RESEARCH ARTICLE

ESTIMATION OF EXTRACELLULAR LIGNINOLYTIC ENZYMES FROM WILD AURICULARIA POLYTRICHA, HELVELLA SP. AND MORCHELLA SP.

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Abstract

Three mushroom samples have been selected for extracellular ligninolytic enzyme activities i.e commercially cultivated Auricularia polytricha, wild Helvella sp. and wild Morchella sp. To investigate the extracellular ligninolytic enzymes, cultures were incubated over a period of 20 days in nutrient rich medium (NRM) and nutrient poor medium (NPM) under static conditions. Samples showed green and brown coloration in solid medium plates; it indicates the presence of laccase and Manganese peroxidase (MnP). Production of extracellular ligninolytic enzymes were observed in all three species in different pattern. The results of the present study allow us to conclude that wild Morchella sp. and Helvella sp. are good for production of ligninolytic enzymes in comparison to commercially cultivated Auricularia polytricha.

Introduction:-

Fungi have a worldwide distribution and grow in a wide range of habitats including extreme environments such as desert area or areas with salt concentrations or ionizing radiation as well as in deep sea sediments. In addition, they have many medicinal uses and are good agents of bioremediation [1,19]. Mushrooms are highly nutritious containing protein (19-35%), low fat content (1.3-2%), relatively large amounts of carbohydrate (51-88%) and fiber (4-20%) in dry mushroom fruit bodies [2]. In the physiological point of view, mushrooms are broadly divided into wood rot, mycorrhizal and litter decomposing fungi. The most important and potent wood-destroying organisms are white rot and brown rot fungi. White rot fungi that play an important role in decomposition of dead trees, especially in degradation of lignin. The white rot fungi belonging to the basidiomycetes that produce various isoforms of extracellular ligninolytic enzymes. The main extracellular enzymes participating in lignin degradation are heme-containing lignin peroxidase (LiP; EC 1.11.1.14), MnP (EC 1.11.1.13) and Cu-containing laccase (EC 1.10.3.2). In addition, enzyme involved in hydrogen peroxide production such as aryl alcohol oxidase (AAO, EC 1.1.3.7) is considered to belong to the ligninolytic system [10].

We have selected three mushrooms strains, Auricularia polytricha, Helvella sp. and Morchella sp. for their extracellular enzymatic studies. The interest of Auricularia polytricha could be easily identified by pilose upper surface which is strongly capitates with dark brown smooth hymenium [13]. Morchella species is one of the most readily recognized of all the edible mushrooms. On the other hand Helvella species they grow in grass as well as humid hardwoods, such as beech, and in hedges and on the talus of meadows. These all are wood decaying WRF produce ligninolytic enzymes.

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Materials and Methods:-
Fresh fruiting bodies of *Morchella* and *Helvella* were collected from forest of Silly forest (1500 M), Himachal Pradesh during rainy season. The fruiting bodies were aseptically collected with the help of knife and carried to laboratory in sterile poly-bags. Mycelial culture of *Auricularia polytricha* was procured from Directorate of Mushroom Research, Solan (H.P.).

Chemicals Used:-
Analytical grade 2,2'-azino-bis (3-ethyl benzothiazoline-6-sulphonic acid) (ABTS, MW 548.7 gmol⁻¹) and Veratryl alcohol (VA, MW 168.19 gmol⁻¹) was purchased from Himedia. All other chemicals used were of analytical grade.

Screening of ligninolytic enzymes through spot test:-
Laccase Spot test:-
Laccase spot test was done for the screening of extracellular ligninolytic enzyme. Test was performed in 90mm diameter Petri-dishes with 20ml of the modified Kirk medium containing 10g of glucose, 2g of KH₂PO₄, 0.5g of MgSO₄.7H₂O, 0.1g of CaCl₂, 2.2g of 2,2-dimethylsuccinate, 0.5g of ammonium tartrate, 0.2g of yeast extract, 0.2g of ABTS, 20g of agar, per liter of medium. The pH was adjusted to 5.0 before autoclaving at 15psi and 120°C for 20 min. The plates were incubated at 25°C for three weeks. The extracellular enzyme ABTS oxidizing activity was measured by the green colour intensity of medium [21].

MnP Spot Test:-
Fungus culture (previously cultured in 2% MEA agar for 8 days) were inoculated in the kirk medium containing 10g of glucose, 2g of KH₂PO₄, 0.5g of MgSO₄.7H₂O, 0.1g of CaCl₂, 2.2g of 2,2-dimethylsuccinate, 0.5g of ammonium tartrate, 20g of agar-agar per liter of medium. For the production of MnP medium was supplemented with 0.04g of MnCl₂.4H₂O per liter of the medium. The pH was adjusted to 5.0 before autoclaving at 15 psi for 20 min at 120°C [22]. Fungal growth was estimated in terms of diameter of fungal colony. Extracellular enzymatic activity was measured by the colour intensity of the medium. Plates were observed once a day for three consecutive weeks.

Extracellular Ligninolytic Enzyme assay:-
The production of extracellular enzymes were carried out in the NRM containing 2g of ammonium tartrate, 10g of glucose, 1 g of KH₂PO₄, 1g of yeast extract, 0.5g of MgSO₄.7H₂O, 5g of KCl, 1ml of solution containing trace elements per liter of medium and the NPM containing 10g of glucose, 2g of KH₂PO₄, 0.2g of yeast extract, 0.1g of peptone, 1ml of solution containing trace elements per liter of medium. Solution of trace elements containing 10mg of Na₂B₄O₇.10H₂O, 7mg of ZnSO₄.7H₂O, 5mg of FeSO₄.7H₂O, 1mg of CuSO₄.5H₂O, 1mg of (NH₄)₆MoO₇.2.4H₂O, 1mg of MnSO₄ dissolved in 100 ml of H₂O [8]. The pH was adjusted to seven before autoclaving at 15 psi and 120°C for 20 min. 150 ml flasks containing 50 ml liquid medium were inoculated with 5-8 days old five mycelium bits (5mm in diameter). Three replicates of flasks of both medium were incubated in static condition in BOD incubator and shaking condition in a rotator shaker at 150 rev/min at 30°C.

The cultures were harvested at the 5th, 10th, 15th and 20th day of incubation. Each sample was centrifuged (10,000 x g for 10 min) at 4°C. The supernatant of liquid culture was used for enzyme assay. The enzymatic reactions were carried out in triplicate and determined using a double beam systronics UV/VIS spectrophotometer. All the enzyme activities were measured at room temperature (20 ± 2°C). The enzymatic activity was expressed as international units (U) defined as the amount of enzyme required to produce 1µmol product min⁻¹ and expressed as Ul⁻¹.

Protein concentration was determined following Bradford method [3]. Protein content in the sample was determined from standard curve and the amount of protein µgml⁻¹ was calculated. Laccase activity was measured following the oxidation of ABTS (ε420, 36,000M⁻¹ cm⁻1). The assay mixture contained 100mM sodium acetate buffer pH 5 and 5mM ABTS [12]. LiP activity was measured by the oxidation of 2mM VA to veratraldehyde (ε 310, 9,300 M⁻¹c⁻¹) in 100mM sodium tartrate buffer (pH3) in the presence of 0.4 mM H₂O₂ (30%) [4]. MnP activity was determined by the production of a Mn³⁺ tartrate complex (ε 238, 6,500M⁻¹ cm⁻¹) from 0.1 mM MnSO₄ in 100mM sodium tartrate buffer pH 4.5 with 0.1 Mm H₂O₂ (30%) [6].

Statistical Analysis:-
All the experimental analysis was carried out in triplicates. The results are expressed as mean values and standard deviation (SD). The results were analyzed using one-way analysis of variance (ANOVA) followed by Tukey’s HSD Test using SAV v.9.1.3 program. Differences at P>0.05 were considered to be significant.
Results and Discussion:-
Identification:-
The samples were identified as *Helvella* sp. and *Morchella* sp. on the basis of their micro and macroscopic features (Fig.1). In Himachal Pradesh there is a huge variation in climatic conditions due to the variations in altitude (450-6500 m). The climatic condition varies from hot and sub-humid tropical (450-900m). Mushrooms have extraordinary capacity to manage climatic conditions.

![Fig.1: Natural view of (A) Helvella sp., (B) Morchella sp.](image)

Spot Test of Extracellular Enzymes:-
With the agar plate screening, selected strains of wood colonizing WRF were tested for spot test. In comparison to the growth on MEA, strains grew noticeably slower on the selected agar media. In particular ABTS had inhibitory effects on fungal growth.

Laccase Spot Test:-
The extracellular ABTS-oxidizing activity of the both fungal strains into the modified Kirk medium is showed in Fig.2. *Morchella* sp. showed high ABTS-oxidizing activity in comparison to *Auricularia polytricha* and *Helvella* sp. The absence of extracellular ABTS-oxidizing activity does not necessarily imply the lack of capacity to produce these oxidative enzymes but could reflect a possible inhibition of their expression; the oxidative enzyme system is not homogeneous; its production and properties depend on the conditions and culture media. Fungal strains oxidized ABTS to the dark green ABTS cation radicals (ABTS) indicating the production of extracellular oxidoreductases [10]. It gave positive reaction immediately after inoculation, formed dark green zone around the mycelial bit and *Auricularia polytricha* and *Helvella* sp. showed green colour after 5 days of inoculation. The green colour preceded the fungal mycelium clearly demonstrating that extracellular mechanisms were responsible for the oxidation [10].

MnP Spot Test:-
All the three selected species *Morchella* sp., *Auricularia polytricha* and *Helvella* sp. showed positive reaction of extracellular MnP and formed brownish colour in medium plate (Fig. 2). *Helvella* sp. and *Morchella* sp. showed dark brown coloration in comparison to *Auricularia polytricha* however mycelial growth was noticeably slow in *Morchella* sp with MnCl₂. MnP plates were evaluated after 3 weeks of incubation for the formation of brownish flecks of manganese oxide (MnO₂) caused by the action of MnP. MnP was thought to play a crucial role during the primary attack on lignin, because it generates highly reactive Mn³⁺ which acts as a low molecular mass redox mediator and forms water soluble lignin fragments. Moreover, there are indications that MnP is even capable of mineralizing lignin up to carbon dioxide [10]. It was assumed that MnP plays an important role in mineralization of lignin.
Fig.2: Extracellular Laccase and MnP activity on agar plate; A- *Auricularia polytricha*, B- *Halvella* sp., C-*Morchella* sp.

**Extracellular Quantitative Protein Determination:**

The Bradford assay is a protein determination method that involves the binding of Coomassie brilliant blue G-250 dye to proteins [3]. When the dye binds to protein, it is converted to a stable un-protonated blue form [23]. This is anionic form of dye. In *Auricularia polytricha* maximum protein concentration was observed on 15th day in NRM and minimum concentration was observed on 5th day NPM (3.57 µg/ml). In *Halvella* sp. maximum protein concentration was observed on 15th day in NPM (16.0 µg/ml) and minimum concentration was observed on 20th day in NPM (2.89 µg/ml). In *Morchella* sp. maximum protein concentration was measured on 15th day in NPM (14.53 µg/ml) and minimum activity was observed on 10th day in NPM (2.3 µg/ml). In all three selected varieties protein concentration was higher on 15th day in both medium.

**Extracellular Ligninolytic enzyme assay:**

**Laccase:** Laccase was first detected in the Japanese lac tree *Toxicodendron verniciflua*. Later, it was found in certain other plants and fungi, but is also found in molds, black yeasts and some bacteria [17]. Laccase has been identified as one of the enzyme that plays a major role in lignin degradation. Laccase only attacks phenolic subunits of lignin, but its substrate range can be extended to non-phenolic subunits by the inclusion of a mediator [17]. In *Auricularia polytricha* maximum laccase activity was observed on 10th day in NRM (63.54UL⁻¹) and minimum was observed on 20th day in NRM (47.21UL⁻¹). In *Halvella* sp. maximum laccase activity was observed on 10th day in NPM (73.17UL⁻¹) and minimum was observed on 20th day in NPM (43.86UL⁻¹) respectively. In *Morchella* sp. maximum laccase activity was observed on 10th day in NRM (67.3UL⁻¹) and minimum was observed in NRM on 20th day (48.63UL⁻¹). It was witnessed that supplementation and incubation conditions also affected the enzymatic
activity. A few WRF gave good laccase activity under high nitrogen condition and medium supplemented with Cu but in the absence of Cu activity was not found and under low nitrogen like *Volvariella volvacea, Piloderma byssinum.* [5]

**AAO:**
The lignin degradation system of WRF is mainly composed of laccase, LiP and MnP. It is considered that, these lignin degrading enzymes do not functions but mutually interact with each other as well as with other oxidases, such as AAO. AAO activity was described for the first time in the fungus *Polystictus versicolor* (a synonym of *Trametes versicolor*) in 1960 [9]. Since then, AAO has been detected and characterized in other white-rot basidiomycetes including *Pleurotus* species, *Bjerkandera adusta* and some ascomycetous fungi [14]. AAO is the main oxidase enzyme is all selected varities. In *Auricularia polytricha* maximum AAO activity was observed on 10th day in NRM (546.4 UL⁻¹) and minimum was observed on 20th day in NPM (395.75UL⁻¹). In *Helvella* maximum AAO activity was observed on 10th day in NPM (509 UL⁻¹); and minimum was observed on 20th day in NRM (369.2UL⁻¹). In *Morchella* sp. maximum AAO activity was observed on 5th day in NRM (550.42UL⁻¹) and minimum activity was observed on 10th day in NPM (468.5UL⁻¹).

**LiP:**
One of the best known ligninolytic enzymes is LiP, which was discovered a little earlier than MnP [15]. LiP is a glycoprotein that contains one mole of iron protoporphyrin IX as a prosthetic group. LiP catalyzes the oxidation of non-phenolic aromatic compounds like veratryl alcohol. In *Auricularia polytricha* maximum LiP activity was observed on 15th day in NRM (44.56UL⁻¹) and minimum activity was observed on 5th day in NRM (1.0 UL⁻¹). In *Helvella* sp. maximum LiP activity was observed on 15th day in NPM (51.38UL⁻¹) and minimum activity was observed on 5th day under static conditions (1.0UL⁻¹). In *Morchella* sp. maximum LiP activity was observed on 15th day in NPM (55.29 L⁻¹) and minimum activity was observed on 20th day (4.29UL⁻¹).

**MnP:**
MnP is a second group of extracellular enzymes secreted by WRF. MnP operate by oxidizing Mn²⁺ to chelated Mn³⁺, which acts as a diffusible oxidant at locations remote from the enzyme active site. MnP is widely distributed in WRF, including *P. chrysosporium, P. radiata, Nematoloma frowardi, P. eryngii* and *B. adusta* [14] and this peroxidase described for the first time in *B. adusta* [14]. In *Auricularia polytricha* maximum MnP activity was observed on 15th day in NPM (46.4UL⁻¹) and minimum activity was observed on 5th day in NPM (1.04 UL⁻¹). In *Helvella* sp. maximum MnP activity was observed on 15th day in NPM and minimum was observed on 5th day in NRM (2.33UL⁻¹). In *Morchella* sp. maximum MnP production was observed on 15th day in NRM (57.32UL⁻¹) and minimum activity was observed on 20th day in NRM (9.73UL⁻¹).

Mushrooms are recognized as important food items since ancient times. Their usage is being increased day by day for their significant role in human health, nutrition and disease. Fungi are ideal food because they have a fairly high content of protein (typically 20-30% dry matter as crude protein) which contains all of the essential amino acids [17]. For carrying out our study, we first selected three species two wild that is *Morchella* sp. and *Helvella* sp. and one commercially cultivated *Auricularia polytricha*. The extracellular laccase, AAO, LiP and MnP are thought to be involved in lignin degradation by WRF, which produce different combination of these extracellular enzymes. Proteins, Laccase and MnP production was maximum in *Helvella* sp. AAO (Aryl Alcohol Oxidase) production was maximum in *Auricularia polytricha* however *Morchella* sp. showed high LiP activity. In WRF laccase besides functioning as lignin degrading enzyme, it was also important in pigment production, polyphenol detoxification, fruity body formation, sporulation and antimicrobial agent [7]. In general Laccase and MnP are more widely distributed among WRF than LiP [20]. Production of enzymes was good in *Morchella* and *Helvella* sp. in comparison to *Auricularia polytricha* (Table-1).

**Table 1:** Extracellular protein (µg/ml) and ligninolytic enzymes (U l⁻¹)

| S.No | Day interval | *Auricularia polytricha* | *Helvella* sp. | *Morchella* sp. |
|------|--------------|--------------------------|----------------|----------------|
|      |              | NRM | NPM | NRM | NPM | NRM | NPM |
| Protein | 5 | 3.57±0.2a | 4.46±0.11ac | 1.8±0.48b | 6.92±1.25a | 2.30±1bc | 5.42±0.2b |
|        | 10 | 7.13±12a | 8.41±0.31cd | 6.2±0.5b | 11.2±0.5b | 13.1±0.3b | 2.3±0.4c |
|        | 15 | 15.43±0.43a | 12.15±0.1bc | 10.43±0.22b | 16±0.5b | 11.36±1b | 14.53±1.6a |
|        | 20 | 8.21±0.67a | 6.26±0.63cd | 6.73±0.26b | 2.89±0.61d | 5.1±1e | 3.95±0.77cd |
| Laccase | 5 | 58.1±0.67c | 52.6±0.49cd | 64.53±0.73b | 69.72±0.51a | 55.6±0.7cd | 64.35±0.3b |
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