Evaluation of Genetic Diversity and Population Structure Analysis among Germplasm of *Agaricus bisporus* by SSR Markers

Hyejin An¹ᵃ, Hwa-Yong Lee¹ᵇ, Hyeran Shin¹, Jun Hyoung Bang¹, Seahee Han¹, Youn-Lee Oh¹ᶜ, Kab-Yeul Jang¹, Hyunwoo Cho¹, Tae Kyung Hyun¹, Jwakyung Sung¹, Yoon-Sup So¹, Ick-Hyun Jo¹ and Jong-Wook Chung¹

¹Department of Industrial Plant Science and Technology, Chungbuk National University, Cheongju, Republic of Korea; ²Department of Forest Science, Chungbuk National University, Cheongju, Republic of Korea; ³Department of Herbal Crop Research, National Institute of Horticultural and Herbal Science, Eumseong, Republic of Korea; ⁴Department of Crop Science, Chungbuk National University, Cheongju, Republic of Korea

**ABSTRACT**

*Agaricus bisporus* is a popular edible mushroom that is cultivated worldwide. Due to its secondary homothallic nature, cultivated *A. bisporus* strains have low genetic diversity, and breeding novel strains is challenging. The aim of this study was to investigate the genetic diversity and population structure of globally collected *A. bisporus* strains using simple sequence repeat (SSR) markers. *Agaricus bisporus* strains were divided based on genetic distance-based groups and model-based subpopulations. The major allele frequency (MAF), number of genotypes (NG), number of alleles (NA), observed heterozygosity (HO), expected heterozygosity (HE), and polymorphic information content (PIC) were calculated, and genetic distance, population structure, genetic differentiation, and Hardy–Weinberg equilibrium (HWE) were assessed. Strains were divided into two groups by distance-based analysis and into three subpopulations by model-based analysis. Strains in subpopulations POP A and POP B were included in Group I, and strains in subpopulation POP C were included in Group II. Genetic differentiation between strains was 99%. Marker AB-gSSR-1057 in Group II and POP B were included in Group I, and strains in subpopulation POP C were included in Group II. Genetic differentiation between strains was 99%. Marker AB-gSSR-1057 in Group II and subpopulation POP C was confirmed to be in HWE. These results will enhance *A. bisporus* breeding programs and support the protection of genetic resources.

1. Introduction

*Agaricus bisporus* is a popular edible mushroom that constitutes approximately 15% of global mushroom production [1]. Consumption of *A. bisporus* has increased in recent years alongside growth of the mushroom market, and *A. bisporus* is considered to be particularly healthy mushroom due to its higher protein, fiber, and amino acid contents and lower calories than other cultivated mushrooms such as *Pleurotus ostreatus* and *Lentinula edodes* [2]. Furthermore, extracts of *A. bisporus* have high antioxidant activities and may help to prevent breast cancer and cardiovascular disease [3,4]. *Agaricus bisporus*, which was first cultivated in France in the seventeenth century [5], is mainly cultivated in North America, Europe, India, and China [6], with recent increases in China and South Korea [5].

The growing demand for mushroom crops has increased the importance of developing new *A. bisporus* cultivars. New cultivars can be developed by selection of favorable traits from existing stocks and by introduction of traits from new genetic resources [7]. Furthermore, use of elite cultivars to introduce traits of interest in breeding programs can facilitate the development of novel cultivars [8,9]. Cultivars of *A. bisporus*, a secondary homothallic fungus, have low genetic diversity [10] owing to genetic erosion caused by the use of limited genetic resources for cultivar development [5,11]. Studies of genetic diversity and population structure have provided essential insights into potential genetic resources for crop breeding [12]. However, as the development of new cultivars from existing cultivars narrows the genetic relationships between breeding parents, accurate information on genetic diversity and relationships among strains is needed to support breeding goals [13]. Phenotypic, biochemical, and molecular markers can all be used to evaluate genetic diversity but, unlike phenotypic and biochemical markers, molecular markers are unaffected by the

**CONTACT** Ick-Hyun Jo ¹ Intron@korea.kr; Jong-Wook Chung ¹ jwchung73@chungbuk.ac.kr

¹These authors contributed equally to this work.

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Molecular markers such as restriction fragment length polymorphisms (RFLPs), random amplified polymorphic DNA (RAPD) markers, amplified fragment length polymorphisms (AFLP), simple sequence repeats (SSRs), and single nucleotide polymorphisms (SNPs) have been used for A. bisporus genotyping in a range of studies [15].

Advances in next-generation sequencing technologies have facilitated the sequencing of whole genomes, and the development of SSR and SNP markers for assessing variability among strains has actively progressed [16]. SSR markers have several advantages over many of the other available marker types, including codominance, high polymorphism levels, reproducibility, reliability, and genome-wide distribution [17]. SSRs in non-coding regions of the genome are particularly valuable as they display higher polymorphism levels than markers in other regions and are extremely useful for analysis of genetic diversity, population structure, and cultivar variability [18–20].

Previous studies investigated the genetic diversity and population structures of several edible mushrooms, including L. edodes [20–22], Flammulina velutipes [23–25], and Auricularia auricula-judae [26]. In A. bisporus, previous studies assessed genetic diversity and population structure using RFLP markers [27], RAPD markers [28,29], SSR markers [30–32], and Inter Simple Sequence Repeat (ISSR) markers [33,34]. An extensive evaluation of commercially cultivated, genetically similar A. bisporus resources is therefore needed to support breeding programs. In this study, 40 SSR markers distributed across the genome were used to examine the genetic diversity and population structure of 156 A. bisporus strains collected from around the world.

2. Materials and methods

2.1. Agaricus bisporus strains and SSR markers

In total, 156 A. bisporus strains collected from markets around the world were used in this study. Strains were deposited and preserved at the Mushroom Division of the National Horticultural Science Academy, Rural Development Administration, Republic of Korea (Table S1). Strains were cultured in Petri dishes loaded with cel-lophane on compost dextrose agar (CDA) for 60 days at 25°C in darkness. Cultured mycelia were harvested, freeze-dried for 4 days, and homogenized. DNA was extracted using a Plant SV mini kit (GeneAll, Seoul, Korea) in accordance with the manufacturer’s instructions, quantified using an Epoch Microplate Spectrophotometer (BioTek, Winooski, VT, USA), and adjusted to a final concentration of 20 ng/µL. Forty SSR markers with high polymorphic information content (PIC) values were selected, and allele counts for the 40 SSR markers were determined using the methods of Lee et al. [31] and An et al. [32] (Table S2).

2.2. PCR and genotyping

For PCR, 20 µL Excel TB 2× Taq Pre-Mix (Inclone Biotech, Yongin, Korea), 2 µL each primer (10 pmol), and 3 µL template DNA were combined in a final reaction volume of 40 µL. PCR was performed at 95°C for 2 min; 30 cycles at 95°C for 20 s, 55°C for 40 s, and 72°C for 45 s; and a final extension at 72°C for 10 min. PCR product sizes were determined using a Fragment Analyzer (Advanced Analytical Technologies Inc., Santa Clara, CA, USA) and by genotyping using Pro Size® 2.0 software (Advanced Analytical Technologies Inc.).

2.3. Data analysis

Genotype data for each individual were scored and imported into PowerMarker ver. 3.25 [35] for calculation of major allele frequency (MAF), number of genotypes (NG), number of alleles (NA), observed heterozygosity (HO), expected heterozygosity (HE), and polymorphism information content (PIC) values. A genotype accumulation curve was plotted using the R studio package “poppr” to determine the power of increasing numbers of SSR markers to distinguish individual genotypes [36]. Agaricus bisporus strains were divided based on genetic distance-based groups and by model-based subpopulations. A phylogenetic tree was constructed using MEGA ver. 5.2 using the unweighted pair inference in accordance with genetic distances determined using the Nei method [37,38]. Population structure was analyzed using STRUCTURE ver. 2.3.1 [39]. The number of subpopulations (K) was assumed to be in the 1–10 range, and the length of burn-in and number of Monte Carlo Markov chains (MCMCs) were 10,000 and 100,000, respectively, with five replicates. STRUCTURE HARVESTER was used to determine delta K from the derived results [40]. Each strain was divided into subpopulations or admixtures (Admix) based on the probability of belonging to each subpopulation. Strains sharing more than 80% of ancestry were assigned to one subpopulation. Strains and subpopulation were presented on an unrooted tree to reveal structured relationships between strains and subpopulations. Analysis of molecular variance (AMOVA) using GenAlex 6.41 was performed to determine the degree of genetic variation and strain differentiation.
within each group, and Hardy–Weinberg equilibrium (HWE) was assessed using “pegas” in the R studio [41,42].

3. Results

3.1. SSR polymorphisms and marker set selection

A population of 156 A. bisporus strains was genotyped using 40 SSR markers. Upon assessment of the 156 strains with the full set of 40 SSR, the major allele frequency (MAF) ranged from 0.20 (AB-gSSR-0182, AB-gSSR-1184) to 0.87 (AB-gSSR-1058), with an average of 0.431. The number of genotypes (NG) ranged from 3 (AB-gSSR-1058) to 33 (AB-gSSR-0182, AB-gSSR-1184) to 0.87 (AB-gSSR-1058), with an average of 0.660.

The polymorphic information content (PIC) value ranged from 0.21 (AB-gSSR-1058) to 0.88 (AB-gSSR-1184), with an average of 0.697. The polymorphic information content (PIC) value ranged from 0.21 (AB-gSSR-1058) to 0.87 (AB-gSSR-1184), with an average of 0.660 (Table 1). A genotype accumulation curve for the population showed that the genotypes of the 156 strains could be distinguished using as few as seven markers (Figure 1). Differences among strains were confirmed by construction of a phylogenetic UPGMA tree based on a combination of the seven markers (Figure S1).

3.2. Phylogenetic relationship and population structure of A. bisporus strains

Phylogenetic analysis of the 156 A. bisporus strains divided the population into two groups: Group I and Group II (Figure 2). Group I comprised 122 strains collected from Australia (1), Belgium (1), Brazil (1), Cambodia (1), Canada (3), China (6), France (5), Germany (9), Indonesia (2), Italy (1), Japan (7), Netherlands (3), New Zealand (2), Peru (1), South Korea (39), Switzerland (1), Thailand (3), UK (9), USA (26), and Vietnam (1). Group II comprised 34 strains from Brazil (3), Canada (6), France (2), Germany (3), Japan (3), Netherlands (2), New Zealand (2), Peru (1), South Korea (8), and USA (4). Grouping was not in accordance with the country origins of the strains (Table S1). The optimal number of subpopulations (K) for division of the A. bisporus population was three (Figure 3(A)). Strains were allocated to a subpopulation when the probability of belonging to that subpopulation was >80% (Figure 3(B)). Subpopulation A (POP A) comprised 51 strains collected from Australia (1), Brazil (1), Canada (2), China (4), Germany (4), UK (2), Italy (1), Japan (3), Cambodia (1), South Korea (21), Netherlands (2), New Zealand (2), Peru (1), and USA (6). Subpopulation B (POP B) comprised 57 strains from Canada (1), Germany (5), France (3), UK (7), Indonesia (2), Japan (3), South Korea (13), Netherlands (1), Thailand (3), and USA (19). Subpopulation C (POP C) comprised 33 strains from Brazil (3), Canada (6), Germany (3), France (2), Japan (3), South Korea (8), Netherlands (2), New Zealand (2), and USA (4). The remaining 15 strains, from Belgium (1), Switzerland (1), China (2), France (2), Japan (1), South Korea (5), Peru (1), USA (1), and Vietnam (1) were included in Admix (Table S1). An unrooted tree displayed the clear subpopulation structure (Figure 3(C)).

Table 1. Genetic diversity parameters of 40 SSR markers with 156 Agaricus bisporus strains.

| Marker       | MAF | NG | NA | HE^a | HO^a | PIC^b |
|--------------|-----|----|----|------|------|-------|
| AB-gSSR-0113 | 0.67 | 12 | 7  | 0.52 | 0.31 | 0.48  |
| AB-gSSR-0139 | 0.73 | 6  | 5  | 0.43 | 0.03 | 0.49  |
| AB-gSSR-0182 | 0.20 | 24 | 13 | 0.85 | 0.58 | 0.83  |
| AB-gSSR-0199 | 0.68 | 14 | 9  | 0.50 | 0.13 | 0.45  |
| AB-gSSR-0238 | 0.32 | 15 | 9  | 0.78 | 0.65 | 0.75  |
| AB-gSSR-0489 | 0.30 | 14 | 10 | 0.81 | 0.20 | 0.78  |
| AB-gSSR-0532 | 0.32 | 19 | 10 | 0.81 | 0.20 | 0.78  |
| AB-gSSR-0564 | 0.64 | 9  | 7  | 0.53 | 0.03 | 0.48  |
| AB-gSSR-0574 | 0.61 | 9  | 6  | 0.57 | 0.46 | 0.54  |
| AB-gSSR-0584 | 0.51 | 18 | 8  | 0.66 | 0.35 | 0.61  |
| AB-gSSR-0603 | 0.28 | 20 | 11 | 0.80 | 0.59 | 0.77  |
| AB-gSSR-0611 | 0.35 | 15 | 9  | 0.77 | 0.48 | 0.74  |
| AB-gSSR-0709 | 0.41 | 12 | 8  | 0.67 | 0.97 | 0.61  |
| AB-gSSR-0713 | 0.35 | 23 | 20 | 0.80 | 0.94 | 0.78  |
| AB-gSSR-0811 | 0.33 | 24 | 13 | 0.79 | 0.88 | 0.77  |
| AB-gSSR-0816 | 0.45 | 14 | 12 | 0.68 | 0.43 | 0.62  |
| AB-gSSR-0837 | 0.41 | 17 | 16 | 0.80 | 0.06 | 0.77  |
| AB-gSSR-0860 | 0.47 | 10 | 10 | 0.56 | 0.02 | 0.46  |
| AB-gSSR-0900 | 0.46 | 23 | 16 | 0.68 | 0.31 | 0.63  |
| AB-gSSR-0913 | 0.30 | 14 | 9  | 0.69 | 0.26 | 0.66  |
| AB-gSSR-0923 | 0.34 | 25 | 13 | 0.80 | 0.59 | 0.77  |
| AB-gSSR-0940 | 0.28 | 13 | 25 | 0.81 | 0.83 | 0.79  |
| AB-gSSR-0959 | 0.36 | 14 | 9  | 0.78 | 0.27 | 0.76  |
| AB-gSSR-1004 | 0.44 | 16 | 12 | 0.74 | 0.41 | 0.71  |
| AB-gSSR-1018 | 0.31 | 23 | 12 | 0.81 | 0.80 | 0.78  |
| AB-gSSR-1036 | 0.68 | 4  | 3  | 0.44 | 0.01 | 0.34  |
| AB-gSSR-1044 | 0.51 | 12 | 9  | 0.65 | 0.22 | 0.60  |
| AB-gSSR-1052 | 0.22 | 27 | 18 | 0.85 | 0.60 | 0.84  |
| AB-gSSR-1057 | 0.40 | 21 | 11 | 0.76 | 0.59 | 0.73  |
| AB-gSSR-1058 | 0.87 | 3  | 3  | 0.23 | 0.00 | 0.21  |
| AB-gSSR-1064 | 0.63 | 13 | 7  | 0.55 | 0.28 | 0.52  |
| AB-gSSR-1080 | 0.41 | 17 | 9  | 0.76 | 0.28 | 0.73  |
| AB-gSSR-1122 | 0.45 | 14 | 7  | 0.71 | 0.83 | 0.67  |
| AB-gSSR-1142 | 0.27 | 17 | 9  | 0.82 | 0.89 | 0.80  |
| AB-gSSR-1180 | 0.42 | 11 | 9  | 0.73 | 0.96 | 0.70  |
| AB-gSSR-1184 | 0.20 | 32 | 19 | 0.88 | 0.79 | 0.87  |
| AB-gSSR-1189 | 0.25 | 21 | 11 | 0.84 | 0.24 | 0.82  |
| AB-gSSR-1202 | 0.36 | 14 | 11 | 0.72 | 0.31 | 0.67  |
| AB-gSSR-1208 | 0.60 | 6  | 6  | 0.53 | 0.00 | 0.45  |
| AB-gSSR-1247 | 0.35 | 15 | 9  | 0.76 | 0.71 | 0.72  |
| Mean         | 0.431 | 16.3 | 10.5 | 0.697 | 0.441 | 0.660 |

^aMAF: major allele frequency.

^bNG: number of genotypes.

^cNA: number of alleles.

^dHE: expected heterozygosity.

^eHO: observed heterozygosity.

^fPIC: polymorphic information content.

^gLee et al. (2018).

^hAn et al. (2019).
3.3. Genetic diversity

Strains were considered according to distance-based grouping. MAF was 0.468 for Group I and 0.541 for Group II, and HO was 0.439 for Group I and 0.449 for Group II. NG and NA values for Group I were approximately double those for Group II, at 13.6 and 9.1 for Group I and 6.1 and 5.8 for Group II, respectively. Diversity index He and PIC values were higher in Group I than in Group II, at 0.664, 0.439, 0.624, and 0.571, 0.449, 0.528 in Group I and 0.547, 0.452, 0.519 in Group II, respectively (Table 2).

Table 2. Genetic diversity indices for distance-based groups and model-based subpopulations of 156 Agaricus bisporus strains.

| Group          | MAF a | NG b | NA c | HE d | HO e | PIC f |
|----------------|-------|------|------|------|------|-------|
| Distance-based |       |      |      |      |      |       |
| Group I (122)  | 0.468 | 13.6 | 9.1  | 0.664| 0.439| 0.624 |
| Group II (34)  | 0.541 | 6.1  | 5.8  | 0.571| 0.449| 0.528 |
| Model-based    |       |      |      |      |      |       |
| POP A (51)     | 0.509 | 7.7  | 6.3  | 0.605| 0.393| 0.553 |
| POP B (57)     | 0.532 | 7.0  | 6.1  | 0.581| 0.470| 0.535 |
| POP C (33)     | 0.547 | 5.6  | 5.5  | 0.562| 0.452| 0.519 |
| Admix (15)     | 0.409 | 7.4  | 6.3  | 0.715| 0.467| 0.677 |

aMAF: major allele frequency. 
bNG: number of genotypes. 
cNA: number of alleles. 
dHE: expected heterozygosity. 
eHO: observed heterozygosity. 
fPIC: polymorphic information content.

3.4. Analysis of molecular variance and HWE

AMOVA was used to examine genetic variation for each group and subpopulation, and a fixed index $F_{st}$ value was determined to confirm differentiation. Upon division of strains into genetic distance-based groups or model-based subpopulations, approximately 100% and 99% of the total genetic variation were observed between strains and $F_{st}$ values were $-0.002$ and $0.013$, respectively (Table 3).

HWE was used to assess the genetic constitution of each group and subpopulation (Figure 4). In the distance-based groups, strains in Group I deviated significantly from HWE for all markers, and those in Group II were in HWE for AB-gSSR-1057. For the model-based subpopulations, strains in POP A and POP B deviated significantly from HWE for all markers, while those in POP C were in HWE only for AB-gSSR-1057. Admix was confirmed to be in HWE for 11 markers (AB-gSSR-0182, AB-gSSR-0532, AB-gSSR-0574, AB-gSSR-0584, AB-gSSR-0811, AB-gSSR-0816, AB-gSSR-0940, AB-gSSR-1018, AB-gSSR-1122, AB-gSSR-1180, and AB-gSSR-1189).

4. Discussion

In this study, it was possible to discriminate 156 globally collected A. bisporus strains with 40 SSR
markers, confirming the efficiency of the SSR markers used in this study. The *A. bisporus* strains were not clustered by geographical distribution. Several studies have reported that the relationship between phylogenetic analysis and geographical distribution of the mushroom strains is not high. In the study of An et al. analyzed 26 *A. bisporus* strains using 170 genomic SSR markers, the stains were not geographical clustered [32] and *L. edodes* wild strains in Korea were not clustered by collected regions [22].

*Agaricus bisporus* resources are widely sold in markets in many countries. A genotype accumulation curve indicated that *A. bisporus* strains could be distinguished using as few as seven SSR markers, and complete recognition of all multilocus genotypes was possible using 96 alleles (AB-gSSR-0182, 13 alleles; AB-gSSR-0238, 9; AB-gSSR-1018, 12; AB-gSSR-1184, 19; AB-gSSR-0913, 9; AB-gSSR-0940, 25; AB-gSSR-1080, 9), indicating that the seven SSR markers had the capacity to accurately identify the 156 globally collected *A. bisporus* strains (Table 1 and Figures 1 and 2). The genotype accumulation curve allowed individual samples to be distinguished from random samples of n loci [43]. The minimum number of markers can be readily determined using the accumulation curve method, potentially helping to refine minimum marker sets needed for sample discrimination.

For the set of 156 *A. bisporus* strains assessed with 40 SSR markers in this study, average MAF, NG, NA, HE, HO, and PIC value were 0.431, 16.3, 10.5, 0.697, 0.441, and 0.660, respectively. Among the diversity indices, HE and HO are sensitive to allele frequency [44], and allele frequency helps to determine the PIC value [45]. These diversity
indices were higher than those reported for other mushrooms. NA and PIC values were 0.47 and 0.47 for \textit{A. auricular-judae}, respectively [26]; NA and PIC values were 2.9 and 0.43 for \textit{F. velutipes}, respectively [25]; and NA, HE, HO, and PIC values were 5.5, 4.9, 0.552, 0.309, and 0.51 for \textit{L. edodes}, respectively [20]. Furthermore, the diversity indices in this study were higher than in recent \textit{A. bisporus} studies. NA, HE, HO, and PIC values were 5, 0.68, 0.53, and 0.62, in an analysis of \textit{A. bisporus} cultivars from USA (6), China (4), Netherlands (2), England (1), Germany (1), and Spain (1), and 13 wild strains from China, with 17 SSR markers conducted by Fu et al. [19]. NG, NA, HE, HO, and PIC values were 6.17, 5.47, 0.619, 0.227, and 0.569 in a study of 26 \textit{A. bisporus} strains with 121 markers conducted by An et al. [32]. The genetic diversity of the \textit{A. bisporus} strains used in this study was higher than in other studies.

Analysis of genetic diversity and population structure provides valuable information on genetic resources for crop breeding [12]. Genetic diversity analysis of cultivars and wild strains is underway to aid the development of new \textit{A. bisporus} cultivars with desirable traits and improved adaptability. Diversity of \textit{A. bisporus} resources, cultivars, and wild strains was assessed using molecular RFLP, SSR, and SNP markers [30, 32, 46–48]. Genetic diversity was also assessed using SSR markers to incorporate disease resistance into breeding programs [18, 19]. Understanding genetic diversity is an important factor in maximizing crop yields and developing sustainable agriculture [49]. The globally collected strains in this study exhibited higher genetic diversity than other \textit{A. bisporus} collections, and the results of this study will facilitate the analysis and use of these diverse strains, as well as cultivars and wild strains, in breeding programs to develop new \textit{A. bisporus} cultivars with desirable traits.

In this study, the genetic diversity (HE and PIC values) of distance-based Group I was higher than that of Group II. Genetic diversity was generally higher for POP A and POP B than for POP C, and
diversity was higher for POP A than POP B, in model-based subpopulations. Consequently, the genetic diversity of Group I, which consisted of strains from POP A and POP B, was high. The 122 strains in Group I were from 20 countries, whereas the 34 strains in Group II were from 10 countries. POP A contained 51 strains from 14 countries, POP B contained 57 strains from 10 countries, and POP C contained 33 strains from 9 countries. Genetic diversity was high in groups and subpopulations, and all groups and subpopulations contained strains from diverse countries.

AMOVA for each grouping method revealed that >99% of genetic variation in *A. bisporus* occurred between individual strains, with little variation observed between groups or subpopulations. This was supportive of high rates of genetic exchange among strains. Factors that can contribute to deviations from HWE include nonrandom mating, migration, mutation, natural selection, and mixing of subpopulations [50]. The AB-gSSR-1057 marker was confirmed to be in HWE in Group II and POP C and was presumed to be indicative of the lower genetic diversity in Group II than in Group I.

In conclusion, this study screened high-efficiency SSR markers for their capacity to discriminate among *A. bisporus* strains collected worldwide. Genetic diversity was assessed using distance-based and model-based analysis. Analysis of genetic diversity not only facilitates germplasm preservation efforts, but can also provide guidance for better use of germplasms in genetic breeding programs [49]. The results of this study will be beneficial for the future development of improved cultivars of *A. bisporus*.

**Disclosure statement**

No potential conflict of interest was reported by the authors.

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