





Lentinula edodes (Berk.) Pegler, also called Xianggu or shiitake, is one of the most popular edible mushrooms worldwide. L. edodes was firstly cultivated at least 900–1,000 years ago in China, and is now the second most widely produced mushroom in the world after Agaricus bisporus (Miles and Chang, 2004). L. edodes is rich in minerals, vitamins, essential amino acids, and lentinan (Chang, 1980). It also has immunomodulatory (Xu et al., 2015), anticancer (Nagashima et al., 2013), and antiviral functions (Di Piero et al., 2010). Moreover, L. edodes could prevent environmental impacts caused by the accumulation of forest and agricultural wastes since it secretes hydrolytic and oxidative enzymes that are responsible for the degradation of organic substrates (Silva et al., 2005).

Breeding elite cultivars is important for sustainable development of the modern mushroom industry. The fruiting body of *L. edodes* is the main target of breeding schemes. Strains with a fast mycelium growth rate (MGR), high precocity, fine morphological characteristics of fruiting body, and high yield are selected as cultivars. The majority of important agronomic traits of *L. edodes* are quantitative traits controlled by multiple genes or quantitative trait loci (QTLs), which are highly influenced by the environment and show a continuous variation (Santoyo et al., 2008). Dissecting the genetic basis of important agronomic traits

TABLE 1 | Statistical characteristics of the 11 agronomic traits of *Lentinula edodes*.

Trait (unit)	Description ^a	Year	Minimum	Maximum	Mean	Range	Sd ^b	CV ^c (%)	r ^d	H ² (%)
DGR-myg (mm/d)	Mycelium growth rate in MYG medium	2013	1.47	6.32	5.51	4.85	0.77	13.99	/	98.82
DGR-sd (mm/d)	Mycelium growth rate in sawdust medium	2013	3.53	4.40	3.96	0.87	0.20	5.08	/	87.84
PD (mm)	Average diameter of pileus, determined	2013	29.54	65.48	45.67	35.94	8.92	19.54	0.398**	85.50
	as the mean of two perpendicular diameters	2014	30.81	67.82	53.03	37.02	7.94	14.96		
PT (mm)	Average thickness of pileus	2013	5.92	15.39	9.81	9.47	2.19	22.30	0.491**	86.91
		2014	5.05	13.90	10.56	8.85	2.02	19.10		
PW (g)	Average weight of pileus	2013	2.80	30.36	11.77	27.56	6.82	57.96	0.420**	84.64
		2014	3.88	28.90	14.58	25.01	5.14	35.23		
SD (mm)	Average diameter of stipe	2013	5.95	17.46	10.03	11.52	2.55	25.43	0.513**	86.96
		2014	7.07	15.66	10.70	8.60	1.82	16.99		
SL (mm)	Average length of stipe	2013	23.09	63.67	38.45	40.58	8.12	21.11	0.544**	89.89
		2014	22.09	62.33	41.28	40.25	7.88	19.09		
SW (g)	Average weight of stipe	2013	0.75	15.34	3.94	14.59	2.89	73.34	0.487**	86.05
		2014	0.93	9.37	4.25	8.45	1.73	40.64		
NF (per/bag)	Average number of total fruiting bodies	2013	2.29	88.25	22.13	85.96	17.39	78.57	0.555**	91.78
	per bag during the whole harvest time	2014	1.90	62.14	13.86	60.24	12.33	88.95		
WF (g/per)	Ratio of yield to the number of fruiting	2013	2.08	39.07	13.89	36.99	8.35	60.08	0.317*	84.15
	bodies harvested during the whole harvest time	2014	4.10	35.95	17.63	31.84	6.58	37.30		
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Y (g/bag) Total weight of fruiting bodies per bag during the whole **limm**vest

markers. A total of 379 polymorphic markers, comprising 328 InDels and 51 SSRs, were then selected. Only 297 markers (249 InDels and 48 SSRs) with a minor allele frequency (MAF) of ≥ 0.05 and missing data $\leq 5\%$ in the 89 strains were utilized for further analysis. These 297 markers were distributed in

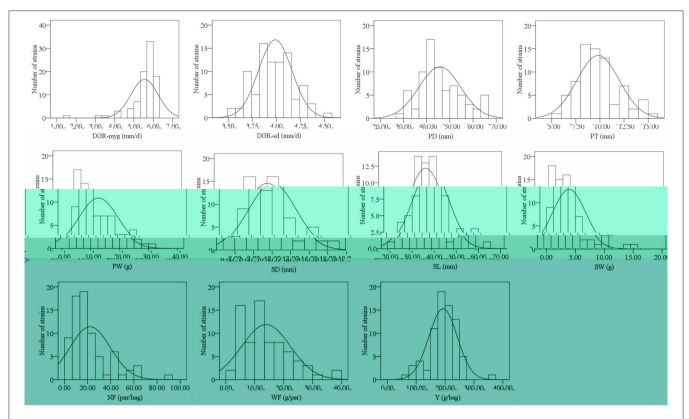


FIGURE 1 | Histograms showing the frequency distribution of 11 agronomic traits in 2013. The y-axis denotes the number of strains, whereas the x-axis indicates value range of traits.

TABLE 2 | A matrix of Pearson correlation coefficients (r) for the 11 agronomic traits in Chinese Lentinula edodes cultivars.

	PD	РТ	PW	SD	SL	sw	NF	WF	Υ
DGR-myg	0.125	0.211	0.223	0.223	0.142	0.197	-0.11	0.209	-0.208
DGR-sd	-0.054	0.004	-0.081	-0.014	-0.108	-0.133	0.07	-0.034	0.091
PD		0.853**	0.927**	0.852**	0.690**	0.756**	-0.733**	0.842**	-0.389**
PT	0.827**		0.813**	0.677**	0.525**	0.596**	-0.615**	0.748**	-0.403**
PW	0.884**	0.755**		0.883**	0.688**	0.828**	-0.711**	0.921**	-0.440**
SD	0.737**	0.753**	0.808**		0.780**	0.915**	-0.683^{**}	0.864**	-0.275*
SL	0.761**	0.669**	0.654**	0.508**		0.851**	-0.629^{**}	0.694**	-0.133
SW	0.821**	0.724**	0.864**	0.827**	0.840**		-0.601**	0.821**	-0.282*
NF	-0.850**	-0.835**	-0.776**	-0.737**	-0.734**	-0.750**		-0.767**	0.472**
WF	0.853**	0.746**	0.880**	0.710**	0.713**	0.834**	-0.780**		-0.475**
Y	-0.618**	-0.634**	-0.579**	-0.551**	-0.500**	-0.561**	0.731**	-0.561**	

The abbreviations are the same as those in Table 1. *P < 0.05; **P < 0.01.

Pearson correlation coefficients are presented here in pairs for agronomic traits measured in 2013 (upper right triangle) and in 2014 (lower left triangle).

Genetic Diversity

A total of 873 alleles were detected from the 297 loci in all 89 strains, and the number of alleles in each locus varied from two to seven with a mean of 2.939 (Supplementary Table S2). The PIC value varied from 0.022 (S676_SSR2) to 0.731 (S95_ID5) with an average of 0.381. In all 89 strains, the average number of alleles from the 48 SSR markers was 3.229, higher than that from the 249 InDel markers (2.884). However, the PIC value of SSRs (0.379)

was comparable to that of InDels (0.381), suggesting a similar genetic variation of both types of markers in the Chinese shiitake cultivars. From all 89 strains, the mean values of Ne, He, I, PPL, H, and PIC were 1.955, 0.456, 0.734, 100%, 0.454, and 0.381, respectively, indicating relatively low genetic variations among Chinese L. edodes cultivars. As for the two groups defined by the NJ tree, all the genetic parameters showed that the genetic diversity in Group A was lower than that in Group B (Table 3).

TABLE 3 | Genetic variability for Lentinula edodes cultivars in China.

Population	Na	Ne	1	Но	He	н	PIC	PPL%
Total	2.939	1.955	0.734	0.545	0.456	0.454	0.381	100.00
Group A	2.162	1.622	0.500	0.444	0.327	0.319	0.262	85.19
Group B	2.582	1.847	0.672	0.580	0.428	0.424	0.352	99.66

Observed number of alleles (Na), effective number of alleles (Ne), percentage of polymorphic loci (PPL), observed heterozygosity (Ho), expected heterozygosity (He), Shannon's information index (I), gene diversity (H), and polymorphism information content (PIC).

The lower genetic diversity in Group A could be due to the smaller sample size.

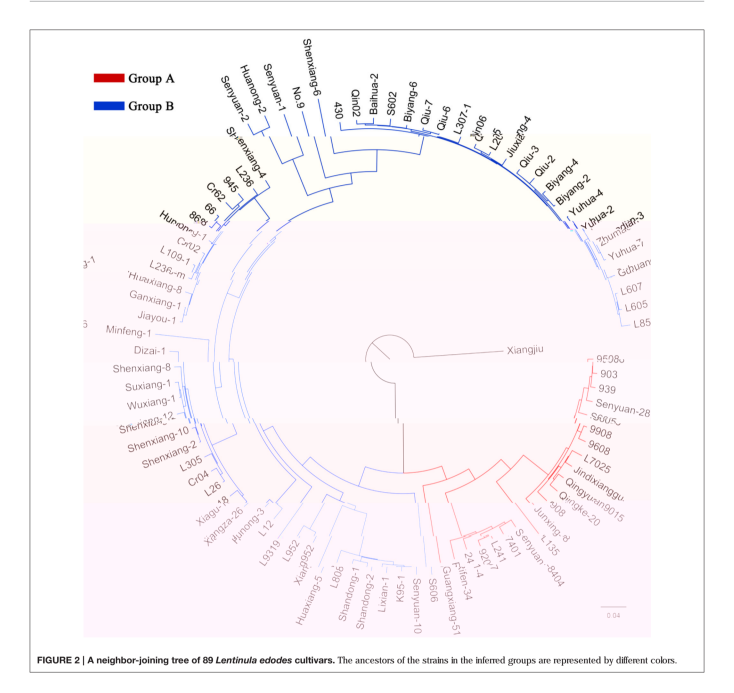
Population Structure and Linkage Disequilibrium

According to the genotyping data, the 89 strains represent 89 unique genotypes, and therefore are not clones. In the NJ tree of *L. edodes*, all strains except Xiangjiu clustered into two distinct groups (Figure 2). Group A consisted of 21 strains and Group B contained 67 strains. PCA also identified two groups congruent with those in the NJ tree (Supplementary Figure S3). The percentages of variation explained by the first 3 axes were 32.7%, 14.1%, and 9.8%.

Analysis of molecular variance (AMOVA) results suggested that the majority of genetic variation was included within populations (75.33%) (Table 4). The overall F_{ST} value across all the strains except Xiangjiu was 0.247, suggesting a great di erentiation among L. edodes cultivars in China.

Model-based STRUCTURE was also utilized to investigate the population structure of the 89 strains. In the analysis of ΔK , a clear maximum was detected for K=2 ($\Delta K=3015$) (Figure 3A). Therefore, two groups were identified in the collection of 89 L. edodes cultivars in China (Figure 3B), which agreed with the results of NJ tree and PCA. None of the 89 strains was assigned exclusively to one group or the other, and all strains shared mixed ancestries from the two groups. These demonstrate that the Chinese shiitake cultivars were genetically closely related.

A total of 19,122 (43.50%) InDel and SSR marker pairs displayed significant LD among all 89 strains ($P \leq 0.001$). The r^2 values among these marker pairs varied from 0.128 to 1, with an average of 0.316 (Supplementary Table S4). At the highly significant threshold of $r^2 \geq 0.2$, 30.43% (13,378) of the marker pairs remained in LD. In this study, owing to the fact that only 73 of these 297 makers were used to construct linkage map in our previous study (Gong et al., 2016), the genomewide LD decay along with the increase of genetic distance were not detected (Supplementary Table S5). The averaged r^2



This is because the two traits were determined in incubators and therefore not a ected by the changing environmental conditions under which the nine traits were measured. Indeed, ANOVA revealed significant di erences of the fruiting body-related traits between the 2 years, suggesting a strong e ect of environmental factors (i.e., year) on these traits.

Here, we observed extensive significant correlations between the nine fruiting body-related traits of L. edodes cultivars, in agreement with previous results detected in two segregating populations and one natural population (Gong et al., 2014b). Yield and yield-component traits of L. edodes were found to exhibit the triangular relationship as displayed in our recent

report (Gong et al., 2014b) and in *A. bisporus* (Foulongne-Oriol et al., 2012a), i.e., yield was positively correlated with NF but negatively correlated with WF; and NF was negatively correlated with WF. For nine fruiting body-related traits and two mycelium growth-related traits, no obvious correlation was observed in 2013, which indicated that the growth of mycelium and development of fruiting bodies may be independently controlled.

A total of 21 molecular markers were identified to be associated with two to four traits by association mapping. For instance, S106_inID1 was associated with four traits (WF, PD, PW, and SW), and S560_ID1 associated with three traits (WF, PW, and PD), partly illustrating the genetic basis of phenotypic

TABLE 4 | Analysis of molecular variance (AMOVA) among and within populations of *Lentinula edodes* cultivars in China.

Source	df	SS	MS	Est. Var.	%var
Among populations	1	1337.155	1337.155	19.955	24.67
Within populations	174	10600.862	60.924	60.924	75.33
Total	175	11938.017		80.880	100.00

df, degree of freedom; SS, sum of squared observations; MS, mean of squared observations; Est. var., estimated variance; %Var., percentage of total variance.

correlation between these traits. The two major reasons for trait correlations are pleiotropy and close linkage between QTLs controlling di erent traits (Mackay et al., 2009; Chen and Lübberstedt, 2010). Our recent work on QTL mapping in two segregating populations also suggested that the co-localization of QTLs underlining di erent traits may be the genetic basis for phenotypic correlation of fruiting body-related traits in *L. edodes* (Gong et al., 2016). Combining evidences from both association mapping and our recent results from QTL mapping, we postulate that the genetic basis of phenotypic correlation in *L. edodes* is the tight linkage of QTLs and pleiotropy.

Multigenic e ects were also observed in this study (Table 5). For instance, nine markers, including S278_ID10, S278_ID36, S278_ID41, and S328_ID5, were associated with PD. Multigenic e ects suggested that these traits in *L. edodes* were complex quantitative traits that were a ected by polygenes.

Genetic Diversity

Understanding the genetic diversity and genetic basis underlying important agronomic traits could improve breeding schemes of L. edodes. In general, the processes of domestication and breeding have a strong impact on the genetic diversity of cultivated species (Font i Forcada et al., 2015). Here, the values of Shannon's information index (I) and polymorphism information content (PIC) revealed by InDel and SSR markers were 0.734 and 0.381, respectively. In a wild population containing 88 Chinese L. edodes strains, the I and PIC values were 0.836 and 0.395, respectively (Xiang et al., 2016). In another study, the PIC value was 0.53 in 89 L. edodes strains from East Asia (Kim et al., 2009). Genetic variation of Chinese L. edodes cultivars is low and was postulated to be derived from a limited number of elite strains (Chiu et al., 1996). Therefore, the wild strains should be introduced into the breeding schemes to diversify the genetic basis of shiitake cultivars in China.

Population Structure and Linkage Disequilibrium

Detailed knowledge on population structure is important to control spurious associations between phenotypes and genotypes in association mapping (Pritchard et al., 2000). Model-based analysis of population structure could provide necessary information in association mapping. Here, population structure analyses based on three methods demonstrated that the Chinese *L. edodes* cultivars could be divided into two unique groups, with Xiangjiu being the sole exception. This strain was proven to be distinct from other *L. edodes* cultivars in previous clustering

analyses (Fu et al., 2010; Liu et al., 2015). Using strains di erent from the current study, the L. edodes cultivars in China was also separated into two main groups (Zhang et al., 2007; Fu et al., 2010; Liu et al., 2012). Therefore, it is reasonable to speculate that L. edodes cultivars in China contained two di erent gene pools, which possibly resulted from domestication and breeding. Great genetic di erentiation existed between the two groups as indicated by a F_{ST} value of 0.247, which is comparable to that in the Chinese wild L. edodes population ($F_{ST} = 0.252$) (Xiang et al., 2016).

A high level of LD among marker pairs was observed in this study. As mentioned before, the narrow genetic base of Chinese shiitake cultivars revealed here and in previous studies (Chiu et al., 1996; Fu et al., 2010) might be one of the factors that could explain the high level of pairwise LD. Also, a small number of tested strains and molecular markers may cause bias of LD estimates. Furthermore, the population structure contributes to increasing LD level. Population structure could create unexpected LD between unlinked loci across the genome (Yan et al., 2011). The mixing of individuals belonging to di erent subpopulations with di erent allele frequencies creates LD, when these subpopulations are admixed to construct a panel of lines for association mapping. Significant LD between unlinked loci results in false-positive associations between a marker and a trait (Soto-Cerda and Cloutier, 2012). In this study, small structured population with narrow genetic base may be the major factors that causes the high level LD and then led to spurious associations between marker alleles and the phenotypes. Therefore, the further association studies require the careful choice of germplasm, as well as a larger number of markers and strains.

Association Mapping

Association mapping has been successfully utilized in crop species, such as maize, cotton, wheat, and rice (Abdurakhmonov and Abdukarimov, 2008). Due to their edible and medicinal values, mushrooms have been consumed by humans for a long time. However, linkage and association mapping in mushroom species are still in their infancy, and information is limited to identify QTLs controlling agronomic traits in mushroom species. Here, we detected 78 marker-trait associations covering 43 molecular markers and four traits. Marker-trait associations detected by this method could provide valuable information for MAS in breeding schemes of *L. edodes*.

Among the 297 markers used here, 73 markers were the same as those used by Gong et al. (2016), while 47 markers were employed by Xiang (2015). Five markers were found to be associated with the same traits as in previous reports (Xiang, 2015; Gong et al., 2016) (Table 5). Gong et al. (2016) revealed that marker S48_ID1 was located in a QTLs-hotspot region in LG2 that was related to PD, PT, PW, SL, SD, SW, and WF. In this study, S48_ID1 was also identified to be associated with WF. S560_ID1 was significantly associated with WF, PW and PD, consistent with previous findings (Gong et al., 2016). S346_ID1 was found to be associated with SW and lie in a QTL hotspot region in LG4 related to SW, SL, and SD (Gong et al., 2016). Three of the six hotspot regions previously identified by Gong

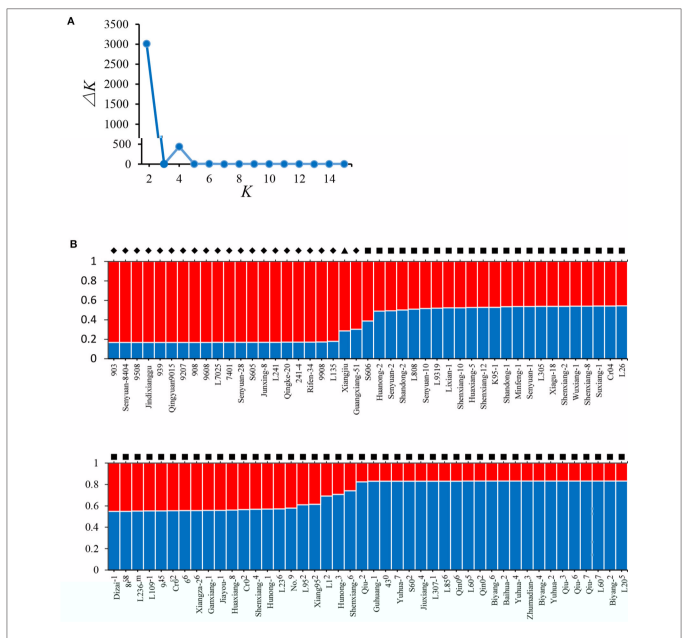


FIGURE 3 | Results of STRUCTURE analysis. (A) Estimation of the number of populations for K ranging from 2 to 15 by ΔK values; (B) Classification of 89 L. eclodes cultivars into two genetic groups. The distribution of the strains assigned to different groups is indicated by the color code (Group A: red, Group B: blue). The y-axis quantifies the cluster membership, and the x-axis lists the different strains. Strains from the different groups defined in the NJ tree are marked in different symbols: \spadesuit , Group B; \blacktriangle , Xiangjiu, excluded from the two groups in the NJ tree.

et al. (2016) were confirmed by this association mapping, thus suggesting that there are reliable regions harboring QTLs related to fruiting body in *L. edodes*. Moreover, S255_ID1 was found to be associated with PW and WF, and S163_E1 associated with PW, in agreement with a previous report (Xiang, 2015).

Five markers were identified to be associated with the same trait in both years. They are S127_ID1 and S328_ID5 for PW, S278_ID10 and S704_inID1 for PW and PD, and S278_ID41 for PD. It is worth mentioning that S278_ID41 is located in a <1 kb position to S278-R/F that lay in a QTL hotspot in MG4 special for SL, PD, and PW (Gong et al., 2016).

Hence, S278_ID41 is also located in this QTL hotspot in MG4.

The foregoing 10 markers were verified by previous studies or detected in both years, suggesting that association

TABLE 5 | Associations between agronomic traits and markers in the Chinese Lentinula edodes cultivars.

Year	Trait	Marker	Scaffolds*	P [#]	<i>P</i> _FDR	R ² (%)
2013	PD	S278_ID41 ^a	Le_N7_S278	8.58×10^{-5}	0.021	22.90
2013	PD	S328_ID5	Le_N7_S328	1.39×10^{-4}	0.021	21.87
2013	PD	$S278_ID10^a$	Le_N7_S278	2.95×10^{-4}	0.024	20.51
2013	PD	S278_ID36	Le_N7_S278	4.4×10^{-4}	0.024	19.32
2013	PD	S106_inID1	Le_N7_S106	$4.7 imes 10^{-4}$	0.024	15.53
2013	PD	S473_ID1	Le_N7_S473	5.07×10^{-4}	0.024	19.25
2013	PD	$S704_{in}ID1^{a}$	Le_N7_S704	5.54×10^{-4}	0.024	19.30
2013	PW	S278_ID41	Le_N7_S278	8.69×10^{-5}	0.026	22.96
2013	PD	S127_ID1	Le_N7_S121	9.59×10^{-4}	0.036	27.57
2013	PW	$S328_ID5^a$	Le_N7_S328	2.4×10^{-4}	0.036	20.79
2013	PW	S473_ID1	Le_N7_S473	$3.6 imes 10^{-4}$	0.036	20.09
2013	PW	S278_ID10 ^a	Le_N7_S278	8.38×10^{-4}	0.044	18.14
2013	PW	S127_ID1 ^a	Le_N7_S121	8.76×10^{-4}	0.044	28.02
2013	PW	S108_ID1	Le_N7_S108	9.52×10^{-4}	0.044	20.70
2013	PW	S443_inID1	Le_N7_S443	0.001	0.044	13.69
2013	PW	S704_inID1 a	Le_N7_S704	0.001	0.044	17.37
2013	PW	S613_inID1	Le_N7_S613	0.001	0.044	22.32
2014	WF	S278_ID10	Le_N7_S278	7.30×10^{-6}	0.001	23.75
2014	WF	S427 ID1	Le_N7_S427	1.16×10^{-5}	0.001	31.32
2014	WF	S560 ID1 ^b	Le_N7_S560	1.26×10^{-5}	0.001	25.96
2014	WF	S443_inID1	Le_N7_S443	1.45×10^{-5}	0.001	22.05
2014	WF	S286_SSR3	Le_N7_S286	1.53×10^{-5}	0.001	21.85
2014	WF	S278_ID36	Le_N7_S278	1.82×10^{-5}	0.001	25.10
2014	WF	S458_ID5	Le_N7_S458	2.38×10^{-5}	0.001	29.95
2014	WF	S488_ID1	Le_N7_S488	2.66×10^{-5}	0.001	27.22
2014	WF	S551_inID1	Le_N7_S551	2.96×10^{-5}	0.001	24.45
2014	WF	S603_ID1	Le_N7_S603	3.30×10^{-5}	0.001	24.40
2014	WF	S267_ID1	Le_N7_S267	3.30×10^{-5}	0.001	24.40
2014	WF	S106_inID1	Le_N7_S106	4.95×10^{-5}	0.001	20.22
2014	WF	S636_inID1	Le_N7_S636	8.10×10^{-5}	0.002	23.38
2014	WF	S255_ID1 ^c	Le_N7_S255	1.04×10^{-4}	0.002	24.47
2014	WF	S48_ID1 ^b	Le_N7_S48	1.29×10^{-4}	0.002	26.54
2014	WF	S131_ID1	Le_N7_S131	1.48×10^{-4}	0.003	26.25
2014	PW	S560_ID1 ^b	Le_N7_S560	1.67×10^{-5}	0.005	27.95
2014	PW	S704_inID1 ^a	Le_N7_S704	4.83×10^{-5}	0.005	25.88
2014	PW	S211_ID1	Le_N7_S211	6.05×10^{-5}	0.005	28.03
2014	PW	S488_ID1		8.14×10^{-5}	0.005	27.37
2014	PW	S767_ID1	Le_N7_S488 Le_N7_S767	8.51×10^{-5}	0.005	20.50
		S255_ID1 ^c		1.33×10^{-4}		
2014	PW		Le_N7_S255	4.03×10^{-4}	0.007	26.29
2014	WF	S127_ID1	Le_N7_S121	4.03×10^{-4} 5.11×10^{-4}	0.007	26.36
2014	WF	S95_ID5	Le_N7_S95	2.06×10^{-4}	0.008	28.33
2014	PW	S328_ID5a	Le_N7_S328		0.009	22.28
2014	WF	S641_SSR1	Le_N7_S641	6.94×10^{-4}	0.011	27.25
2014	PW	S258_ID5_2	Le_N7_S258	3.10×10^{-4}	0.011	27.03
2014	PW	S126_ID1	Le_N7_S126	3.29×10^{-4}	0.011	21.08
2014	PW	S163_E1 ^c	Le_N7_S163	5.19×10^{-4}	0.015	28.67
2014	PW	S534_ID1	Le_N7_S534	5.68×10^{-4}	0.015	25.57
2014	WF	S206_ID1	Le_N7_S206	0.001	0.017	24.25
2014	PW	S278_ID36	Le_N7_S278	8.19×10^{-4}	0.020	19.12
2014	PW	S457_ID1	Le_N7_S457	9.42×10^{-4}	0.021	29.09

(Continued)

TABLE 5 | Continued

Year	Trait	Marker	Scaffolds*	P [#]	P_FDR	R ² (%)
2014	PW	S95_ID5	Le_N7_S95	9.80×10^{-4}	0.021	29.38
2014	SW	S127_ID1	Le_N7_S121	3.98×10^{-4}	0.024	29.51
2014	SW	S258_ID5_2	Le_N7_S258	4.00×10^{-4}	0.024	26.97
2014	SW	S90_E1	Le_N7_S90	4.03×10^{-4}	0.024	17.11
2014	SW	S278_ID33	Le_N7_S278	4.03×10^{-4}	0.024	17.11
2014	SW	S553_ID5	Le_N7_S553	4.03×10^{-4}	0.024	17.11
2014	SW	S108_ID1	Le_N7_S108	4.97×10^{-4}	0.024	23.66
2014	PD	S278_ID41 ^a	Le_N7_S278	1.81×10^{-4}	0.025	20.71
2014	PD	S560_ID1 ^b	Le_N7_S560	1.96×10^{-4}	0.025	20.64
2014	PD	S278_ID10 ^a	Le_N7_S278	3.31×10^{-4}	0.025	16.92
2014	PD	S704_inID1 ^a	Le_N7_S704	3.43×10^{-4}	0.025	20.10
2014	PW	S278_ID3	Le_N7_S278	0.001	0.027	23.89
2014	WF	S178_ID1	Le_N7_S178	0.002	0.030	13.11
2014	WF	S704_inID1	Le_N7_S704	0.002	0.030	16.55
2014	SW	S32_E1	Le_N7_S32	7.16×10^{-4}	0.030	28.04
2014	SW	S636_inID1	Le_N7_S636	8.54×10^{-4}	0.031	19.22
2014	SW	S178_ID1	Le_N7_S178	0.001	0.033	15.25
2014	SW	S346_ID1 ^b	Le_N7_S346	0.001	0.033	14.73
2014	SW	S470_ID1	Le_N7_S470	0.001	0.037	14.22
2014	PW	S127_ID1 ^a	Le_N7_S121	0.002	0.039	24.68
2014	WF	S470_ID1	Le_N7_S470	0.003	0.039	12.07
2014	PW	S90_E1	Le_N7_S90	0.003	0.039	12.71
2014	PW	S278_ID33	Le_N7_S278	0.003	0.039	12.71
2014	PW	S553_ID5	Le_N7_S553	0.003	0.039	12.71
2014	PW	S278_ID10 ^a	Le_N7_S278	0.003	0.040	12.78
2014	SW	S160_ID1	Le_N7_S160	0.002	0.044	13.48
2014	SW	S278_ID10	Le_N7_S278	0.002	0.044	13.75
2014	PW	S35_inID1	Le_N7_S35	0.003	0.045	24.87

^aMarker detected in both years;

bi-parental population, association mapping investigates genetic variations in a natural population, and thus can evaluate many alleles simultaneously (Abdurakhmonov and Abdukarimov, 2008). Therefore, this study confirmed the feasibility and reliability of association mapping in *L. edodes*.

The 10 aforementioned markers resided on or were close to 24 annotated genes of the $L.\ edodes$ reference genome (Supplementary Table S6). These genes could be potential candidate genes related to agronomic traits. By checking the RNA-seq expression levels of the 24 candidate genes in shiitake strain L54 (unpublished data), 13 of them (54.17%) were found to be significantly up-regulated or down-regulated during the transition from mycelium to primordium, with fold changes >2 (Supplementary Figure S5). Hence, it is reasonable to speculate that most of these candidate genes are involved in the development of fruiting body.

Despite the importance of edible mushrooms, research on their breeding and production are still very limited as compared to other crops, which may be partly due to the lack of knowledge of their genetics and breeding system (Chakravarty, 2011). Linkage mapping is usually conducted in purpose-created segregating populations, such as progeny of selected parents. However, the resolution of linkage mapping is hampered by the limited number of recombination in the segregating population. Moreover, there are some particularities in linkage mapping in edible mushrooms. Construction of genetic map is performed using a haploid progeny, whereas the phenotypic evaluation of some traits is only possible at the dikaryotic stage, after crossing haploid progenies with the compatible monokaryotic tester (Foulongne-Oriol, 2012). For each locus, these mushrooms have the same allele from the tester line in one nucleus and contain the allele from one of the parents in the other nucleus (Gao et al., 2015). The identified QTLs reflect the allelic substitution e ect of the segregating allele with their

^bmarkers detected by Gong et al. (2016);

^cmarkers detected by Xiang (2015);

P FDR: P value after FDR correction;

R²: Phenotypic variation explained by each marker;

^{*}Scaffolds names are derived from the L54A reference genome.

^{*}Original P values detected by TASSEL 3.0.

TABLE 6 | Markers associated with at least two traits.

Marker	Number of associated traits	Traits ^a
S106_inID1	2	PD (2013); WF (2014)
S108_ID1	2	PW (2013); SW (2014)
S127_ID1	4	PD, PW (2013); WF, SW, PW (2014)
S178_ID1	2	WF, SW (2014)
S255_ID1	2	WF, PW (2014)
S258_ID5_2	2	PW, SW (2014)
S278_ID10	4	PD, PW (2013); WF, PD, PW, SW (2014)
S278_ID33	2	SW, PW (2014)
S278_ID36	3	PD (2013); WF, PW (2014)
S278_ID41	2	PD, PW (2013); PD (2014)
S328_ID5	2	PW, WF (2014)
S443_inID1	2	PW (2013); WF (2014)
S470_ID1	2	SW, WF (2014)
S473_ID1	2	PD, PW (2013)
S488_ID1	2	WF, PW (2014)
S553_ID5	2	SW, PW (2014)
S560_ID1	3	WF, PW, PD (2014)
S636_inID1	2	WF, SW (2014)
S704_inID1	3	PD, PW (2013); PW, WF, PD (2014)
S90_E1	2	SW, PW (2014)
S95_ID5	2	PW, WF (2014)

^aNumber in brackets indicates the year when the trait-marker associations were detected.

interactions with the constant allele from the tester nucleus, and such constraints may lead to inconsistencies in QTL detection (Foulongne-Oriol, 2012). Alternatively, association mapping compares favorably to linkage mapping for the dissection of natural variation by using diverse germplasms, such as those derived from the wild populations, germplasm collections or subsets of breeding germplasm (Zhu et al., 2008). Because many generations have passed and more recombinations occurred, the resolution of association mapping is considerably higher than that in simple bi-parental populations (Rafalski, 2010). However, the complex population structure is one of the sources of false positives in association mapping. For instance, the complex and heterogeneous population structure of a S. cerevisiae population was reported to lead to a high type I error rate in association mapping (Connelly and Akey, 2012). A discernable population structure was also reported in shiitake cultivars here. Moreover, the allele frequency distribution has considerable impact on the detection power of association mapping. Here, only the markers with a (MAF) >0.05 were utilized for association analysis.

Overall, the combination of linkage mapping and association mapping could be a more powerful strategy for dissecting the

REFERENCES

Abdurakhmonov, I. Y., and Abdukarimov, A. (2008). Application of association mapping to understanding the genetic diversity of plant germplasm resources. Int. J. Plant Genomics. 2008:574927. doi: 10.1155/2008/574927 genetic architectures of quantitative traits in edible mushrooms. To begin with, the genomic regions underlying quantitative traits of interest could be defined by linkage mapping. Then, based on the results of linkage mapping, fine mapping could be performed by candidate gene-based association analysis to identify the loci or genes for the traits of interest. In addition, the utilization of nested association mapping populations (Yu et al., 2008) could be a promising approach for genetic dissection of quantitative traits in edible mushrooms.

CONCLUSION

In summary, we reported here the genetic diversity, population structure and association mapping of agronomic traits in a Chinese L. edodes population containing 89 cultivars by using 297 genome-wide markers. A narrow genetic base with a discernable population structure was observed in the Chinese shiitake cultivars. In association mapping, a total of 43 markers were detected to be significantly associated with four traits. Five of these marker-trait associations were verified by previous studies and another five of them were significantly detected in cultivation tests performed in two consecutive years. Our results have highlighted the significant potential of LD-based association mapping of complex agronomic traits in shiitake with consideration of the population structure. Associations identified here could provide insights into the genetic architecture of important agronomic traits, thus paving a way toward implementation of MAS in L. edodes.

AUTHOR CONTRIBUTIONS

YX and YB conceived and designed the experiments; CL and LZ performed the experiments; WN and HK developed the molecular markers; CL, WG, and ZY analyzed the phenotypic and genotypic data; YX, WG, and MC wrote the paper.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb. 2017.00237/full#supplementary-material

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. B 57, 289–300.

Bradbury, P. J., Zhang, Z., Kroon, D. E., Casstevens, T. M., Ramdoss, Y., and Buckler, E. S. (2007). TASSEL: software for association mapping

- of complex traits in diverse samples. *Bioinformatics* 23, 2633–2635. doi: 10.1093/bioinformatics/btm308
- Celik, I., Camci, H., Kose, A., Kosar, F. C., Doganlar, S., and Frary, A. (2016). Molecular genetic diversity and association mapping of morphine content and agronomic traits in Turkish opium poppy (*Papaver somniferum*) germplasm. *Mol. Breed.* 36, 1–13. doi: 10.1007/s11032-016-0469-8
- Chakravarty, B. (2011). Trends in mushroom cultivation and breeding. Aust. J. Agric. Eng. 2, 102–109. Available online at: https://search.informit.com.au/documentSummary;dn=683694909801019:res=IELENG
- Chang, S. (1980). Mushrooms as human food. Bioscience 30, 399–401. doi: 10.2307/1308002
- Chen, Y., and Lübberstedt, T. (2010). Molecular basis of trait correlations. *Trends Plant Sci.* 15, 454–461. doi: 10.1016/j.tplants.2010.05.004
- Chiu, S. W., Ma, A. M., Lin, F. C., and Moore, D. (1996). Genetic homogeneity of cultivated strains of shiitake (*Lentinula edodes*) used in China as revealed by the polymerase chain reaction. *Mycol. Res.* 100, 1393–1399. doi: 10.1016/S0953-7562(96)80069-4
- Conesa, A., Götz, S., García-Gómez, J. M., Terol, J., Talón, M., and Robles, M. (2005). Blast2GO: a universal tool for annotation, visualization and analysis in functional genomics research. *Bioinformatics* 21, 3674–3676. doi: 10.1093/bioinformatics/bti610
- Connelly, C. F., and Akey, J. M. (2012). On the prospects of whole-genome association mapping in *Saccharomyces cerevisiae*. *Genetics* 191, 1345–1353. doi: 10.1534/genetics.112.141168
- Cook, J. P., McMullen, M. D., Holland, J. B., Tian, F., Bradbury, P., Ross-Ibarra, J., et al. (2012). Genetic architecture of maize kernel composition in the nested association mapping and inbred association panels. *Plant Physiol.* 158, 824–834. doi: 10.1104/pp.111.185033
- Dalman, K., Himmelstrand, K., Olson, Å., Lind, M., Brandström-Durling, M., and Stenlid, J. (2013). A genome-wide association study identifies genomic regions for virulence in the non-model organism *Heterobasidion annosum* s.s. *PLoS ONE* 8:e53525. doi: 10.1371/journal.pone.0053525
- Di Piero, R. M., de Novaes, Q. S., and Pascholati, S. F. (2010). E ect of *Agaricus brasiliensis* and *Lentinula edodes* mushrooms on the infection of passionflower with Cowpea aphid-borne mosaic virus. *Braz. Arch. Biol. Technol.* 53, 269–278. doi: 10.1590/S1516-89132010000200004
- Evanno, G., Regnaut, S., and Goudet, J. (2005). Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study. *Mol. Ecol.* 14, 2611–2620. doi: 10.1111/j.1365-294X.2005.02553.x
- Fan, L., Pan, H., Soccol, A. T., Pandey, A., and Soccol, C. R. (2006). Advances in mushroom research in the last decade. Food Sci. Biotechnol. 44, 303–311. Available online at: http://mychagaworld.com/pdf/Advances%20in%20Mushroom%20Research%20in%20the%20Last%20Decade.pdf
- Font i Forcada, C., Oraguzie, N., Reyes-Chin-Wo, S., Espiau, M. T., and Fernández i Martí, A. (2015). Identification of genetic loci associated with quality traits in almond via association mapping. PLoS ONE 10:e0127656. doi: 10.1371/journal.pone.0127656

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- Santoyo, F., González, A. E., Terrón, M. C., Ramírez, L., and Pisabarro, A. G. (2008). Quantitative linkage mapping of lignin-degrading enzymatic activities in *Pleurotus ostreatus*. Enzyme Microb. Technol. 43, 137–143. doi:10.1016/j.enzmictec.2007.11.007
- Silva, E. M., Machuca, A., and Milagres, A. M. (2005). E ect of cereal brans on *Lentinula edodes* growth and enzyme activities during cultivation on forestry waste. *Lett. Appl. Microbiol.* 40, 283–288. doi: 10.1111/j.1472-765X.2005.01669.x
- Slovak, R., Göschl, C., Su, X., Shimotani, K., Shiina, T., and Busch, W. (2014). A scalable open-source pipeline for large-scale root phenotyping of arabidopsis. Plant Cell 26, 2390–2403. doi: 10.1105/tpc.114.124032
- Soto-Cerda, B. J., and Cloutier, S. (2012). "Association mapping in plant genomes," in *The Genetic Diversity in Plants*, ed M. Caliskan (Rijeka: InTech Open Access Publisher). 29–54.
- Talas, F., Wüerschum, T., Reif, J. C., Parzies, H. K., and Miedaner, T. (2012). Association of single nucleotide polymorphic sites in candidate genes with aggressiveness and deoxynivalenol production in *Fusarium graminearum* causing wheat head blight. *BMC Genet.* 13:14. doi: 10.1186/1471-2156-13-14
- Wen, Z., Tan, R., Yuan, J., Bales, C., Du, W., Zhang, S., et al. (2014). Genome-wide association mapping of quantitative resistance to sudden death syndrome in soybean. BMC Genomics 15:809. doi: 10.1186/1471-2164-15-809
- Xiang, X. (2015). Association Analysis of Important Quantitative Traits based on Candidte-Gene Markers in Lentinula edodes. Master's thesis, Huazhong Agricultural University, Wuhan.
- Xiang, X., Li, C., Li, L., Bian, Y., Kwan, H. S., Nong, W., et al. (2016). Genetic diversity and population structure of Chinese *Lentinula edodes* revealed by InDel and SSR markers. *Mycol. Prog.* 15, 1–13. doi: 10.1007/s11557-016-1183-y
- Xu, X., Yang, J., Ning, Z., and Zhang, X. (2015). Lentinula edodes-derived polysaccharide rejuvenates mice in terms of immune responses and gut microbiota. Food Funct. 6, 2653–2663. doi: 10.1039/C5FO00689A

- Yan, J., Warburton, M., and Crouch, J. (2011). Association mapping for enhancing maize (Zea mays L.) genetic improvement. Crop Sci. 51, 433–449. doi: 10.2135/cropsci2010.04.0233
- Yeh, F. (1997). Population genetic analysis of codominant and dominant markers and quantitative traits. Belg. J. Bot. 129, 157.
- Yu, J., and Buckler, E. S. (2006). Genetic association mapping and genome organization of maize. Curr. Opin. Biotechnol. 17, 155–160. doi:10.1016/j.copbio.2006.02.003
- Yu, J., Holland, J. B., McMullen, M. D., and Buckler, E. S. (2008). Genetic design and statistical power of nested association mapping in maize. *Genetics* 178, 539–551. doi: 10.1534/genetics.107.074245
- Zhang, R., Huang, C., Zheng, S., Zhang, J., Ng, T. B., Jiang, R., et al. (2007). Strain-typing of Lentinula edodes in China with inter simple sequence repeat markers. Appl. Microbiol. Biotechnol. 74, 140–145. doi: 10.1007/s00253-006-0628-7
- Zhu, C., Gore, M., Buckler, E. S., and Yu, J. (2008). Status and prospects of association mapping in plants. *Plant Genome* 1, 5–20. doi:10.3835/plantgenome2008.02.0089

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