

Cellulase Production by *Curvularia pallescens* isolated from textile effluent

Abstract: Make in format as aim, location and duration of study, design of study, result and interpretation; leave the space before mentioning the units

Effects of pH, temperature, incubation time, source of carbon and nitrogen were tested in submerged fermentation process in production of cellulose by *Curvularia pallescens* isolated from textile effluent. The production medium was prepared in distilled water, supplemented with 4.5% wheat bran, 0.05% KCl, 0.2% KH₂PO₄, (carbon source), yeast extract (nitrogen source), maintained with pH of 5.5 and incubated at 28⁰C for 120h was found optimal for production of cellulose.

Keywords: Cellulase, *Curvularia pallescens*, textile effluent, submerged fermentation, wheat bran

INTRODUCTION: number the references and put in square bracket sequentially

Cellulases are important industrial enzymes and find application in several industrial processes (Kang *et. al.*, 2004). Currently the most important application is the bio-bleaching of pulp, the production of dissolving pulp, the treatment of wastewater. The cost of production and low yields of these enzymes are the major problems for industrial application. Therefore, investigations on the ability of the cellulose hydrolyzing microbial strains to utilize inexpensive substrate have been done (Kang *et. al.*, 2004; Haltrich *et. al.*, 1996). The enzyme is commercially used after extracting from many microorganisms especially fungal source (Hanif *et. al.*, 2004; Kang *et. al.*, 2004) of mostly terrestrial origin but less from marine sources.

Therefore, in the present study, the enzyme was attempted in a fungus, *Curvularia pallescens* isolated from textile effluent for maximizing its production under optimal conditions in submerged fermentation by using inexpensive substrate wheat bran.

MATERIALS AND METHODS: number the references and put in square bracket

Sequential; leave the space before mentioning the units

Organism and culture condition

Curvularia pallescens was isolated from textile effluent using serial dilution and spread plate method (Graca *et. al.*, 1997).

All the enzyme production studies were carried out under submerged conditions in the media contained Wheat bran 4.5%, yeast extract 1.5%, glucose 1%, NH₄Cl 0.25%, Thiamine dichloride

38 0.05%, KH_2PO_4 0.2%, $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ 0.05%, CaCl_2 0.01%, KCl 0.05%.. 10 agar plugs of 8mm
39 diameter of the fungus grown for 7 days on PDA culture plates were inoculated in 100ml of the
40 medium. The flasks were incubated at 28°C under shaker conditions at 120 rpm. Cultures were
41 harvested on 5th day and assayed for cellulase activity.
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43 **Optimization of the medium**

44 Standardization of the optimum condition for the growth of the isolated organism as well as for
45 cellulase production was determined by varying temperature and pH of the specially designed
46 media, carbon and nitrogen sources, inoculum size, incubation period, mechanical shaking with
47 different speed during incubation.
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49 **Cellulase assay**

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51 The test fungus was assayed for total cellulolytic activity by filter paper assay (FPA) (Mandel *et al.*, 1976); endoglucanase (Cx) activity by carboxymethyl assay (CMC), cellobiohydrolase (C1) activity by cotton assay and β -glucosidase activity by using p-nitrophenyl- β -D-pyranoglucosidase (PNPG) method (Rosenberg *et al.*, 1975).
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56 1 unit of FPA , CMCase and cotton activity was defined as the amount of enzyme that releases 1
57 micromole of glucose from the substrate per minute and 1 unit of β -glucosidase was defined as
58 the amount of enzyme required to liberate 1 micromole of 4-nitrophenol per minute.
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61 **RESULTS : Add discussion along with result; leave the space before mentioning of units;**

62 **give the sketch of fungus**

63 Glucose favoured cellobiohydrolase and endoglucanase activity in *C. pallescens* (30.35 and
64 21.24U/ml respectively) where as sucrose and fructose proved to be best for FPA (61.35 U/ml)
65 activity and β -glucosidase (6.97 U/ml) activity respectively (figure-1).

66 Organic nitrogen sources used for optimization were peptone, malt and yeast. *C. pallescens*
67 showed maximum FPA activity 38.59 U/ml, cellobiohydrolase activity 30.35 U/ml, β -
68 glucosidase activity 3.08 U/ml in the presence of yeast where as endoglucanase activity 33.71
69 U/ml reported higher with malt extract (figure-2).

70 $(\text{NH}_4)_2\text{SO}_4$ was reported as best inorganic nitrogen source for cellobiohydrolase, endoglucanase
71 and β -glucosidase activities at 57.18, 56.82 and 6.77 U/ml repectively. FPA activity was shown
72 highest at 87.59 U/ml with NaNO_3 (figure-3).

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74 The test fungus achieved maximum FPA activity followed by cellobiohydrolase, endoglucanase
75 and β -glucosidase activity at 46.76, 42.06, 26.94 and 3.56 U/ml respectively at pH 5.5 (figure-
76 4). Temperature of 28⁰C produced maximum cellulase activity. Highest activity recorded was of
77 FPA (38.94 U/ml), followed by cellobiohydrolase (30.29 U/ml), endoglucanase (22.41 U/ml),
78 and β -glucosidase (3.98 U/ml) (figure-5). FPA activity 38.65 respectively was obtained
79 maximum for *C. pallescens* after 168 hrs whereas cellobiohydrolase, endoglucanase and β -
80 glucosidase activities 40.29, 57.41 and 2.98 U/ml respectively were recorded highest at 120 hrs
81 of incubation (figure-6).

82 Media containing various amounts of inoculi were used for studying the effect of inoculum size
83 on lignocellulolytic activity. Results are shown in figure-7. Reported maximum FPA,
84 Cellobiohydrolase, endoglucanase and β -glucosidase activities 37.94, 30.01, 22.24 and 3.98
85 U/ml by inoculation 10 disc of 8mm size in the production medium. *C. pallescens* also gave
86 maximum cellulase production at 120rpm. Endoglucanase activity was observed highest
87 followed by FPA, cellobiohydrolase and β -glucosidase activities as 38.59, 30.35, 27.41 and 1.91
88 U/ml respectively (figure-8).

89 **DISCUSSION: Add this section to result at relevant places; references mentioned in this**

90 **section to be numbered sequentially in square bracket**

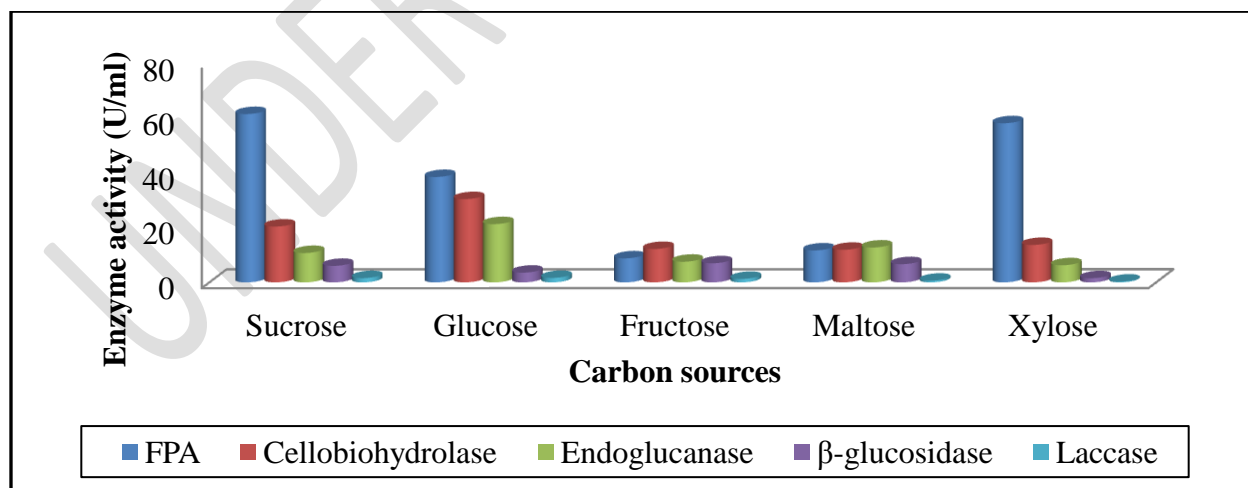
91 The media optimization is an important aspect to be considered in the development of
92 fermentation technology. The production of primary metabolites by microorganisms is highly
93 influenced by their growth, which is determined by the availability of the nutrients in the
94 substrates. Garcia *et al.*, (2002) reported that submerged fermentation for aerobic
95 microorganisms is well known and widely used method for the production of cellulase and
96 xylanase. Chellapandi and Jani (2009) reported enhanced endoglucanase production by soil
97 isolates of *Fusarium* sp. and *Aspergillus* sp. through submerged fermentation process. Papinutti
98 and Lechner (2008) studied influence of the carbon source on the growth and lignocellulolytic
99 enzyme production by *Morchella esculenta*. Arora and Sehgal (2010) reported production of
100 cellulase and xylanase by *Scopulariopsis acremonium* through submerged fermentation using
101 shake flask cultivation media. The effect of process parameters such as effect of temperature, pH

102 and inoculum size was investigated. Maximum cellulase and xylanase having an enzyme activity
103 of 694.45 and 931.25 IU, respectively, were produced at 30°C incubation temperature. The pH
104 optimum to achieve these enzyme activities was 5.5 with an inoculum size of 1×10^5 spores ml⁻¹
105 of tween – 80.

106 Gupta *et al.*, (1990) studied microbial proteins and cellulase production from cellulosic
107 materials by *Coprinus atramentarius* and reported the optimum pH for protein production and
108 extracellular enzymes (cellulase and xylanase) by *C. atramentarius*, utilizing cellulose to be 6
109 and optimum temperature 30°C. The resulting enzyme activities were endoxylanase as 7.2 IU ml⁻¹,
110 exoglucanase as 1.0 IU ml⁻¹ and xylanase as 5 IU ml⁻¹. Li *et al.*, (2006) reported pH of 4.14
111 was reported to be optimum for the production of endoxylanase production by *Aspergillus*
112 *awamori* under submerged fermentation which gave an enzyme activity of 28.25 U ml⁻¹.

113 Shear stress within the medium, which is directly related to the stirrer speed, has also been
114 shown to have a marked influence on enzyme production by *Thermomyces lanuginosus* SSBP
115 (Reddy *et al.*, 2002; Singh *et al.*, 2000). Acharya *et al.*, (2008) reported maximum cellulase
116 production by *Aspergillus niger* in submerged fermentation at 120 rpm. However Ojumu *et al.*,
117 (2003) observed maximum cellulase production by *Aspergillus flavus* Linn isolate NSPR 101 at
118 agitation of rate 200 rpm.

119 **Figure-1: Optimization of carbon source for lignocellulases production by *Curvularia pallescens***



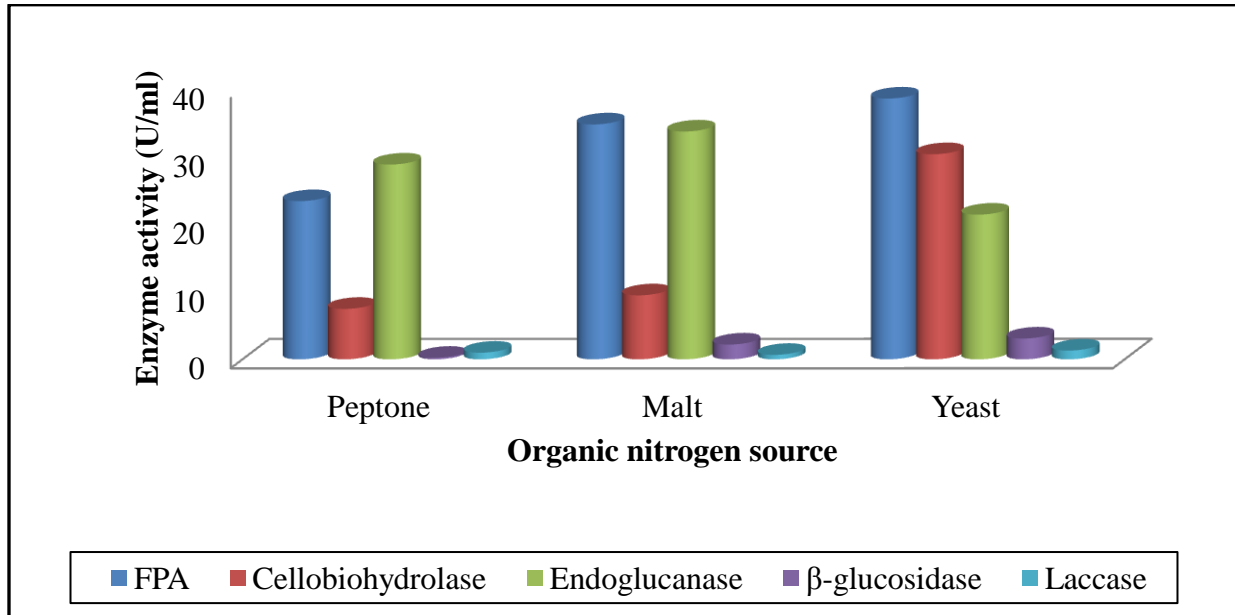
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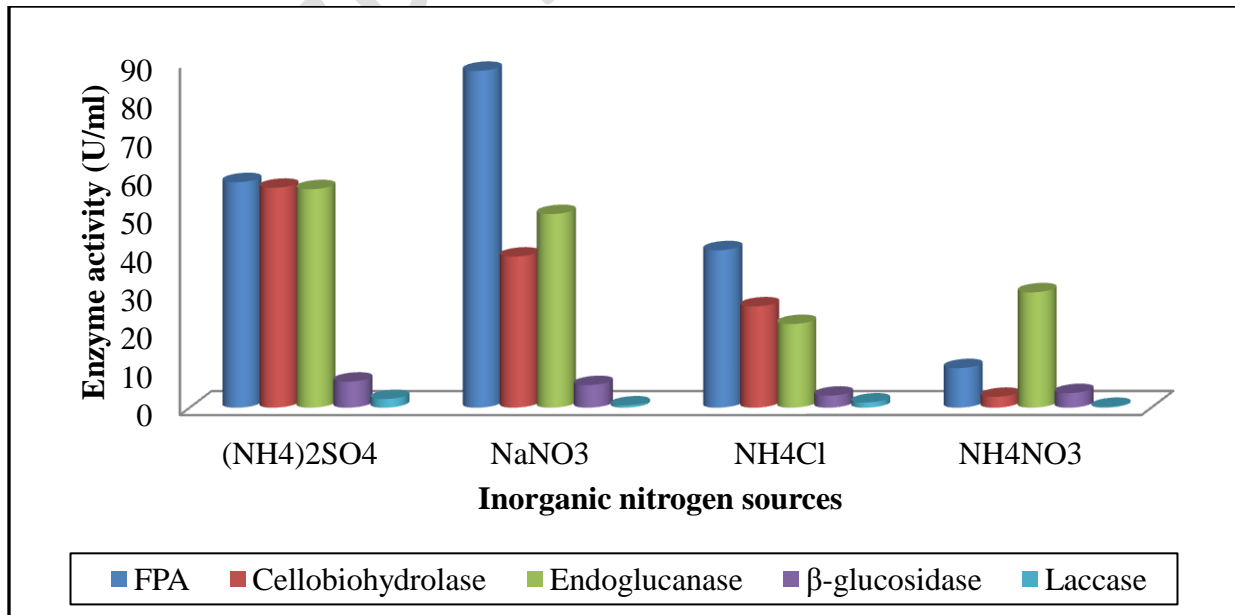
124 **Figure-2: Optimization of nitrogen source (organic) for lignocellulases production by *Curvularia***
125 ***pallescens***



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128 **Figure-3: Optimization of nitrogen source (inorganic) for lignocellulases production by *Curvularia***
129 ***pallescens***

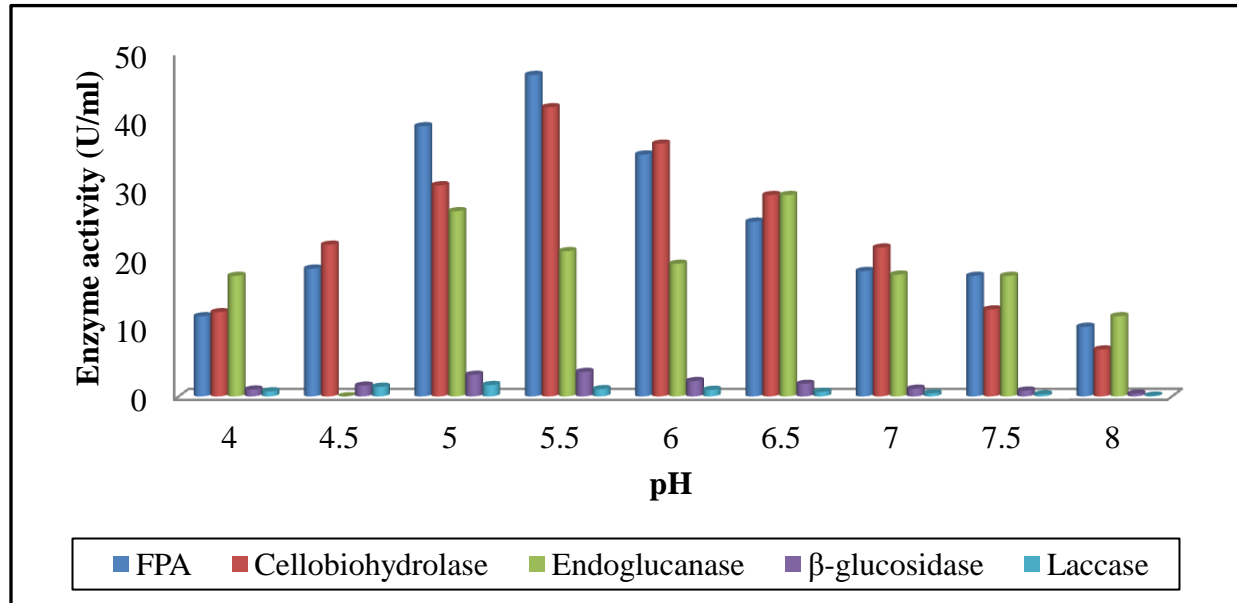


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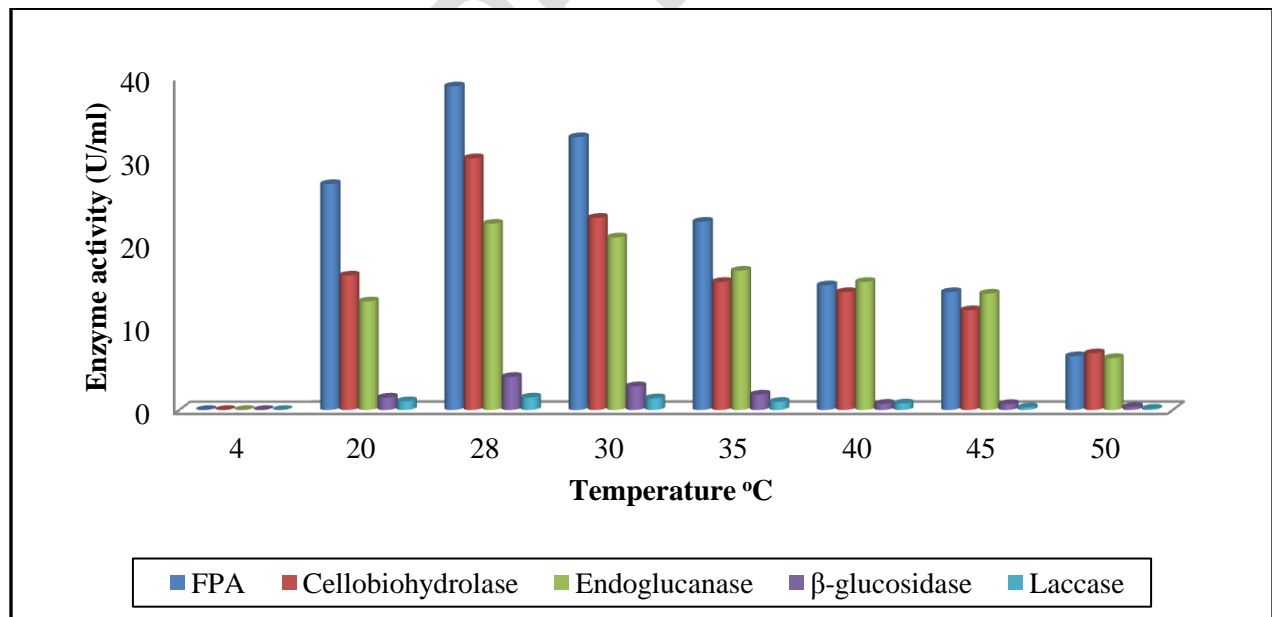
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133 **Figure-4: Optimization of pH for lignocellulases production by *Curvularia pallescens***



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135 **Figure-5: Optimization of temperature for lignocellulases production by *Curvularia pallescens***



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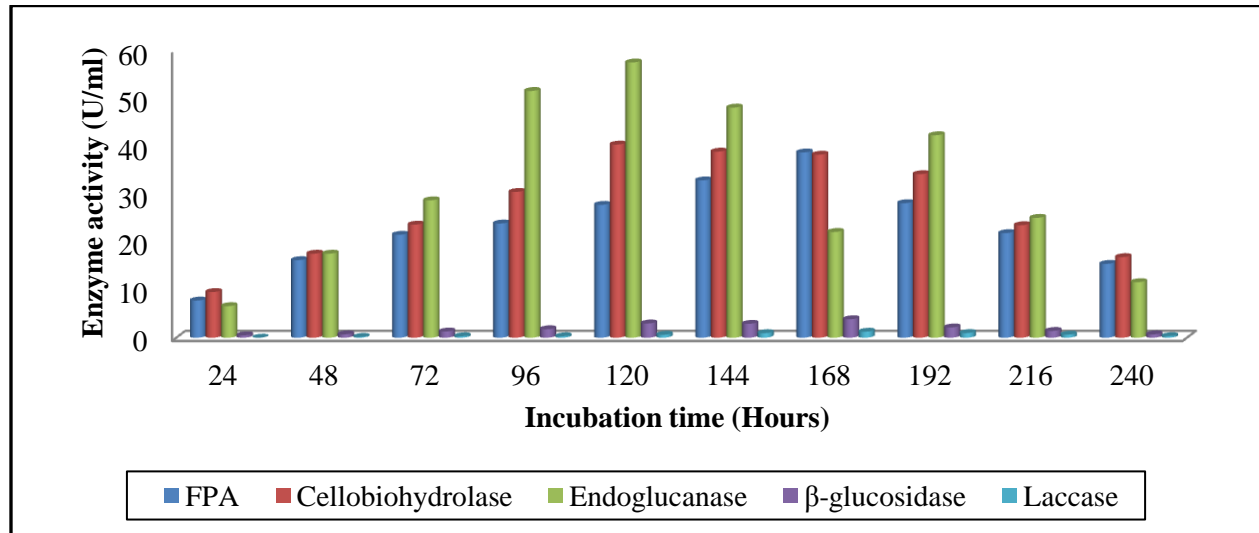
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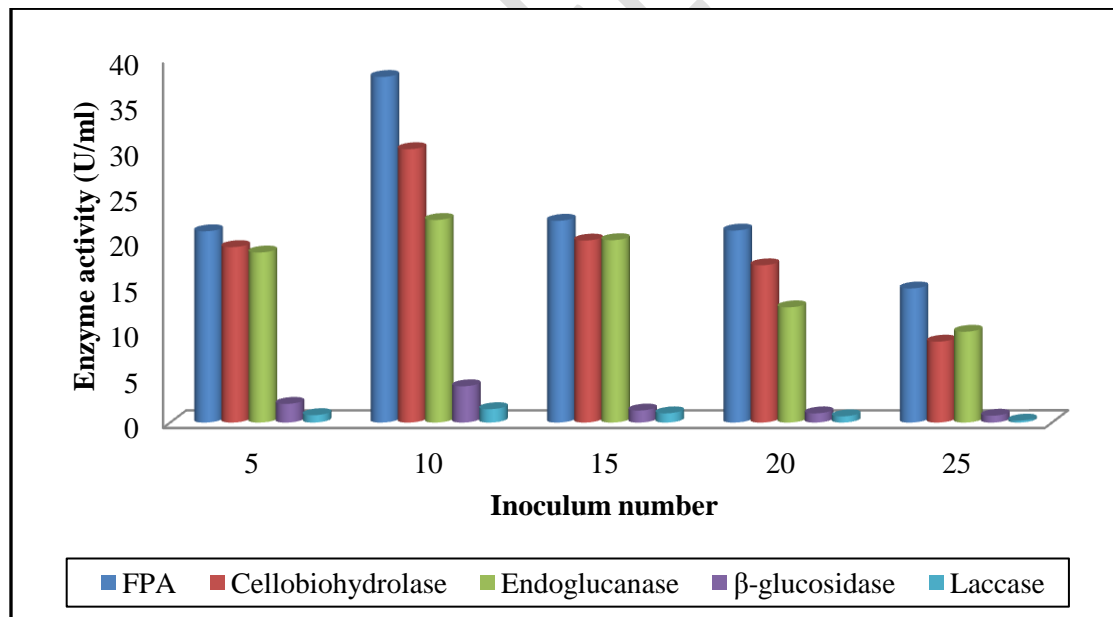
141 **Figure-6: Optimization of incubation time for lignocellulases production by *Curvularia pallescens***



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144 **Figure-7: Optimization of inoculum size for lignocellulases production by *Curvularia pallescens***



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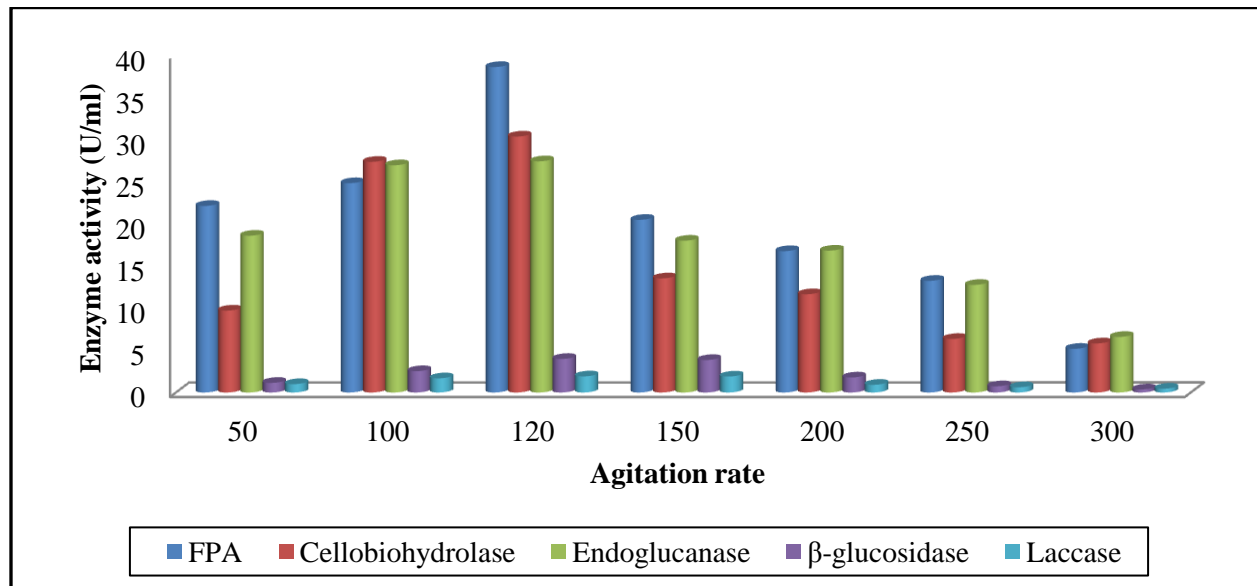
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151 **Figure-8: Optimization of agitation rate for lignocellulases production by *Curvularia pallescens***



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153 **CONCLUSION: Add a paragraph**

154 **REFERENCES: Present sequentially as numbers from introduction, method and**
155 **discussion**

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