fat buildup and atherosclerosis probability, antioxidation, antibiosis, antiancancer, antidiabetics (Lee & Foo, 2013; Su, Wang, Song, Bai, & Li, 2016), hypolipidemic (Deng, Lin, Shyur, & Lin, 2015; Way et al., 2009), and antirheumatic (Lyu et al., 2013; Nyambe & Williamson, 2016).

Pu-erh tea is normally made from sun-dried green tea leaves (Camellia sinensis var. assamica (JW Masters) Kitamura) by microbial solid-state fermentation (pile fermentation) at high temperature (about 50°C) and high humidity condition (Lv, Zhang, Lin, & Liang, 2013; Zhang et al., 2016). Due to the participation of microorganism, the storage of Pu-erh tea could be regarded as postfermentation.
Pu-erh tea could be stored in natural condition to make pu-erh aged tea after a long-term storage for years, and quality increases with aging time (Xu et al., 2019).

Although Pu-erh tea has been processed, stored, and consumed for hundreds of years, its microbiological properties, particularly fungal community associated with Pu-erh tea storage, are not completely understood. To date, just several actinomycetes and bacteria are detected in Pu-erh tea storage, such as *Actinoplanes aurantiacus*, *Actinoplanes pallidoaurantiacus*, *Actinoplanes purpeobrunneus*, *Streptomyces bacillaris*, and *Streptomyces cinereus* (Chen, Chan, Chang, Liu, & Chen, 2009; Chen, Liu, & Chang, 2010). *Aspergillus* spp. and *Penicillium* spp. are dominant fungi found in pu-erh tea storage (Tian, Zhu, Wu, Wang, & Liu, 2013).

The macroscopic yellow cleistothecia called “golden flowers” are traditionally judged as a quality standard of Fu-zhuan tea (Li et al., 2019). Due to the difference in processing techniques, golden flowers were hardly found in solid-state fermentation of pu-erh tea. However, in a particular storage environment, the golden flowers similar to that in Fu-zhuan tea could appear in Pu-erh tea occasionally named as Jinhua pu-erh tea (JPT) with higher sensory quality, which was chosen in this work to investigate the correlation between filamentous fungi and chemical compositions in the storage. In this paper, five filamentous fungal strains were isolated from pu-erh tea and identified as *Aspergillus niger*, *Aspergillus pallidofulvus*, *Aspergillus sesamica*, *Penicillium manginii*, and *Aspergillus tamari* based on ITS, β-tubulin, and calmodulin gene sequences, respectively. *A. pallidofulvus*, *A. sesamica*, and *P. manginii* were dominant fungi and generated macroscopic yellow cleistothecia after a long-term storage in relative humid environment. Dominant fungi improved tea quality in aroma and taste, and had significant (*p* < .05) impacts on specific chemical compounds during the storage.

### MATERIALS AND METHODS

#### 2.1 Samples

Pu-erh tea samples: JPT and General Pu-erh tea (GPT) were processed and offered by Kunming Dapu Tea CO. Ltd, Yunnan Province, China. Both pu-erh teas were made from identical pu-erh ripen tea (loose tea) after they were autoclaved, compressed,
and dried, and were stored at different natural conditions in Kunming, Yunnan Province, China. The processing and storage of Pu-erh teas are shown in Figure 1. After a long-term storage for about 15 years, macroscopic golden cleistothecia appeared on the surface and in the inside of the Pu-erh tea stored at relative humid environment that was defined as JPT; GPT stored at relative dry environment was defined as control without visible golden flowers.

### 2.2 Chemicals and reagents

Standards of (+)-catechin (C), (−)-epicatechin (EC), (−)-epigallocatechin (EGC), (−)-epicatechin 3-O-gallate (ECG), (−)-epigallocatechin 3-O-gallate (EGCG), gallic acid (GA), and caffeine were purchased from Sigma-Aldrich Co. Ltd. SP fungal DNA Kit, DNA marker, polymerase chain reaction (PCR) spread reagent, and PCR primers, ITS1 (5′-TCCGTAGGTGAACCTGCGG-3′) and ITS4 (5′-TCCTCCGCTTATTGATAGC-3′); Bt2a (5′-GGTAACCAAATCGGTGCTGCTTTC-3′) and Bt2b (5′-ACCCTCAGTGTAGTGACCCTTGGC-3′); and CF1L (5′-GCTGACTCGTTGACCGAAGAG-3′) and CF4 (5′-ATTTTTGCATCATGAGCTGAAC-3′), were purchased from TaKaRa Biotechnology Co. Ltd. Acetonitrile, methanol, and acetic acid for high-performance liquid chromatography (HPLC) were purchased from Mreda Biotechnology Co. Ltd.

### 2.3 Isolation and calculation of filamentous fungi

Tea samples were used to isolate the fungi using potato dextrose agar (PDA) medium, and they were counted by dilution-plating method (Wang, Peng, & Gong, 2011). The colony-forming units were calculated by per dry weight gram of tea leaves after cultivation at 30°C for 2 days (Wang, Gong, Chisti, & Sirisansaneeyakul, 2015).

### 2.4 Fungal identification

The colony morphological characteristics and conidia structure of isolated filamentous fungi were observed after cultivation on PDA medium and Czapek yeast extract agar medium at 25°C for 7 days (Zhou, Ma, Wang, & Xia, 2018). The filamentous fungus grew aero- bically as pure cultures in 20 ml of Czapek Dox medium in 125-ml shake flasks at 30°C, 250 rpm, for 2 days. Fresh cells were obtained by centrifugation at 1,700 g for 5 min and freeze-dried at −80°C (Wang et al., 2015; Zhao et al., 2013). DNA was extracted by using SP Fungal DNA Kit. The fungal primers ITS1 and ITS4, Bt2a and Bt2b, and CF1L and CF4 were used in the PCR to amplify the ITS, β-tubulin, and calmodulin regions, respectively (Abe et al., 2008).

The final volume of 50 μl, containing 1.0 μl of template DNA, 5 μl of 10× buffer, 5 μl of dNTPs (2.5 mM), 0.5 μl of Taq polymerase, 1.0 μl (10 μM) of each primer, and 36.5 μl of sterile distilled water, was used to implement amplifications (Wang et al., 2015; Zhou et al., 2018). The PCR procedure was as follows: pre-degeneration at 95°C for 5 min, degeneration at 94°C for 1 min, annealing at 54°C for 1 min, extension at 72°C for 1.5 min, with 35 cycles, and extension at 72°C for 10 min (Abe et al., 2008). It was stored at 10°C in the end of reaction process.

The PCR was produced in an ABI 3730 Automatic DNA Sequencer (Applied Biosystems) (Abe et al., 2008; Zhou et al., 2018). The received sequences were sent to GenBank of NCBI to seek similar sequences of type strain by using BLASTn (Wang et al., 2015). Multiple sequence alignment was carried out by using Clustal X for Windows. The evolution distance was calculated through a Kimura 2-parameter of the MEGA 4.0 Soft. Neighbor-joining method was used to establish phylogenetic trees.

### 2.5 Sensory evaluation

The sensory panel composed of seven panelists selected from 10 professional tea tasters, and the selection was based on evaluation

### TABLE 1 Sensory evaluation and score of General pu-erh tea (control) and Jinhua pu-erh tea

| Factors          | Sensory evaluation                     | Score | Score |
|------------------|----------------------------------------|-------|-------|
|                  | GPT (control)                          | JPT   |       |
| Appearance (a)   | Normal brick and well-compressed        | Obvious golden flower and well-compressed | 88 ± 1.73 | 87.3 ± 1.53 |
| Liquor color (b) | Bright brownish red                     | Bright thick red                          | 90.3 ± 1.15 | 91.0 ± 2.0 |
| Aroma (c)        | Stale pure flavor                       | Stale and harmonious arohid flavor        | 90.3 ± 1.15 | 94.7 ± 1.53* |
| Taste (d)        | Stale mellow, pure, and thick           | Stale mellow and smooth                    | 87.0 ± 1.0  | 94.0 ± 1.0** |
| Infused leaves (e)| Fat, thick, and even                    | Fat, thick, and even                      | 83.0 ± 1.0  | 83.6 ± 0.58  |
| Total score      | -                                      | 88.0 ± 0.72 | 91.2 ± 1.53* |

Note: Score was present by mean value ± SD of three replications. Total score was the summation of products of factors and score coefficient. Total score = 25%*(a) + 10%*(b) + 25%*(c) + 30%*(d) + 10%*(e).

*Abbreviations: GPT, General pu-erh tea; JPT, Jinhua pu-erh tea.

*There is significant difference in p < .05 level, and **a highly significant difference in p < .01 level using the independent t test of SPSS 20.0.
performance of consistency and reliability (Qin et al., 2013). Sensory evaluation was based on five factors, including appearance (a), liquor color (b), aroma (c), taste (d), and infused leaves (e) according to China National Institute of Standardization (CNIS) GB/T 23776-2018 (Gong et al., 2018). The evaluation procedures were as follows: A total of 100–150 g of tea samples were prepared for the evaluation of appearance; 3 g of samples was infused in 150 ml boiled water for 5 min; and the liquor color, aroma, taste, and infused leaves were evaluated after they were infused, respectively. Total score was estimated as follows:

\[
\text{Total score} = 25\% \times (a) + 10\% \times (b) + 25\% \times (c) + 30\% \times (d) + 10\% \times (e).
\]

2.6 | Determination of chemical compositions

Moisture content and water extract content in each pu-erh tea sample were developed according to CNIS GB/T 8304-2013 (Wang et al., 2013) and GB/T 8305-2013 (Zhou, Xu, Lu, Wang, & Sha, 2013), respectively. Contents of tea polyphenols (TP) and total free amino acids were determined by using the spectra photometric method based on FeSO₄ and the ninhydrin assay (Liang, Zhang, & Lu, 2005), respectively. The main tea pigments, including theaflavins (TF), thearubigins (TR), and theabrownins (TB), were analyzed using the spectra photometric method described by Wang et al. (2011). Water-soluble sugar (WSS) content was determined by visible spectrophotometer method with anthraquinone on 620 nm (Liang et al., 2005). Contents of GA and catechins, including C, EC, EGC, ECG, and EGCG, were determined by HPLC using an Agilent 1200 Series system with Agilent C₁₈ Chromatogram column (250 mm × 4.6 mm, 5 μm) (Wang et al., 2016). Caffeine content was determined by Agilent 1200 HPLC equipment using Agilent C₁₈ Chromatogram column (250 mm × 4.6 mm, 5 μm) with solvent A (100% acetonitrile) and solvent B (0.2% v/v acetic acid–water solution) as mobile phase (Tan et al., 2012).

2.7 | Statistical analysis

Each pu-erh tea sample was measured in three replications. All data are presented as mean value ± standard deviation (SD). The independent t test using SPSS 20.0 for Windows was carried out to determine whether the significant difference at \( p < .05 \) level or the highly significant difference at \( p < .01 \) level exists between JPT and GPT (control). Bivariate correlation analysis using SPSS 20.0 for Windows was carried out to determine effects of each dominant fungus on chemical composition after a long-term storage.

### TABLE 2 A comparisons of the chemical components in General pu-erh tea (control) and Jinhua pu-erh tea

| Samples | General pu-erh tea (control) | Jinhua pu-erh tea |
|---------|-----------------------------|------------------|
| Moisture content (%) | 9.59 ± 0.48 | 10.55 ± 0.26* |
| Water extract content (%) | 34.00 ± 1.51 | 36.33 ± 0.86 |
| TF (mg/g) | 3.63 ± 0.208 | 4.37 ± 0.208* |
| TR (mg/g) | 36.43 ± 2.22 | 44.13 ± 1.21** |
| TB (mg/g) | 131.7 ± 7.15 | 107.1 ± 11.13* |
| TP (mg/g) | 85.73 ± 6.95 | 70.73 ± 2.15* |
| C (mg/g) | 4.44 ± 1.14 | 4.14 ± 1.06 |
| EC (mg/g) | 13.7 ± 1.16 | 12.61 ± 0.98 |
| EGC (mg/g) | 14.99 ± 1.32 | 13.67 ± 0.91 |
| ECG (mg/g) | 10.33 ± 2.89 | 6.16 ± 1.11 |
| EGC (mg/g) | ND | ND |
| Total catechins (mg/g) | 43.46 ± 4.18 | 36.6 ± 3.96 |
| GA (mg/g) | 12.81 ± 1.27 | 15.04 ± 1.36 |
| Caffeine (mg/g) | 38.42 ± 1.09 | 38.93 ± 0.76 |
| WSS (mg/g) | 33.40 ± 2.10 | 40.93 ± 2.17* |
| Free amino acids (mg/g) | 6.43 ± 0.78 | 6.07 ± 1.17 |

Note: The content of total catechins was the summation of C, EC, EGC, ECG, and EGCG. All data were present by mean value ± SD of three replications.

Abbreviations: C, (+)-catechin; EC, (−)-epicatechin; EGC, (−)-epicatechin 3-O-gallate; ECG, (−)-epicatechin 3-O-gallate; GA, gallic acid; ND, not detected; TB, theabrownins; TF, theaflavins; TR, thearubigins; WSS, water-soluble sugars.

* There is a significant difference at \( p < .05 \) levels, and **A highly significant difference at \( p < .01 \) levels using the independent t test of SPSS 20.0.

### TABLE 3 Colony characteristics of filamentous fungi isolated from pu-erh teas

| Isolate | Shape | Surface | Color | Exudates | Cleistothecia |
|---------|-------|---------|-------|----------|---------------|
| GPTS1   | Circular | Rough  | Black | None     | None          |
| JPTS1   | Circular | Rough  | Dark yellow colonies with white edges | Yellow sclerotium | Yellow |
| JPTS2   | Irregular | Rough  | Light yellow | Yellow sclerotium | Golden |
| JPTS3   | Circular | Rough  | Greyish-green center with yellow patches | Red pigment | Yellow |
| GPTS2   | Irregular | Rough  | Hazel green with gray back | None | None |
RESULTS

3.1 Quality characteristic analysis of different pu-erh teas

Quality characteristics were calculated by sensory evaluation based on five sensory factors. Significant difference analysis was carried out by using independent $t$ test of SPSS 20.0. Results of sensory evaluation in GPT (control) and JPT are shown in Table 1. Due to both pu-erh teas made from identical pu-erh ripen tea (loose tea) after they were autoclaved, compressed, and dried, quality characteristics in appearance, liquor color, and infused leaves were similar with no significant differences ($p > .05$). However, due to the difference in storage environment, particularly the appearance of macroscopic golden cleistothecia, JPT had better aroma ($p < .05$) and taste ($p < .01$). Therefore, storage environment had significant ($p < .05$) impact on aroma and taste.

3.2 Differences in chemical compositions between GPT and JPT

To better understand the effect of storage environment on quality components, with GPT as control, main chemical components were determined in three replications. A comparison of isolated chemical components is presented in Table 2. The relative humid environment enhanced moisture content significantly ($p < .05$) in JPT. Compared with the control, the results showed significant ($p < .05$) difference of TF, TB, TP, and WSS contents and a highly significant difference...
of TR content in JPT, which illustrated that storage environment had significant ($p < .05$) impact on TF, TB, TP, and WSS contents, and a highly significant ($p < .01$) impact on TR, while it had no significant ($p > .05$) impacts on water extract content, GA, free amino acids, total catechins, and each catechin contents. The macroscopic golden cleistothecia in JPT enhanced WSS and TF contents significantly ($p < .05$) and TR content highly significantly ($p < .01$), while it decreased TP and TB contents significantly ($p < .05$). We speculated that filamentous fungi caused significant differences in given chemical compositions.

3.3 Biological identification of filamentous fungi
In this paper, about nine different fungal strains were isolated from pu-erh teas. Five filamentous fungi could be detected with a high abundance and named as GPTS1, JPTS1, JPTS3, and JPTS2, and GPTS2, respectively, which were superior in numbers. Others were occasionally detected in pu-erh teas, which belonged to the genera *Aspergillus*, *Penicillium*, and *Saccharomyces* according to colony morphological characteristics and conidia structures. The colony morphological characteristics of isolated filamentous fungi are shown in Figure 2.
in Table 3. Strain GPTS1 was identified as A. niger based on colony characteristics and conidial structure, which has been described in our previous study (Zhou et al., 2018). Strains JPTS1, JPTS2, and JPTS3 could generate yellow or golden cleistothecia in a particular state. PCR method was used to amplify the ITS, β-tubulin, and calmodulin regions, respectively. Based on PCR-amplified sequences (Figures S1–S4), phylogenetic trees of isolated filamentous fungi are built and shown in Figure 2. In the phylogram for Aspergillus species, strain JPTS1 was clustered with A. pallidofulvus and showed a 99.9% of identity to the tested strain NRRL4789 (NR137468), strain JPTS2 was clustered with A. sesamica and showed a 99.8% of identity to the tested strain CBS137324 (KJ775437), and strain GPTS2 was clustered with A. tamarii and showed a 99.9% of identity to the tested strain NRRL20818 (AF433058). In the phylogram for Penicillium species, strain JPTS3 was identified as P. manginii and showed a 99.6% of identity to the tested strain CBS253.31 (MH855205). In conclusion, five filamentous fungi were isolated from pu-erh teas and identified as A. niger, A. pallidofulvus, A. sesamica, A. tamarii, and P. manginii, respectively. Among them, A. pallidofulvus, A. sesamica, and P. manginii could generate macroscopic cleistothecia that were similar to the golden flowers appearing in JPT.

3.4 | Distribution of fungal consortium in pu-erh teas

The distribution and difference of fungal consortium are presented in Figure 3. The distribution of fungal consortium in JPT had obvious difference than in control (Figure 3a). All isolated filamentous fungi were detected in JPT, which occupied about 95.38% in total fungi count. However, P. manginii was not detected in GPT (control), and the isolated filamentous fungi occupied about 82.25% in count. Fungal flora in JPT had significant \( p < .05 \) increase due to a high moisture content (Figure 3b). Particularly, A. pallidofulvus and A. sesamica had a significant \( p < .05 \) or a highly significant \( p < .01 \) increase in JPT, respectively. Only A. niger had a significant \( p < .05 \) decrease in JPT. The fungal flora, particularly the dominant fungi found in JPT including A. pallidofulvus, A. sesamica, and P. manginii, caused significant differences of given chemical components in JPT after a long-term storage. Correlation analysis was conducted later to investigate the relationship between each isolated filamentous fungus and chemical components, that is TF, TR, TB, TP, GA, and WSS, respectively.

3.5 | Effects of isolated filamentous fungi on chemical compositions

In this paper, A. niger, A. pallidofulvus, A. sesamica, A. tamarii, and P. manginii were isolated from pu-erh teas. Particularly, A. pallidofulvus, A. sesamica, and P. manginii were dominating fungi in JPT, which generate yellow or golden color cleistothecia in the storage. The relationships between each isolated filamentous fungus and chemical compound were evaluated by bivariate correlation analysis using SPSS 20.0 for Windows. The results are presented in Table 4. Table 4 shows that moisture content in pu-erh tea had significantly \( p < .05 \) positive correlations with A. pallidofulvus \( r = .830 \) and P. manginii \( r = .840 \), respectively. However, fungal community in pu-erh tea storage had no significant \( p > .05 \) correlations with water extract content, caffeine, free amino acids, and catechins including C, EC, EGC, ECG, and EGCG, respectively. The isolated filamentous fungi had positive or negative correlations with TP, TF, TR, TB, GA, and WSS. Specifically, A. niger had a significantly \( p < .05 \) positive correlation with TP \( r = .819 \), a significantly \( p < .05 \) negative correlation with WSS \( r = −.827 \), and a highly significantly \( p < .01 \) negative correlation with TR \( r = .931 \), respectively. A. pallidofulvus and P. manginii had highly significantly \( p < .01 \) positive correlations with WSS \( r = .921 \) and 0.923, respectively, which enhanced WSS content through degradation of water-insoluble cellulose. Conversely, A. sesamica had a highly significantly \( p < .01 \) negative correlation with TB \( r = −.929 \) and a significantly \( p < .05 \) positive correlation with GA \( r = .834 \), respectively, which caused significant \( p < .05 \) decrease in TB and a slight increase in GA in JPT. A. tamarii had no significant \( p > .05 \) impact on given chemical components. Therefore, A. niger, A. pallidofulvus, A. sesamica, and P. manginii caused significant \( p < .05 \) or highly significant \( p < .01 \) differences of TP, TF, TR, TB, and WSS in JPT after a long-term storage.

4 | DISCUSSION

Solid-state fermentation plays an important role in the processing of pu-erh tea (Abe et al., 2008). Fungi have profound impact on substance metabolisms and show correlation with quality formation of pu-erh tea (Zhao et al., 2013, 2019). So far, A. niger, Aspergillus tubingensis, Aspergillus fumigatus, Aspergillus acidus, Aspergillus awamori, A. tamarii, Aspergillus sydowii, Blastobotrys adeninivorans, Candida tropicalis, Fusarium graminearum, Pichia farinosa, Rasamsonia byssochlamydoides, Rasamsonia emersonii, Rasamsonia cylindrospora, Rhizomucor pusillus, Rhizomucor tauricus, and Thermomyces lanuginosus have been detected in solid-state fermentation of pu-erh tea as dominant fungi (Wang et al., 2011; Zhang et al., 2016; Zhao et al., 2015; Zhou et al., 2018). In this study, five filamentous fungi were isolated from pu-erh teas and identified as A. niger, A. pallidofulvus, A. sesamica, A. tamarii, and P. manginii, respectively. The detection of A. niger and Aspergillus tamarii in pu-erh tea storage has a close connection with solid-state fermentation of pu-erh tea.

The strain generating yellow color cleistothecia is termed as the “golden flower fungus” (Mo, Zhu, & Chen, 2008; Mao, Wei, Teng, Huang, & Xia, 2017). The species within the genera Aspergillus, Eurotium, and Penicillium are main fungal taxa isolated from the postfermentation of dark tea (Mo et al., 2008). Aspergillus cristatus and Eurotium cristatum are golden flower fungi involved in the fermentation of Chinese fu-zhuan tea (Ge et al., 2016; Zou et al., 2014). Numerous macroscopic yellow cleistothecia similar to that in other dark tea also could appear in pu-erh tea after a long-term
Distribution (a) and differences (b) of isolated filamentous fungi in pu-erh tea type. GPT, General pu-erh tea. JPT, Jinhua pu-erh tea. About nine different kinds of fungal strains were isolated from pu-erh tea samples based on PDA medium, and those colony-forming units were calculated. Five filamentous fungi were isolated and identified. *Aspergillus pallidofulvus*, *Aspergillus sesamicoa*, and *P. manginii* were dominant fungi in count and generated macroscopic golden cleistothecia after a long-term storage in relative humid environment. Others were insignificant fungal strains in count, which belonged to the genera *Aspergillus*, *Penicillium*, and *Saccharomyces* according to the colony morphological characteristics and conidia structures. * indicates there is a significant difference at \( p < .05 \) level, and ** indicates a highly significant difference at \( p < .01 \) level using the independent \( t \) test of SPSS 20.0.

### TABLE 4: Correlation coefficients of each isolate strain and chemical components

| Indicators          | Aspergillus niger | Aspergillus pallidofulvus | Aspergillus sesamicola | Penicillium manginii | Aspergillus tamarii | Others |
|---------------------|-------------------|---------------------------|------------------------|----------------------|---------------------|--------|
| Moisture content    | 0.544             | 0.830*                    | 0.686                  | 0.840*               | 0.230               | −0.417 |
| Water extract content| −0.784           | 0.785                     | 0.568                  | 0.810                | −0.037              | −0.229 |
| TF                  | −0.762            | 0.834*                    | 0.876*                 | 0.851*               | 0.236               | 0.389  |
| TR                  | −0.931**          | 0.851*                    | 0.804*                 | 0.806                | −0.079              | 0.416  |
| TB                  | 0.561             | −0.702                    | −0.929**               | −0.722               | −0.076              | −0.163 |
| TP                  | 0.819*            | −0.773                    | −0.801                 | −0.671               | 0.022               | −0.693 |
| C                   | −0.134            | −0.273                    | −0.097                 | −0.501               | 0.396               | −0.812*|
| EC                  | 0.186             | −0.448                    | −0.668                 | −0.595               | −0.361              | 0.250  |
| EGC                 | 0.279             | −0.592                    | −0.668                 | −0.711               | −0.636              | 0.096  |
| ECG                 | 0.702             | −0.636                    | −0.728                 | −0.563               | 0.029               | −0.733 |
| Total catechins     | 0.563             | −0.519                    | −0.781                 | −0.446               | 0.173               | −0.703 |
| GA                  | −0.409            | 0.527                     | 0.834*                 | 0.482                | −0.246              | −0.018 |
| Caffeine            | 0.059             | 0.243                     | 0.469                  | −0.018               | −0.154              | −0.061 |
| WSS                 | −0.827*           | 0.921**                   | 0.772                  | 0.923**              | 0.215               | −0.016 |
| Free amino acids    | 0.224             | −0.463                    | −0.068                 | −0.576               | −0.732              | 0.550  |

Abbreviations: C, (+)-catechin; EC, (−)-epicatechin; ECG, (−)-epicatechin 3-O-gallate; EGC, (−)-epigallocatechin; EGGC, (−)-epigallocatechin 3-O-gallate; GA, gallic acid; TB, theabrownins; TF, theaflavins; TP, tea polyphenols; TR, thearubigins; WSS, water-soluble sugars.

*A significant correlation at \( p < .05 \) level, **A highly significant correlation at \( p < .01 \) level.
storage, which was defined as JPT. With GPT made from the same pu-erh ripen tea (loose tea) with identical processing as the control, through comparison of fungal consortium, A. pallidofulvus, A. sesamica, and P. manginii were dominant fungi found in JPT and generate the appearance of macroscopic yellow cleistothecia after a long-term storage, which had a significantly ($p < .05$) positive correlation with moisture content in pu-erh tea cake.

The individual numbers of fungi in pu-erh tea decreased significantly after solid-state fermentation through autoclaving, compressing, and drying (Tian et al., 2013). The fungal flora is maintained at a relatively low level. Due to a long-term storage for years, dominant fungi also had significant impact on chemical compositions during the storage, which caused significant ($p < .05$) differences of TF, TB, TP, and WSS contents, and a highly significant ($p < .01$) difference of TR between JPT and GPT (Table 2). Under effects of dominant fungi, TP could be oxidized to TF and TR, and TB had a significant decrease during the storage, which were different from the changes of chemical composition contents in solid-state fermentation of pu-erh tea (Chen, Cui, Li, Sheng, & Lv, 2013).

Aspergillus spp. including A. tubingensis, A. fumigatus, and A. marvanovae could convert TP to TB in solid-state fermentation and submerged fermentation (Wang, Gong, Chisti, & Sirisansaneeyakul, 2014; Wang et al., 2015). A. niger and A. fumigatus could degrade ester catechins into nonester catechins and GA during solid-state fermentation (Qin, Li, Tu, Ma, & Zhang, 2012). Additionally, A. sydowii could convert caffeine to theophylline and had significant impacts on WSS and free amino acid contents (Zhou, Ma, Ren, Xia, & Li, 2020; Zhou et al., 2019). Effects of each filamentous fungus on chemical compositions in the storage were investigated through the bivariate correlation analysis (Table 4). The results exhibited that the isolated dominant fungi had significant ($p < .05$) or highly significant ($p < .01$) impacts on given chemical compositions, including TP, TF, TR, TB, WSS, and GA after a long-term storage.

5 | CONCLUSIONS

Our present work describes fungal diversity and differences in chemical compositions between GPT and JPT to study effects of dominant fungi in pu-erh tea storage. Due to the appearance of macroscopic yellow cleistothecia, JPT had better aroma ($p < .05$) and taste ($p < .01$) (Table 1) and had significant ($p < .05$) differences of TF, TB, TP, and WSS contents, and a highly significant ($p < .01$) difference of TR compared with the control (Table 2). A. niger, A. pallidofulvus, A. sesamica, A. tamarii, and P. manginii were isolated from pu-erh teas and identified. A. pallidofulvus, A. sesamica, and P. manginii were dominant fungi and generated macroscopic yellow cleistothecia in JPT after a long-term storage. The difference in fungal community should be ascribed to moisture content in tea cake and storage environment. Bivariate correlation analysis showed that the filamentous fungi had significantly positive or negative correlations with the differences of TF, TB, TP, WSS, and TR contents (Table 4). Particularly, A. pallidofulvus and P. manginii exhibited a highly significantly positive correlation ($p < .01$) with WSS and a significantly ($p < .05$) positive correlation with TF, while A. sesamica showed a highly significantly negative correlation ($p < .01$) with TB and a significantly ($p < .05$) positive correlation with GA, respectively. Those results indicated that the isolated filamentous fungi had significant impact on given chemical compositions in the storage, which provides reference for the application of A. pallidofulvus, A. sesamica, and P. manginii in tea.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

AUTHOR CONTRIBUTIONS

B. Zhou and C. Ma designed the research and wrote the paper. C. Ma and C. Zheng participated in the performance of the research. X. Liu conducted the statistical analysis. T. Xia contributed to the writing and data analysis. All authors read and approved the final manuscript.

ETHICAL APPROVAL

The study did not involve any human or animal testing.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the
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