INTRODUCTION

Mushrooms have been appreciated across the world for their unique flavour and have been valued by mankind as a culinary wonder (Patel and Goyal, 2012). They have been consumed by man for their delicious taste and pleasing flavour and are the cheapest source of protein particularly for the vegetarians. Besides being rich source of proteins, mushrooms also contain fats, vitamins, crude fiber, carbohydrates and some essential minerals, which are the key factors for the normal functioning of the body (Sharma et al., 2017). They also contain certain important vitamins like thiamine, ascorbic acid, nicotinic acid, riboflavin, pantothenic acid and biotin. This low cost vegetable is not only packed with essential nutrients but also has the unique properties to fight many deadly diseases (Thakur and Singh, 2014). Cultivation of mushrooms is an eco-friendly activity, which represents unique exploitation of the microbial technology for the bioconversion of the unused lignocellulosic wastes into nutritious food. In India, cultivation of edible mushrooms started way back in 1961, but emphasis is more on the temperate mushroom, Agaricus bisporus. However, our country can excel in the mushroom industry only by cultivating and commercializing some of the tropical and subtropical mushrooms. One such edible mushroom of the tropical region is Macrocybe, which belongs to the Family Tricholomataceae of the Order Agaricales (Kirk et al., 2008). It is characterized by large fleshy sporophores, excellent shelf life, ability to grow at temperature above 30°C, high biological efficiency and easiness in post-harvest handling. Macrocybe resembles another edible summer mushroom, Calocybe very much as both have conspicuous large, saprophytic sporophores but it differs from Calocybe as it lacks siderophilous granulation in the basidia and differs at DNA level (Razaq et al., 2016). Macrocybe has seven well recognised species widely distributed in the tropical regions of the world (Pegler et al., 1998). However, from India, only five wild species of Macrocybe viz., M. crassa, M. pachymeres, M. gigantea, M. lobayensis and M. titans have been reported so far and cultivation of only M. gigantea has been attempted (Mohanan, 2011; Pamitha, 2014;
MATERIALS AND METHODS

Procurement of mushroom strain
Pure culture of M. gigantea was procured from Directorate of Mushroom Research (DMR), Solan, Himachal Pradesh and Sher-e-Kashmir University of Agricultural Sciences and Technology (SKUAST), Jammu. The strain used for fruiting ability was maintained on potato dextrose agar (PDA) and malt extract agar (MEA) medium at room temperature and subculturing was done regularly after three months to sustain their fruiting vigour.

Preparation of spawn
The spawn was prepared by using the method outlined by Munjal (1973). Grains of three different cereals viz., bajra (Pennisetum glaucum L.), wheat (Triticum aestivum L.) and maize (Zea mays L.) were collected from the local market, thoroughly washed and soaked in water for 5-6 hours. After that, the grains were boiled for 30 minutes, drained, air-dried and mixed with 2 percent gypsum (calcium sulphate) and 0.5% chalk powder (calcium carbonate) to maintain pH at 6.7 and keep the grains separate from each other. After proper mixing, the grains were put in thoroughly cleaned bottles (500 ml) and fresh polypropylene bags up to 3/4th of their capacity, plugged with non-absorbant cotton and autoclaved at 15lb/sq. inch for 2 hours. After cooling, the sterilized grains were inoculated with 3-5 uniform sized mycelium discs of M. gigantea and incubated at 28±2°C till the mycelium covered the surface of the grains completely. Downward linear growth of the mycelium started in each bottle/bag within 3-4 days, which completely covered the grains within 15-20 days (Fig. 1).

Collection and preparation of substrates
Agricultural wastes like wheat straw (T. aestivum L.) and paddy straw (Oryza sativa L.) were collected from the local fields of Jammu (India) and chopped into 5-7 cm pieces. Thereafter, they were subjected to hot water (80-90°C) dip treatment in a drum for 40-50 minutes. Excess water was drained off and the agrowastes were allowed to cool at room temperature and dry till 60% moisture was left. Substrates that on squeezing did not leak out drops of water were considered saturated with approximately 60% moisture level and were considered ready for use.

Spawning and spawn run
For the process of spawning, transparent polythene bags (9×16 inch size) were taken and filled up to 3/4th of their capacity with alternate layers of the pasteurized substrate and spawn (5%). The spawned bags were then transferred to hanging nets kept in the mushroom house. A temperature of 25-35°C and relative humidity of 80-90% was maintained in the mushroom house for spawn run. Each treatment was performed three times.

Casing of bags
After completion of the spawn run, the bags were opened and encased with 2-4 cm thick layer of farm yard manure (FYM), sterilized in an autoclave at 15 lbs psi for 60 minutes. Casing layer was kept moist by sprinkling water as and when required.

Cropping and harvesting
After casing, the bags were retained in the hanging nets. After a few days, the mycelium of M. gigantea sprouted in the form of needle shaped primordia, some of which matured into large sporophores (Fig. 2). The sporophores on maturation were harvested by twisting lightly. The sporophore count and its corresponding weight were measured after every harvest.

Biological efficiency (B . E)
It is an estimate of the ability of mushrooms to convert substrate into fruiting bodies. It was calculated as per the following formula:

\[ B.E = \frac{\text{Fresh weight of mushroom yield}}{\text{Dry weight of substrate used}} \times 100 \]  

Statistical analysis
Data were subjected to Analysis of Variance (ANOVA) and the least significant difference was determined at the 5% level by Duncan Post Hoc test using SPSS software package (Srikram and Supapvanich, 2016).

RESULTS AND DISCUSSION

During these studies, observations were made on various aspects like spawn development, growth behaviour, pinhead number, sporophore number, yield potential and biological efficiency of M. gigantea cultivated on...
wheat straw and paddy straw waste.

Assessment of duration required for spawn development of *M. gigantea* on different cereal grains

Results depicted in Fig. 3 showed that minimum time for spawn preparation was recorded on bajra grains (14.3 days) followed by wheat grains (17.6 days), whereas maximum time was taken by maize grains (20.3 days). Variation in the time period required for spawn preparation may be attributed to the size of the grains. During the present investigation, the small sized grains of bajra get quickly enveloped with the mushroom mycelium than the larger grains of wheat and maize. Other workers have used wheat grains (Akhtar *et al.*, 2018; Bharti, 2019), maize grains (Akhtar *et al.*, 2018) and paddy grains (Pamitha, 2014; Duong *et al.*, 2017) for spawn preparation of *M. gigantea* and have reported requirement of 15 to 25 days for its preparation. However, in view of the present observation, bajra grains are recommended for spawn preparation as it will affect the total duration required for the production of first flush of *M. gigantea* sporophores.

Assessment of growth behaviour

During the study, duration required for complete spawn run, pinhead formation and sporophore maturity of *M. gigantea* on two agrowastes (wheat straw and paddy straw) inoculated with three different grain spawns were assessed.

As depicted in Table 1, minimum period for spawn run (16.3 days) was recorded on wheat straw inoculated with bajra grain spawn. It was further observed that on paddy straw, all the three types of spawns made on bajra, wheat and maize grains showed non-significant differences in spawn run and took more time (up to 20.3 days). Earlier, few other researchers have also used wheat grain based spawn and they reported complete spawn run of *M. gigantea* on different agrowastes within 15-21 days of spawning (Kushwaha *et al.*, 2016; Bharti 2019).

Statistical analysis of the data presented in Table 1 revealed that emergence of primordia took place equally well on wheat and paddy straw inoculated with bajra, wheat and maize grain spawn. However, minimum period required for primordial emergence after farm yard manure (FYM) casing was recorded on the paddy straw inoculated with bajra grain spawn (14.6 days). Earlier, Kushwaha *et al.* (2016) and Bharti, (2019) have recorded the emergence of pinheads on the agrowastes within 16 to 21 days of the casing. In contrast to these findings, Akhtar *et al.* (2018) reported that pinhead initiation of *Tricholoma giganteum* (Syn. *Macrocybe gigantea*) on the wheat straw took place after
31-45 days of spawning by using wheat grain and maize grain based spawn.

Similarly, during the present investigation, non-significant differences were recorded with respect to the types of grain spawn used and the number of days required for obtaining mature sporophores (first flush). On the tested agrowastes inoculated with different spawns, minimum of 9.0 days and a maximum of 10.3 days were recorded for obtaining first flush after pinhead formation. However, in contrast to the present results, Bharti (2019) reported that by using wheat grain spawn, 13 to 17.87 days were required for the production of first flush of *M. gigantea* sporophores. Earlier, Duong *et al.* (2017) while cultivating another edible species of *Macrocybe*, *M. titans* by using paddy grain spawn found sporophores ready for harvesting within 7-10 days of pinhead formation.

**Assessment of pinhead and sporophore number**

Data presented in Table 2 showed non-significant differences with respect to number of pinheads on agrowastes spawned with three different types of spawn. Maximum (218.3) and minimum number (190.0) of pinheads were obtained on wheat straw inoculated with wheat grain spawn and bajra grain spawn respectively. The number of developing sporophores was also recorded as only few of the pinheads attained maturity. A perusal of data presented in Table 2 showed that paddy straw inoculated with bajra grain spawn proved to be the best as it resulted in the formation of the highest number of sporophores (20.6). This may be due to multilateral enzyme system of *M. gigantea*, which bio-degrades a large range of lignocellulosic wastes (Kumla *et al.*, 2020). The lowest number of sporophores (17.0) were recorded on wheat straw inoculated with maize grain spawn (Table 2). Statistically, differences were non-significant in term of sporophore number. Earlier, Kushwaha *et al.* (2016) recorded very high sporophore number of *M. gigantea* (54-110). However, Inyod *et al.*
Z. mays, a new record species for Vietnam (UGC The authors are thankful to the Department of Botany.

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Conflict of interest
The authors declare that they have no conflict of interest.

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Table 3. Sporophore yield (g) and biological efficiency (%) of M. gigantea on agrowastes inoculated with different grain spawns.

| Spawn prepared on | Wheat straw | Paddy straw | Biological efficiency (%) |
|------------------|-------------|-------------|--------------------------|
| Bajra grain      | 331.3 ± 3.93| 343.6 ± 2.96| 66.26                    |
| Wheat grain      | 332.0 ± 3.05| 327.0 ± 1.52| 66.4                     |
| Maize grain      | 320.0 ± 4.04| 334.6 ± 4.37| 64.0                     |
| F-value          | 3.39        | 6.90        |                          |
| P-value          | 0.05        | 0.05        |                          |

The values given are mean (n=3) ± standard error. Means followed by the same letter in the same column are not significantly different at 5% probability level according to the Duncan Post Hoc test.

(2016) recorded very few sporophores (4.35-5.75) of another species of Macrocybe, M. crassa that was cultivated on rubber tree sawdust.

Assessment of sporophore yield and biological efficiency

Data given in Table 3 revealed non-significant differences in the yield of M. gigantea on wheat straw inoculated with three different grain spawns. Maximum yield (343.6g/500g of dry substrate) was recorded on paddy straw spawned with bajra grain spawn, whereas minimum yield (320.0g) was obtained on wheat straw spawned with maize grain spawn. These results indicated that the grains used for spawn production significantly affect the sporophore yield of M. gigantea.

Biological efficiency obtained on the agrowastes was in accordance with the sporophore yield, being maximum (68.7%) on paddy straw and minimum on wheat straw (64.0). An increase in the biological efficiency of M. gigantea on paddy straw may be due to the appropriate nutrient content of the substrate, which affected the growth and formation of sporophores. Pamitha (2014), Kushwaha et al. (2016) and Bharti, (2019) have also reported similar range of yield and biological efficiency (724g/1000g of dry substrate, 72.4%; 4493.3g/6400g of dry substrate, 70.1%; 651.87g/1000g of dry substrate, 65.1% respectively) of M. gigantea on different agrowastes.

Conclusion

It can be concluded from the present study that bajra grains P. glaucum L.) are better than wheat grains (T. aestivum L.) and maize grains (Z. mays L.) for the spawn production of M. gigantea as it gets ready in minimum time period (14.3 days), shows quick run (16.3 days) on the agrowastes and produces maximum sporophores (20.6) within few days of casing with farm yard manure.

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