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28 Province in order to find the difference of dominant fungi among the four regions.  
29 Through high throughput sequencing we detected *Ascomycota* accounts for absolute  
30 dominant phylum; *Dothideomycetes*, *Sordariomycetes*, *Tremellomycetes*,  
31 *Microbotryomycetes*, and *Eurotiomycetes* were dominant classes; *Capnodiales*,  
32 *Hypocreales*, and *Pleosporales* were the main orders; *Cladosporiaceae*,  
33 *Pleosporaceae*, *Nectriaceae*, *Clavicipitaceae*, *Tremellaceae*, *Phaeosphaeriaceae*,  
34 *Trimorphomycetaceae*, *Sporidiobolaceae*, *Bionectriaceae*, and *Trichocomaceae* were  
35 major family; *Cladosporium*, *Epicoccum*, *Fusarium*, and *Alternaria* were the most  
36 abundant phylotypes at genus level; *Epicoccum\_nigrum*, *Gibberella\_zeae*, and  
37 *Fusarium\_proliferatum* were the dominant fungal species. Great fungal diversity was  
38 observed in the rice samples harvested in the four major Japonica rice-growing  
39 regions in Heilongjiang province. However, no significant difference in diversity was  
40 observed among the four regions, likely due to the relatively close geographical  
41 proximity leading to very similar climatic conditions. Since some of the fungal  
42 species produce mycotoxins, it is necessary to take precautions to ensure the rice is  
43 stored under safe conditions to prevent the production of mycotoxins. This is the first  
44 report on investigation of field fungal diversity in freshly harvested Japonica rice in  
45 Heilongjiang Province in China.

46

47 **Keywords:** Field fungi; diversity; japonica rice; high through-put sequencing

48

## 49 1. INTRODUCTION

50 Rice (*Oryza sativa L.*) is a major food crop in China and more than 65% of the  
51 populace consumes rice as staple food. China ranks No.1 in total annual rice  
52 production in the world and accounts for around 1/3 of the global paddy rice  
53 production [1]. Heilongjiang province ranks 1st in Japonica rice cultivation in China  
54 with total production of 30 million tons in 2018 [2]. In addition, rice cultivated in  
55 Heilongjiang province is very famous for its high quality and excellent flavor due to  
56 optimal environmental conditions suitable for rice growing. The rice produced in this  
57 province is well received throughout the country and is even exported to many  
58 regions around the world. However, rice is prone to invasion by fungi and  
59 contamination by their mycotoxins. Fungi play a key role in rice safety and  
60 understanding the fungi community structure is of great importance when taking  
61 appropriate measures to ensure rice safety. Fungi infect rice crops early in the field

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62 and may produce mycotoxins during this period. Consumers are concerned about this  
63 issue and consequently it is necessary to investigate the status of fungi contamination  
64 of rice in Heilongjiang province. Rice is widely cultivated in China under different  
65 climatic conditions and is extensively contaminated by various fungi. However, little  
66 information is currently available on the fungal diversity of field fungi, especially  
67 aflatoxigenic fungal contamination of rice in the main cultivating regions of  
68 Heilongjiang province. The objective of this study was to investigate fungal diversity  
69 of freshly harvested rice in the four main cultivation regions of Heilongjiang province  
70 through high-throughput sequencing and FUNGuild in order to find the abundance  
71 difference of dominant fungi among the four regions.

72

## 73 **2. MATERIALS AND METHODS**

### 74 **2.1 Materials**

75 Twelve rice samples were harvested from four regions in Heilongjiang's major  
76 rice cultivation areas as indicated in Fig. 1: Wuchang city (three samples of rice  
77 specie Daohuaxiang No 2, three repetitions, marked as WC-1, WC-2, WC-3, WC-4,  
78 WC-5, WC-6, WC-7, WC-8, and WC-9), Jiamusi city (three samples of rice specie  
79 Longjing No. 31, three repetitions, marked as JMS-1, JMS-2, JMS-3, JMS-4, JMS-5,  
80 JMS-6, JMS-7, JMS-8, and JMS-9), Zhaoyuan county (three samples of rice specie  
81 Songjing No. 3, three repetitions, marked as ZY-1, ZY-2, ZY-3, ZY-4, ZY-5, ZY-6,  
82 ZY-7, ZY-8, and ZY-9), and Tailai county (three samples of rice specie Songjing No. 9,  
83 three repetitions, marked as TL-1, TL-2, TL-3, TL-4, TL-5, TL-6, TL-7, TL-8, and  
84 TL-9). During September 26-29 of 2017, around 2 kg of each sample was cut using a  
85 reaping hook from rice fields, put into plastic bags, and sealed tightly. After arriving  
86 at the lab, 30 spikes of rice were manual threshed from each sample and three 50 g  
87 paddy rice samples were weighed from each sample into 1000 mL Erlenmeyer flasks  
88 with a 500 mL PBS buffer ( $\text{KH}_2\text{PO}_4$  0.27 g,  $\text{NA}_2\text{HPO}_4$  1.42 g, NaCl 8 g, KCl 0.2 g,  
89 diluted with 800 mL distilled water, adjusted pH value of 7.4, constant volume of 1 L ,  
90 and sterilized). They were labeled as three replicates of one sample. These samples  
91 were shaken with a vibrator for 30 minutes, subjected to sucking filtration, and  
92 filtered through 0.45  $\mu\text{m}$  water membranes. The residues were collected from the  
93 membranes using medicinal ladles and transferred into 1 mL microcentrifuge tubes  
94 and preserved by cryopreservation using liquid nitrogen. All operations were  
95 conducted in a sterile room and masks and gloves were worn to guarantee the samples

96 would not be contaminated by environmental fungi. The samples were then  
97 transported to Guangzhou Gene Denovo Bio-Tech Ltd. Co. (Guangzhou, China)  
98 under dry ice conditions to perform high throughput sequencing of the PCR products.  
99 The obtained data was assembled into sequence tags and subject to BLAST in  
100 GenBank for microbe classification, followed by OTU, and diversity and inter-sample  
101 comparative analyses.

102

103



104

105 Fig. 1 Distribution of samples collecting locations in Heilongjiang province of China.  
106 a. Illustration of the geographical location of Heilongjiang province in China. b.  
107 Distribution of samples collecting locations in Heilongjiang province

108

## 109 2.2 DNA extraction and PCR amplification

110 Microbial DNA was extracted from stool samples using the E.Z.N.A. stool DNA  
111 Kit (Omega Bio-Tek, Norcross, GA, U.S.) according to manufacturer's protocols. The  
112 ITS region of the Eukaryotic ribosomal RNA gene was amplified by PCR( 95 °C for 2  
113 min, followed by 27 cycles at 98 °C for 10 s, 62 °C for 30 s, and 68 °C for 30 s and a  
114 final extension at 68 °C for 10 min ) using primers ITS3\_KYO2F 5'-  
115 GATGAAGAACGYAGYRAA -3' and ITS4R 5'- TCCTCCGCTTATTGATATGC-3',  
116 where the barcode is an eight-base sequence unique to each sample. PCR reactions  
117 were performed in triplicate 50 µL mixture containing 5 µL of 10 × KOD Buffer, 5 µL  
118 of 2.5 mM dNTPs, 1.5 µL of each primer (5 µM), 1 µL of KOD Polymerase, and 100  
119 ng of template DNA.

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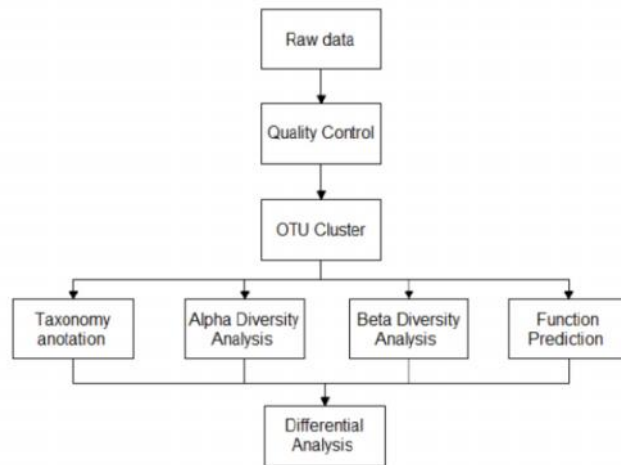
## 120 2.3 Illumina Hiseq2500 sequencing

121 Amplicons were extracted from 2% agarose gels and purified using the AxyPrep  
122 DNA Gel Extraction Kit (Axygen Biosciences, Union City, CA, U.S.) according to  
123 the manufacturer's instructions and quantified using QuantiFluor™-ST (Promega,  
124 U.S.). Purified amplicons were pooled in equimolar and paired-end sequenced (2 ×  
125 250) on an Illumina platform according to the standard protocols.

## 126 2.4 Bioinformatics analysis

127 Bioinformatics analysis was conducted according to Fig.2.

128



ITS analysis flow chart

129

130

Fig.2 ITS analysis flow chart

131

## 132 2.5 Quality control and reads assembly

### 133 2.5.1 Reads filtering

134 Raw data containing adapters or low-quality reads would affect the following  
135 assembly and analysis. Hence, to get high-quality clean reads, raw reads were further  
136 filtered according to the following rules:

137 1) Removing reads containing more than 10% of unknown nucleotides (N);

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138 2) Removing reads containing less than 80% of bases with quality (Q-value) > 20.

139 **2.5.2 Reads assembly**

140 Paired-end clean reads were merged as raw tags using FLSAH [3] (v 1.2.11) with  
141 a minimum overlap of 10 bp and mismatch error rates of 2%.

142 **2.5.3 Raw tag filtering**

143 Noisy sequences of raw tags were filtered by QIIME [4] (V1.9.1) pipeline under  
144 specific filtering conditions [5] to obtain high-quality clean tags.

145 **2.5.4 Chimera checking and removal**

146 Clean tags were searched against the reference database  
147 ([http://drive5.com/uchime/uchime\\_download.html](http://drive5.com/uchime/uchime_download.html)) to perform Reference-based  
148 chimera checking using UCHIME algorithm

149 ([http://www.drive5.com/usearch/manual/uchime\\_algo.html](http://www.drive5.com/usearch/manual/uchime_algo.html)). All chimeric tags were  
150 removed and finally obtained effective tags for further analysis.

151 **2.5.5 OTU cluster**

152 The effective tags were clustered into operational taxonomic units (OTUs)  
153 of  $\geq 97$  % similarity using UPARSE [6] pipeline. The tag sequence with the highest  
154 abundance was selected as a representative sequence within each cluster. Between  
155 groups, Venn analysis was performed in R to identify unique and common OTUs.

156 **2.5.6 Taxonomy classification**

157 The representative sequences were classified into organisms by a naive Bayesian  
158 model using RDP classifier [7] (Version 2.2) based on UNITE [8] Database  
159 (<https://unite.ut.ee/>). The abundance statistics of each taxonomy and a phylogenetic  
160 tree was constructed in a Perl script and visualized using SVG. Biomarker features in  
161 each group were screened by Metastats and LEfSe software.

162 **2.5.7 Alpha diversity analysis**

163 Chao1, Simpson and all other alpha diversity indices were calculated in QIIME.  
164 OTU rarefaction curve and Rank abundance curves were plotted in QIIME. Statistics  
165 of between-group Alpha index comparison was calculated by a Welch's t-test and a  
166 Wilcoxon rank test in R. Alpha index comparisons among groups was computed by a

167 Tukey's HSD test and a Kruskal-Wallis H test in R.

### 168 **2.5.8 Beta diversity analysis**

169 The weighted and unweighted unifrac distance matrix was generated by QIIME.  
170 Multivariate statistical techniques including PCA, principal coordinates analysis  
171 (PCoA) and NMDS of (Un)weighted unifrac distances were calculated and plotted in  
172 R. Statistics of Welch's t-test, Wilcoxon rank test Tukey's HSD test, Kruskal-Wallis H  
173 test, Adonis (also called Permanova) and Anosim test was calculated using R.

### 174 **2.5.9 Functional prediction**

175 The functional group (guild) of the OTUs was inferred using FUNGuild [9]  
176 (v1.0).

## 177 **3. RESULTS AND DISCUSSION**

### 178 **3.1 Fungal diversity and richness in single rice samples and comparison of these** 179 **indexes among the four regions**

180 Total fungal ITS tags (106951, 108190, 111294, 105520, 108264, 113010,  
181 115999, 104584, 108716, 123025, 108835, and 119401) were recovered from 12  
182 (Rice 1, Rice 2, Rice 3, Rice 4, Rice 5, Rice 6, Rice 7, Rice 8, Rice 9, Rice 10, Rice  
183 11, and Rice 12) samples, respectively. The library samples were then clustered into  
184 fungal Operational Taxonomic Units (OTUs) at 97% similarity (Table 1).

185

186 Table 1 Community richness, diversity and coverage indexes for the four regions\*

Region	OTU	Chao 1	Ace	Good	Shannon	Simpson
				Coverage (%)		
Wuchang	510±53 <sup>a</sup>	710±49 <sup>a</sup>	709±55 <sup>a</sup>	99.8±0.0 <sup>a</sup>	3.49±0.53 <sup>a</sup>	0.76±0.07 <sup>a</sup>
Jiamusi	379±45 <sup>b</sup>	531±72 <sup>b</sup>	541±77 <sup>b</sup>	99.8±0.0 <sup>a</sup>	3.24±0.34 <sup>a</sup>	0.75±0.07 <sup>a</sup>
Zhaoyuan	396±53 <sup>a</sup>	592±102 <sup>b</sup>	585±90 <sup>b</sup>	99.8±0.0 <sup>a</sup>	3.33±0.36 <sup>a</sup>	0.76±0.04 <sup>a</sup>
Tailai	383±64 <sup>a</sup>	636±87 <sup>b</sup>	537±87 <sup>b</sup>	99.8±0.0 <sup>a</sup>	3.43±0.30 <sup>a</sup>	0.78±0.06 <sup>a</sup>

187 Data represents mean±SD. Data followed by the same superscript letter in the same  
188 column are not significantly different.



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189

190 The Chao and ACE are abundance indexes; the Simpson and Shannon are  
191 diversity indexes. Higher values of Chao (richness estimate) and ACE indicate more  
192 community richness. The Shannon (diversity index) and Simpson values indicate the  
193 community diversity, and higher Shannon and Simpson values indicate greater  
194 community diversity. The good coverage value indicates the depth of sequencing. The  
195 good sequencing coverage in all the four regions almost reached 99.8%, which  
196 indicated that almost all fungi have been detected. The number of OTUs determined  
197 in the four regions showed that Wuchang got the maximum value, whereas Jiamusi  
198 obtained the minimum value. An OTU is usually recognized as a genus of  
199 microorganism. Consequently, a total of 507, 564, 459, 374, 398, 366, 440, 409, 338,  
200 443, 316, and 390 fungal genera were identified in the 12 rice samples, respectively.

201 To compare fungal diversity and richness among the four regions, data was  
202 statistically analyzed and presented in Table 1. As seen in the table, no significant  
203 difference was found in the five indexes of the four regions. This is probably due to a  
204 close physical proximity among the four regions resulting in a lack of significant  
205 differences in environmental conditions. Since Jiamusi city is around 2° latitude north  
206 of the other three regions and greater than 1 °C of daily average minimum and  
207 maximum temperature lower than the other three regions, the observed numbers of  
208 fungal genera were the lowest as a result of cooler temperatures.

209

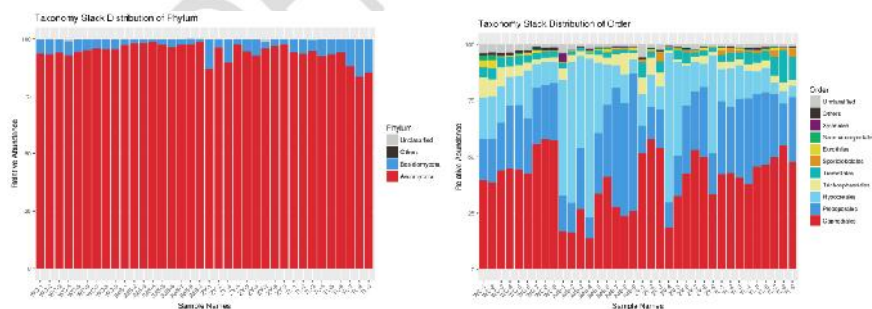
### 210 **3.2 Fungal community composition**

211 For the 12 rice samples, fungal community compositions were detected at seven  
212 levels: Domain, Phylum, Class, Order, Family, Genus, and Species. Fig. 3  
213 demonstrates the taxonomy stack distribution of the Phylum, Order, Genus, and  
214 Species of the identified fungi. Due to limited space, only the results of fungi  
215 community compositions at the Phylum, Order, Genus, and Species levels were  
216 presented in the Figure. Of the classifiable sequences, two Phyla were identified as  
217 seen in Fig. 3a: *Ascomycota* and *Basidiomycota*, in which *Ascomycota* accounts for  
218 absolute dominance. At Class level, *Dothideomycetes*, *Sordariomycetes*,  
219 *Tremellomycetes*, *Microbotryomycetes*, *Eurotiomycetes*, and *Saccharomycetes* were  
220 identified, where *Dothideomycetes* and *Sordariomycetes* account for absolute  
221 dominance. At Order level, *Capnodiales*, *Pleosporales*, *Hypocreales*, *Tremellales*,  
222 *Trichosphaeriales*, *Sporidiobolales*, and *Eurotiales* were determined, in which

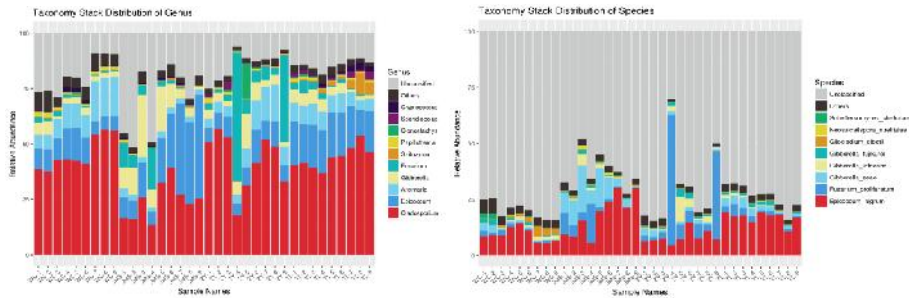


223 *Capnodiales*, *Hypocreales*, and *Pleosporales* were dominant (Fig. 3b). At Family  
 224 level, *Cladosporiaceae*, *Pleosporaceae*, *Nectriaceae*, *Clavicipitaceae*, *Tremellaceae*,  
 225 *Phaeosphaeriaceae*, *Trimorphomycetaceae*, *Sporidiobolaceae*, *Bionectriaceae*, and  
 226 *Trichocomaceae* were detected, and *Cladosporiaceae*, *Pleosporaceae* as well as  
 227 *Nectriaceae* are the dominant families (data not shown in Fig. 3). As seen in Fig. 3c,  
 228 *Cladosporium* is the absolute dominant genus; followed by *Epicoccum*, *Alternaria*,  
 229 *Gibberella*, and *Fusarium*; they are less abundant phylotypes at the genus level  
 230 detected in these rice samples. From Fig. 3d, it can be observed that unclassified  
 231 species account for large portions of the whole bars. This is due to limitations of the  
 232 UNITE Database which prevents classification of large amounts of species.  
 233 *Epicoccum\_nigrum*, *Gibberella\_zeae*, and *Fusarium\_proliferatum* are the dominant  
 234 fungal species found in the determined samples. High proportions of  
 235 *Fusarium\_proliferatum* were detected in two samples (ZY-4 and ZY-9). *Gibberella*  
 236 *zeae*, usually known by the name of its anamorph *Fusarium\_graminearum*, is  
 237 identified as a plant pathogen which causes *Fusarium* head blight and can produce  
 238 toxins, particularly deoxynivalenol (DON). *Fusarium\_spp.* produces a diverse number  
 239 of secondary metabolites, including some fatal mycotoxins [10], and they are attributed  
 240 as the most important toxigenic fungi in the Northern temperate areas [11]. Among  
 241 the *Fusarium\_spp.* isolated from rice, *F. proliferatum* and *F. verticillioides* were proven  
 242 to be the most abundant Fumonisin producers [12]. *Fusarium\_proliferatum* can occur  
 243 in a wide range of plants, including rice and produce mycotoxins such as fumonisin  
 244 [12,13].

245



246



247

248 Fig.3 Taxonomy stack distribution of genus and species of the 12 rice samples in Heilongjiang  
 249 province. a. At Phylum level; b. At Order level; c. At Genus level; d. At Species level.

250

251 Through a naive Bayesian model using RDP classifiers based on UNITE  
 252 Database analysis of the assembled sequences, it was found that in the rice samples  
 253 *Epicoccum nigrum* and *Fusarium proliferatum* were dominant, where *Epicoccum*  
 254 *nigrum* is a plant pathogen and endophyte. *Fusarium proliferatum* is a fungal plant  
 255 pathogen and usually infects asparagus. Huang et al [14,15]. isolated pathogens of  
 256 rice spikelet rot disease from infected rice samples collected from Zhejiang province  
 257 in Southern China and identified *Fusarium proliferatum* as one of the pathogens. Liu  
 258 [16] confirmed that *Fusarium proliferatum* was the main pathogen which induced rice  
 259 spikelet rot disease. Hou [17] demonstrated that *Fusarium proliferatum* was one of the  
 260 five determined *Fusarium* and accounted for 63.4% of the total detected strains.  
 261 Furthermore, they also determined that *Fusarium proliferatum* produced mycotoxins.  
 262 Du et al [18]. detected *Penicillium*, *Aspergillus*, and *Fusarium* as the major fungal  
 263 genus in Huaidao No. 5 rice freshly harvested in 2013 and indicated that *Penicillium*  
 264 and *Aspergillus* are the dominant fungi genus. A great difference exists between their  
 265 result and ours, likely because Huaidao No. 5 was planted in Jiangsu Province which  
 266 is located on the east coast of China and has a climate type of subtropical monsoon  
 267 climate to temperate monsoon climate, while Heilongjiang Province is located in  
 268 northeastern China with a temperate continental monsoon climate. Consequently, the  
 269 rice fungal communities in these two provinces are rather dissimilar.

270

271 Table 2 Fungal diversity and abundance (%) of rice samples at genus and species levels  
 272 collected from the four regions\*

Levels	Fungal Strains	Wuchang	Jiamusi	Zhaoyuan	Tailai
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Genus	<i>Cladosporium</i>	45.83±7.58 <sup>a</sup>	24.40±8.36 <sup>b</sup>	42.80±12.83 <sup>a</sup>	43.91±8.18 <sup>a</sup>
	<i>Epicoccum</i>	9.65±3.32 <sup>a</sup>	23.16±18.15 <sup>bc</sup>	8.72±2.97 <sup>a</sup>	18.26±2.69 <sup>ac</sup>
	<i>Alternaria</i>	11.96±4.61 <sup>a</sup>	2.62±0.66 <sup>b</sup>	8.93±4.89 <sup>ad</sup>	7.48±1.74 <sup>cd</sup>
	<i>Gibberella</i>	3.25±1.49 <sup>a</sup>	10.97±8.52 <sup>bc</sup>	5.49±4.22 <sup>ac</sup>	4.11±1.93 <sup>a</sup>
	<i>Fusarium</i>	1.16±0.73 <sup>a</sup>	4.83±6.06 <sup>a</sup>	13.90±20.35 <sup>a</sup>	2.05±1.72 <sup>a</sup>
	<i>Saitozyma</i>	0.27±0.09 <sup>a</sup>	0.39±0.26 <sup>a</sup>	0.18±0.07 <sup>a</sup>	2.83±3.01 <sup>b</sup>
	<i>Papiliotrema</i>	1.31±0.77 <sup>a</sup>	0.32±0.25 <sup>bc</sup>	0.79±0.64 <sup>ac</sup>	0.93±0.33 <sup>ac</sup>
	<i>Clonostachys</i>	0.26±0.17 <sup>a</sup>	0.23±0.21 <sup>a</sup>	2.01±5.15 <sup>a</sup>	0.30±0.40 <sup>a</sup>
	<i>Sporidiobolus</i>	0.17±0.03 <sup>a</sup>	0.12±0.08 <sup>a</sup>	0.75±1.19 <sup>ac</sup>	1.61±1.17 <sup>bc</sup>
	<i>Cryptococcus</i>	0.74±0.82 <sup>a</sup>	0.47±0.24 <sup>a</sup>	0.18±0.11 <sup>ab</sup>	1.23±1.08 <sup>ac</sup>
	Unclassified	30.91±18.50 <sup>a</sup>	28.46±7.62 <sup>a</sup>	13.94±6.60 <sup>a</sup>	14.26±1.95 <sup>a</sup>
Species	<i>Epicoccum_nigrum</i>	9.22±3.09 <sup>a</sup>	18.26±9.18 <sup>b</sup>	8.22±2.92 <sup>a</sup>	17.05±2.66 <sup>b</sup>
	<i>Fusarium_proliferatum</i>	1.10±0.74 <sup>a</sup>	4.47±5.74 <sup>a</sup>	13.83±20.36 <sup>a</sup>	2.01±1.72 <sup>a</sup>
	<i>Gibberella_zeae</i>	1.25±1.16 <sup>a</sup>	9.04±6.68 <sup>b</sup>	1.68±0.92 <sup>a</sup>	2.60±1.31 <sup>a</sup>
	<i>Gibberella_intricans</i>	1.02±0.75 <sup>a</sup>	1.16±1.58 <sup>a</sup>	2.54±3.38 <sup>a</sup>	0.55±0.51 <sup>a</sup>
	<i>Gibberella_fujikuroi</i>	0.94±0.83 <sup>a</sup>	0.62±0.45 <sup>a</sup>	1.22±1.07 <sup>a</sup>	0.94±0.62 <sup>a</sup>
	<i>Gliocladium_cibotii</i>	1.68±1.50 <sup>a</sup>	0.10±0.09 <sup>b</sup>	0.12±0.09 <sup>b</sup>	0.11±0.03 <sup>b</sup>
	Unclassified	79.89±3.94 <sup>a</sup>	63.15±7.83 <sup>bc</sup>	69.74±18.42 <sup>ac</sup>	73.52±5.39 <sup>ac</sup>

273 \* Values followed by the same superscript letter in the same row are not significantly different.

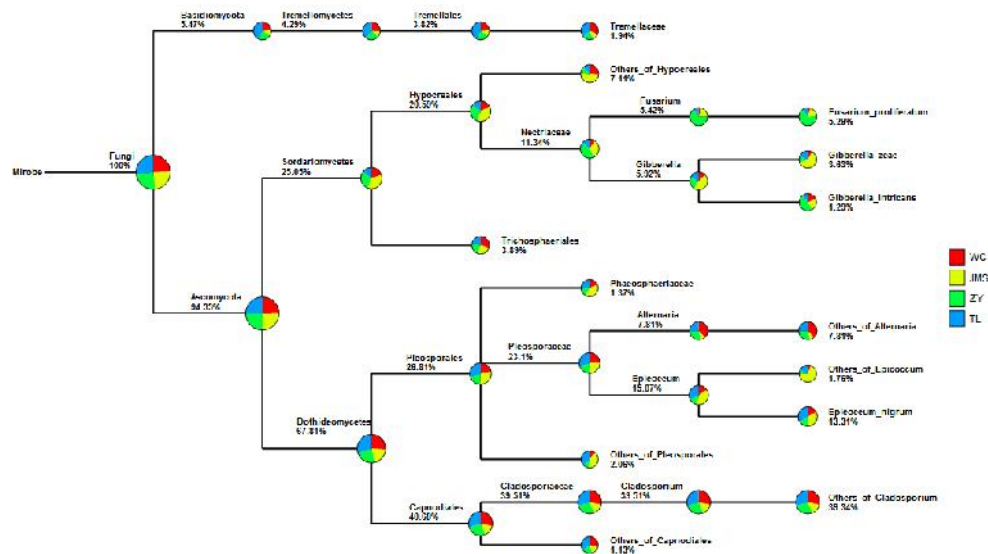
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275 As seen in Table 2, for the four regions the dominant fungi at Genus level are  
 276 *Cladosporium*, *Epicoccum*, *Alternaria*, *Gibberella*, and *Fusarium*, and almost no  
 277 significant difference in their abundance was observed among the five Genera in the  
 278 four regions. *Cladosporium* has been the most frequently found species in the four  
 279 regions. However, the abundance of *Cladosporium* in Jiamusi was significantly lower  
 280 than those of in the other three regions. *Cladosporium* is recognized as a psychrophile

281 hence it is more adaptable to cool temperature condition. The cause of its low  
 282 abundance in Jiamusi in comparison to the other three regions is still unclear.  
 283 *Cladosporium* has been proven to be a potentially pathogenic mycotoxin-producing  
 284 fungus frequently occurring in outdoor environments [19]. In addition, the proportion  
 285 of *Epicoccum* in Jiamusi was greater than those in the other three regions. *Epicoccum*  
 286 is a plant pathogen and widespread fungus which produces coloured pigments.  
 287 Therefore, rice in Jiamusi region has a higher probability of contamination by  
 288 coloured pigments which will in turn reduce rice quality.

289 The dominant fungi at the species level are *Epicoccum\_nigrum*,  
 290 *Fusarium\_proliferatum*, and *Gibberella\_zeae*. Like above, almost no significant  
 291 difference in their abundance was found among the three species in the four regions.  
 292 This is probably due to the relatively close geographical proximity of the four regions  
 293 resulting in similar climatic conditions.

294



295

296 Fig. 4 Multi rice samples taxonomy analysis tree on the species level

297

298 As seen in Fig. 4, fungal community of the four regions (composed of 12 rice  
 299 samples) was mainly composed of two Phylum, *Ascomycota* and *Basidiomycota*,  
 300 which account for 94.33% and 5.47%, respectively. At the species level, others of  
 301 *Cladosporium* accounts for 39.34% of the total species, followed by *Epicoccum*  
 302 *nigrum* which accounts for 13.31%, others of *Alternaria*, *Fusarium proliferatum*,

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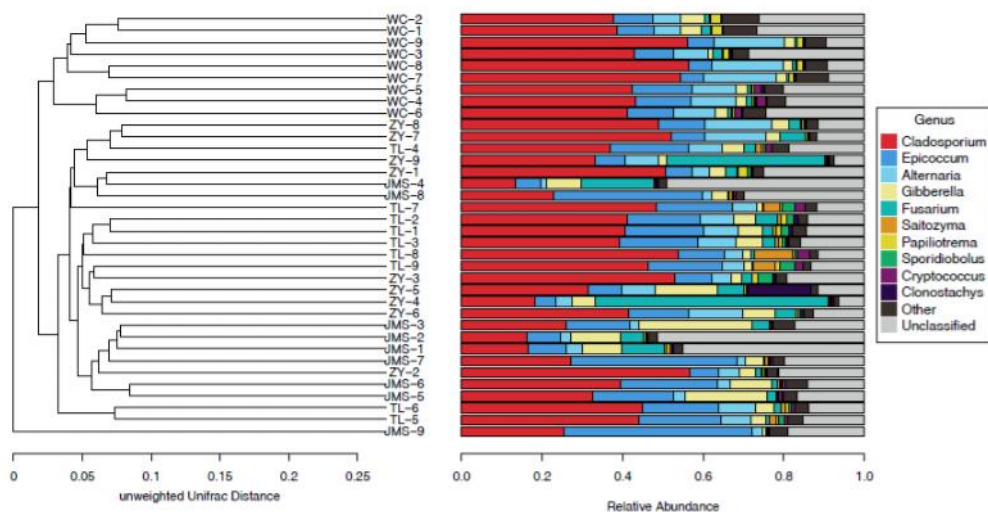
303 *Gibberella zeae*, and others of *Epicoccum* account for 7.81%, 5.29%, 3.63%, and  
304 1.76%, respectively. The least is *Gibberella intricans*, which accounts for 1.29%. For  
305 the mycotoxin-producing species, the proportion of *Fusarium proliferatum* accounts  
306 for absolute dominant in Zaoyuan region in comparison with the other three regions,  
307 and this probably is a hint that rice planted in Zhaoyuan has a greater potential to be  
308 contaminated by mycotoxins especially Fumonisin. The proportion of *Gibberella zeae*  
309 and others of *Epicoccum* in Jiamusi region account for absolute dominant compared  
310 with the other three regions. Since *Gibberella zeae* is the sexual stage of *Fusarium*  
311 *graminearum*, it has the possibility of producing deoxynivalenol (DON) and nivalenol  
312 (NIV) [20]. In addition, the proportion of *Gibberella intricans* in Zhaoyuan is the  
313 biggest in comparison with the other three regions. Although non-toxigenic fungi and  
314 yeasts themselves may only cause spoilage without safety issues, the damage they  
315 caused still not to be ignored.

316

### 317 **3.3 Cluster analysis of the 12 rice samples**

318 As seen in [Fig. 5](#), the fungi of the 3 rice samples (three replicates for each sample)  
319 from Wuchang city were clustered into one group. This is probably a result of near  
320 geographical proximity among the three spots where the rice samples were collected  
321 resulting in a similar fungal community. However, not all the fungi from the same  
322 region can be clustered into one group. Many regions have rice fields with varying  
323 soil types, water resources, types of fertilization, rice varieties, and other  
324 environmental factors which might increase the possibility of fungal diversity and  
325 make it difficult to cluster the fungi of rice samples from the same region into one  
326 group. Nevertheless, most of the fungi of rice samples from the same region can be  
327 clustered into the same group.

328



329

330 Fig. 5 Unweighted Pair-Group Method with Arithmetic Means (UPGMA) analysis of microbial  
 331 community structure based on ITS gene amplicon sequencing data.

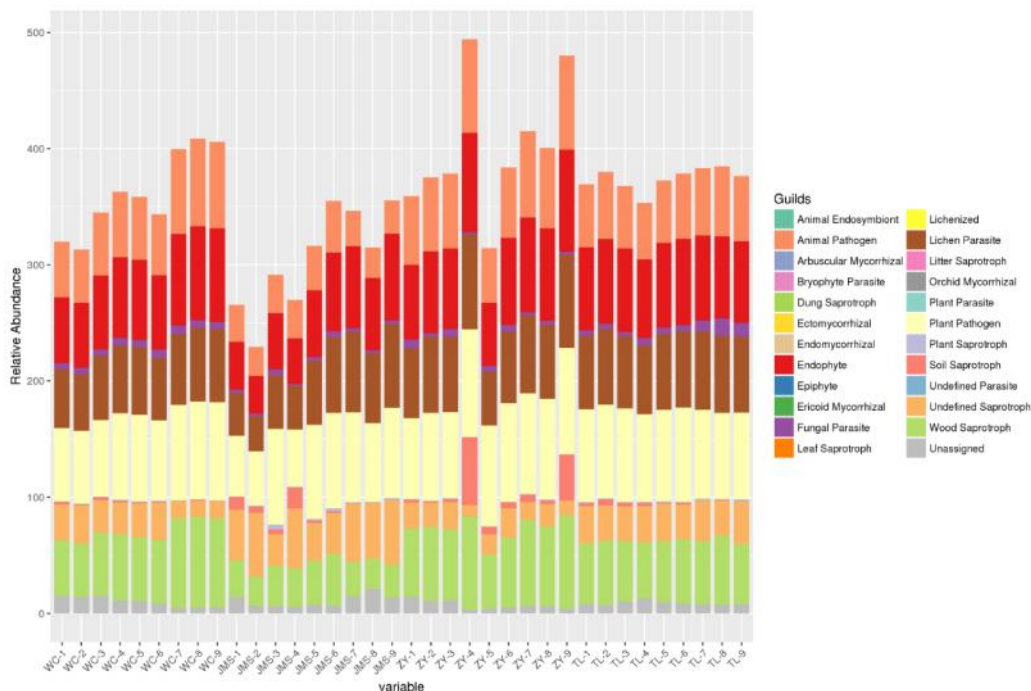
332

### 333 3.4 Fungal communities and functional guilds analysis

334 Fungal communities and functional guilds of the rice samples detected in the four  
 335 regions are shown in Fig. 6. As seen in this figure, an open environment enables the  
 336 rice to be a plant host to a wide range of environmental fungi. The most abundant  
 337 phylotypes are seen to be plant pathogen, endophyte, fungal parasite, undefined  
 338 saprotroph, wood saprotroph, soil saprotroph, as well as animal pathogen. For all the  
 339 rice samples, plant pathogen, lichen parasite, soil saprotroph, wood saprotroph, and  
 340 endophyte account for the largest proportions. For the mycotoxigenic fungi species,  
 341 they are in the category of plant pathogen.

342





343

344 Fig. 6 Stacks of Guilds of the 12 rice samples from Heilongjiang province

345

346 Around 70% of all major crop diseases were induced by fungal plant pathogens.  
 347 Furthermore, 15% of global agricultural production was destroyed through yield  
 348 losses and mycotoxin contamination [21]. Plant pathogens, especially mycotoxigenic  
 349 fungi are considered to be the most harmful class of plant pathogens by far. As a  
 350 cosmopolitan genus of filamentous ascomycete fungi, *Fusarium* includes a number of  
 351 toxin-producing plant pathogens of agricultural importance [22]. For the rice freshly  
 352 harvested in Heilongjiang province, the *Fusarium proliferatum* determined likely  
 353 includes mycotoxigenic species, although a **fungi toxicity** test has not been conducted  
 354 yet.

355

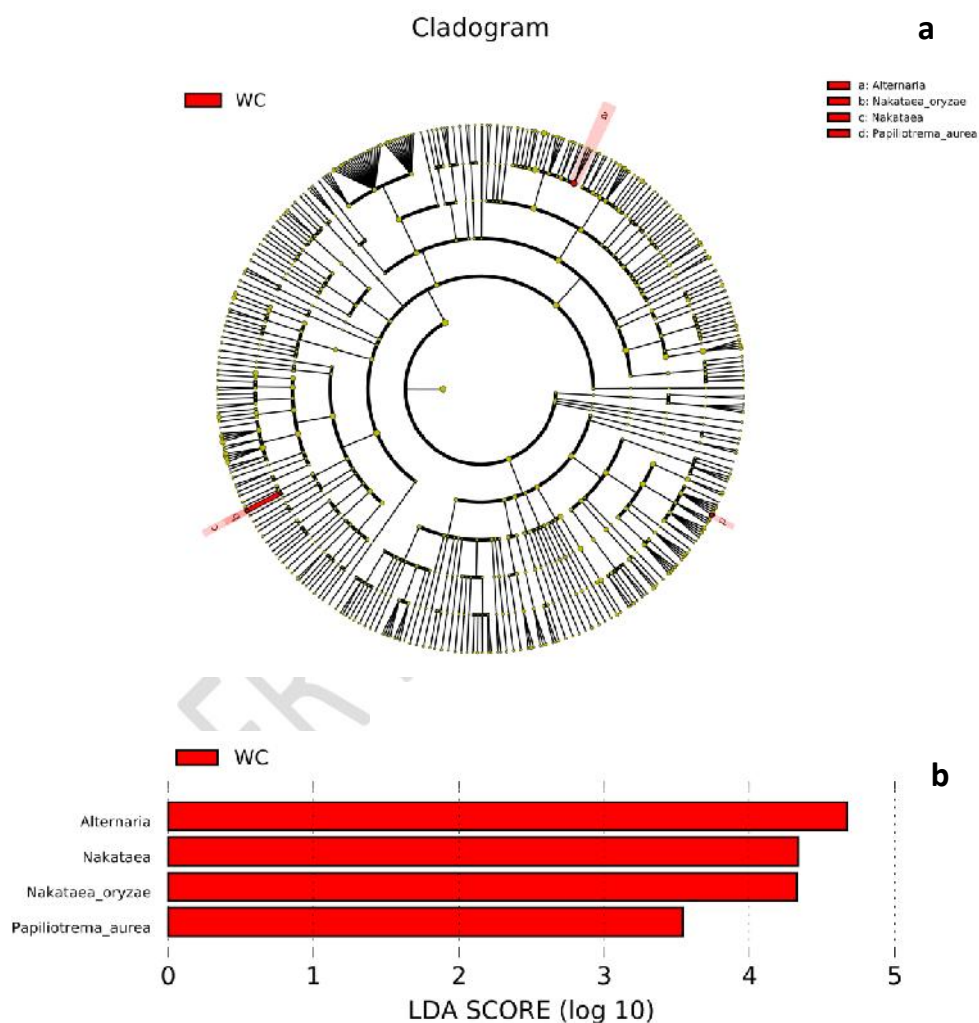
### 356 3.5 LEfSe analysis

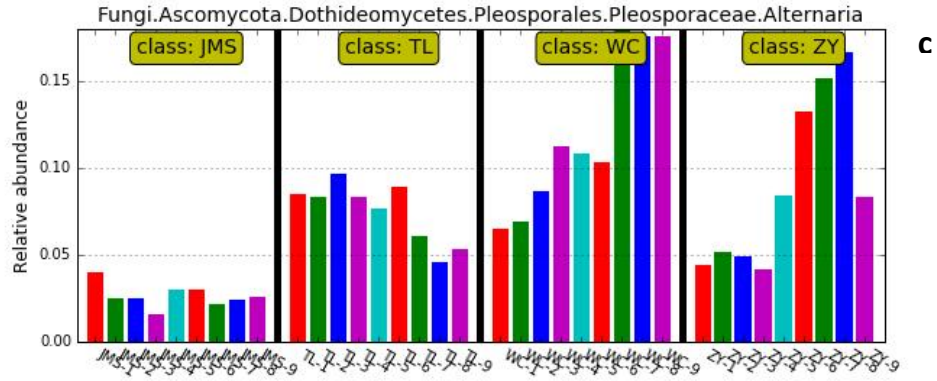
357 Key phylotypes of rice fungi microbiota representing the four regions identified  
 358 using linear discriminant analysis (LDA) effect size (LEfSe) are shown in Fig. 7. As  
 359 seen in Fig 7a, the Cladogram indicates that the numbers of four fungi genera and  
 360 species in Wuchang city are significantly greater than those of in the other three  
 361 regions; they are *Alternaria*, *Nakataea*, *Nakataea\_oryzae*, and *Papiliotrema\_aurea*.



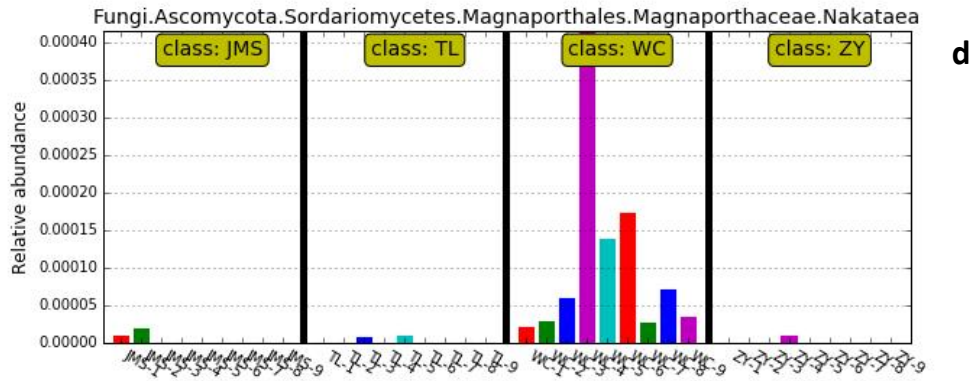
362 Their LDA scores are greater than 3 (Fig. 7b) and they might be considered as  
363 specific fungi associated with Wuchang region. Fig. 7c, d, e, and f illustrate the  
364 relative abundance of the four fungi given above in the four regions. Consequently, it  
365 might be possible to develop biomarkers using the four fungi given above to  
366 distinguish rice from Wuchang region.

367

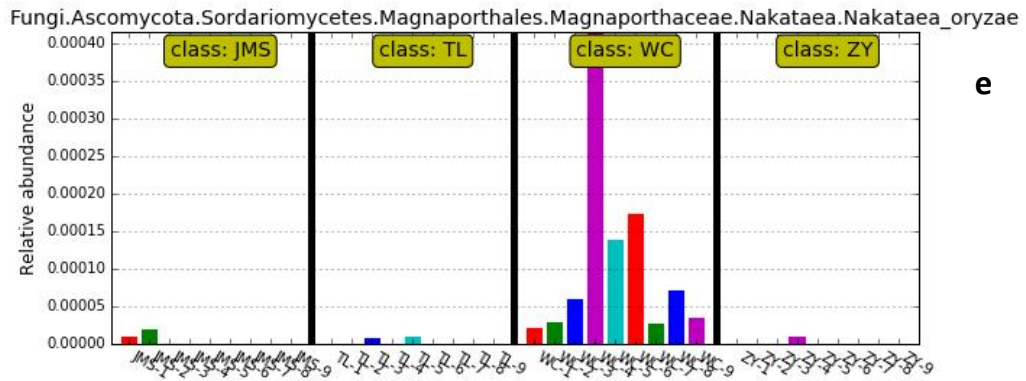




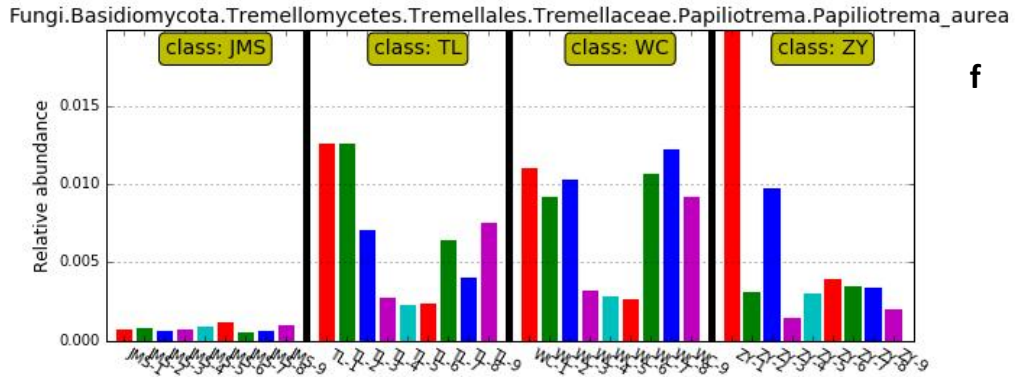
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373

374 Fig. 7 Cladogram, LDA score, and relative abundance of fungi of rice samples from the four  
 375 regions. a. Cladogram; b. LDA score; c. relative abundance of *Alternaria* of rice samples from  
 376 the four regions; d. relative abundance of *Alternaria* of rice samples from the four regions; e.  
 377 a relative abundance of *Nakataea\_oryzae* of rice samples from the four regions; f. a relative  
 378 abundance of *Papiliotrema\_aurea* of rice samples from the four regions

379

#### 380 4. CONCLUSION

381 To explore the potential of fungi contamination as well as mycotoxin production,  
 382 it is necessary to investigate field fungal diversity in rice in Heilongjiang province  
 383 through high throughput sequencing of freshly harvested rice samples. Our results  
 384 indicate that *Cladosporium* accounts for an absolute dominant at the genus level and  
 385 *Epicoccum\_nigrum*, *Fusarium\_proliferatum*, and *Gibberella\_zeae* are relatively  
 386 abundant fungi species, in which *Fusarium\_proliferatum* has the potential to produce  
 387 mycotoxins such as fumonisin. Rice planted in Zhaoyuan has the greatest potential to  
 388 produce fumonisin whereas rice grown in Jiamusi is most likely to be contaminated by  
 389 DON and NIV in comparison with the other three regions. Consequently, it is  
 390 necessary to take adequate measures to prevent mycotoxin production during rice  
 391 storage, as well as related damage induced by non-mycotoxins-producing fungus  
 392 growth and reproduction. In addition, *Alternaria*, *Nakataea*, *Nakataea\_oryzae*, and  
 393 *Papiliotrema\_aurea* are the specific fungi genera and species which can distinguish  
 394 rice planted in Wuchang from the other three regions.

395

396

397 COMPETING INTERESTS

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398 Authors have declared that no competing interests exist.

399

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UNDER PEER REVIEW