Mutagenesis and screening of *Penicillium oxalicum* (Dal 5) for enhancing cellulase production

Ankita Singh¹, B. S. Chadha²*

¹Senior Resident, ²Professor, Dept. of Microbiology, Pandit Jawahar Lal Nehru Government Medical College & Hospital, Chamba, Himachal Pradesh, ³Guru Nanak Dev University, Amritsar, Punjab, India

*Corresponding Author:*
Email: singhankitahp5@gmail.com

**Abstract**

In the modern era due to increasing fuel demand, prices of crude oil is increasing day by day. So the need for bioenergy sources is increasing in coming years. Of all the biomass present in the world lignocelluloses as industrial, forest and agricultural residues constitutes the major portion. The present study was focused to determine potential of fungus *Penicillium oxalicum* (Dal 5) for cellulase production on different commonly available lignocellulosic substrates (Rice straw, Wheat straw, Corn cob, Baggase, Sorghum straw), organic nitrogen source (Corn steep liquor).

**Keywords:** Bioenergy, Buffer, Cellulase, Enzyme, Substrate.

**Introduction**

Rising prices of crude oil is increasing day by day due to increase in fuel demand so the alternative source of bioenergy is increasing sharply. So lignocelluloses is identified as the prime source of biofuel and other valuable products. Lignocelluloses is agricultural, industrial and forest contributes to the majority of biomass in the world (Howard et al., 2003). For the production of industrial important products from lignocellulosic biomass, bioconversion of cellulose and hemicellulolytic components is necessary into fermentable sugars. So variety of microorganisms bacteria and fungi has the ability to degrade the lignocellulosic biomass to glucose and xylose monomers. Some cellulases exist as discrete multienzyme complexes, called cellulosomes which consists of multiple subunits cellulolytic enzymes on the other hand of filamentous fungi is non-complexed that consists of exoglucanases or cellobiohydrolases (CBH I and CBH II). Enzymes synergistically act to catalyze the hydrolysis of cellulose (Wen et al., 2005). Cellulose units are interlinked by β -1, 4-glycosidic bonds is a liner polymer of D-glucose units, which is renewable biomass available on earth.

1. Endoglucanase (EC 3.2.1.74) which attack cellulose in amorphous zone and release oligomers.
2. Cellulobiohydrolase (EC 3.2.1.91) that liberate cellulobiase from reducing and non-reducing ends.
3. β-Gucosidase (EC 3.2.1.21) which hydrolyse cellolobiase to glucose.

Cellulose In plant cell walls, is entangled with and shielded by hemicellulose, is group of complex polysaccharides composed of different glyco-units and glycosidic bonds. Degradation of hemicellulose, converts hemicellulose into valuable saccharides, is done by interdependent and synergistic hemicellulases. Xylan, β-glucan, polygalacturonan, xylrogucan, arabinoxylan, mannan, galactan, galactomannan, arabinan, etc are common hemicelluloses, they are targets of xylanase, β-glucanase, polygalacturonase, xylrogucanase, mannanase, arabinase, galactanase, glucuronidase, acetyl xylan esterase, and other enzymes (Wen et al., 2005).

Temperature, pH, adsorption are physical parameters, nitrogen, phosphorus, chemical factors phenolic compounds and some inhibitors can inhibit bioconversion of lignocellulose. Cellulases production by microbial cells is done by genetic and biochemical which also includes catabolite repression, induction, or end product inhibition (Cheng et al., 2009). Many efforts made to increase the production of cellulases through strain improvement by mutagenesis (Kaur et al., 2012). Mutagenized and genetically modified organism produces high levels of cellulases (Mandel and Andreotti 1978; Nevalainen, 1980; Durand, 1988; Szengyel, 2000). The objective of these efforts has been to obtain high levels of cellulases in order to degrade waste cellulose. However, despite the effort of many laboratories, only few commercially efficient enzyme complexes have been produced. To resolve problems, of high cost of enzymes production many researches are done in isolating new microorganisms for production of higher specific activity and greater efficiency (Stahlberg 1991). The present study was focussed to determine potential of fungus *Penicillium oxalicum* (Dal 5) for cellulase substrates of lignocellulosic (Rice straw, Baggage, Wheat straw, Sorghum straw, Corn cob), organic nitrogen source (Corn steep liquor).

**Materials and Methods**

2.1. Microorganisms and culture media

*Penicillium oxalicum* (Dal 5) wild type strain and mutants were maintained on YpSS medium of following composition.
2.2 Slide culturing for morphological/microscopic examination

A sheet of sterile filter paper along with U-shaped glass paper rods were placed at the bottom of petridish. The filter paper was moistened with sterile distilled water. A glass slide was placed on the glass rods along with cover slips. The entire assembly was autoclaved. A 5 mm thin agar block (YpSS) was placed on the slides aseptically using forceps. The block was inoculated with spores of *P. oxalicum* and cover slip was placed over it. After that the assembly was incubated at 30°C for 48 h. Then the slide was observed under microscope (10X) and (40X) to examine mycelial growth as well as sporulation.

2.3 Cultures are screened for production of hemicellulase/cellulase different organic nitrogen sources and lignocelluloses sources

Cellulase/hemicellulase production is studied by shake flask culture of *P. oxalicum*. The cellulase production medium 50 ml in 250 ml Erlenmeyer flasks contained (% w/v) of -

- Cellulose powder: 3%
- Rice straw: 1%
- Wheat bran: 1%
- KH₂PO₄: 2.4%
- (NH₄)₂SO₄: 0.5%
- MgSO₄: 0.12%
- Tween 80: 0.1%
- Trace elements
- CuSO₄: 100mg/10 ml
- MnSO₄: 100 mg/10 ml
- ZnSO₄: 100 mg/ml

Filter sterilized 0.4% urea was added after autoclaving.

To study the effect of different carbon sources, rice straw in the medium was replaced with either of sorghum straw, baggase, wheat bran, wheat straw or corn cob. Furthermore the effect of adding CSL in place of (NH₄)₂SO₄ in enzyme production was also studied.

Production medium was inoculated with 4 stubs of actively growing *P. oxalicum* culture taken from the petriplate and incubated for 7 days at 30°C. After 7 days contents of flasks were centrifuged at 10,000 rpm for 10 min. Then supernatent was taken and pellet was discarded. The enzyme extract was suitably diluted (2, 5, 10, 20, 50 and 100 X) using sodium acetate buffer (50mM, pH 5.0).

<table>
<thead>
<tr>
<th>YpSS (pH 6.0)</th>
<th>Composition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soluble starch</td>
<td>1.5</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>0.4</td>
</tr>
<tr>
<td>KH₂PO₄</td>
<td>0.2</td>
</tr>
<tr>
<td>K₂HPO₄</td>
<td>0.23</td>
</tr>
<tr>
<td>MgSO₄</td>
<td>0.05</td>
</tr>
<tr>
<td>Citric acid</td>
<td>0.057</td>
</tr>
</tbody>
</table>

### 2.4 Estimation of enzyme activities on different carbon sources and organic nitrogen sources:

1. Activity of β - Glucosidase:

   **Substrate:** In 50 mM sodium acetate buffer of pH 5.0, 4 nitrophenyl β-D glucopyranoside (pNPG) (3mM) is prepared.

   The enzyme assay was performed using micro titre plate method. The reagent blank contained buffer (100µl) whereas the substrate blank contained pNPG (25µl) and buffer (75µl). The enzyme blank was prepared by adding buffer (75µl) and diluted enzyme (25µl). The reaction mixture contained buffer (50µl), suitably diluted enzyme (25µl), and substrate (25µl). The ELISA plate was wrapped in foil and incubated 50°C for 30 min. The reaction was stopped by adding NaOH glycine buffer 100µl. The developed yellow color was read using microplate reader at 405nm (BIORAD plate reader). The amount of p-nitrophenol released was quantified using the pNP standard. The amount of enzyme required to release 1μl of pNP per min The β-glucosidase activity (units/ml) and was expressed under assay conditions.

2. Cellobiosidase activity (CBH I+ CBH II)

   **Substrate:** 4 nitrophenyl β-D-cellobioside prepared in sodium acetate buffer pH 5.0.

   The enzyme assay was performed using micro titre plate method. The reagent blank contained buffer (100µl) whereas, the substrate blank contained pNPG (25µl) and buffer (75µl). The enzyme blank was prepared by adding buffer (75µl) and diluted enzyme (25µl). The reaction mixture contained buffer (50µl), suitably diluted enzyme (25µl), and substrate (25µl). The ELISA plate was wrapped in foil and is incubated for 30 min at 50 degree temperature. NaOH glycine buffer 100µl is added to stop the reaction. The developed yellow color was read using microplate reader at 405nm (BIORAD plate reader). The amount of p-nitrophenol released was quantified using the pNP standard. The amount of enzyme required to release 1μl of pNP per min under assay conditions.

3. Cellbiohydrodase activity

   **Substrate:** 4 nitrophenyl β-D-lactopyranoside (pNPL 3mM) prepared in sodium acetate buffer pH 5.0. The enzyme assay was performed using micro titre plate method. Buffer (100µl) is reagent blank, pNPL (25µl) and buffer (75µl) is substrate blank. The enzyme blank contained buffer (75µl) and enzyme (25µl). The reaction mixture contained buffer (50µl), enzyme (25µl) and substrate (25µl). The ELISA plate was wrapped in foil and incubated 50°C for 1h. The reaction was stopped by adding NaOH glycine buffer 100µl. The developed yellow color was read using microplate reader at 405nm (BIORAD plate reader). The
amount of p-nitrophenol released was quantified using the pNP standard. The cellobiohydrolase activity amount of enzyme required to release 1µl of pNP per min under assay conditions

4. β- Xylosidase activity

**Substrate**: 4 nitrophenyl β-D-Xyloside prepared in sodium acetate buffer pH 5.0.

The enzyme assay was performed using micro titre plate method. The reagent blank contained buffer (100µl) whereas the substrate blank contained pNPG (25µl) and buffer (75µl). The enzyme blank was prepared by adding buffer (75µl) and diluted enzyme (25µl). The reaction mixture contained buffer (50µl), suitably diluted enzyme (25µl), and substrate (25µl). The ELISA plate was wrapped in foil and incubated 50°C for 10 min at 50°C. Stop reaction by adding NaOH glycine buffer 100µl. The developed yellow color was read using microplate reader at 405nm (BIORAD plate reader). The amount of p-nitrophenol released was quantified using the pNP standard. The xyloside activity was expressed as the amount of enzyme required to release 1µl of pNP per min under assay conditions.

5. Arabinofuranosidase activity

**Substrate**: 4 nitrophenyl α-L-Arabinofuranoside prepared in acetate buffer pH 5.5.

The enzyme assay was performed using micro titre plate method. The reagent blank contained buffer (100µl) whereas, the substrate blank contained pNPG (25µl) and buffer (75µl). The enzyme blank was prepared by adding buffer (75µl) and diluted enzyme (25µl). The reaction mixture contained buffer (50µl) suitably diluted enzyme (25µl) and substrate (25µl). The ELISA plate was wrapped in foil and incubated 50°C for 30 min. The reaction was stopped by adding NaOH glycine buffer 100µl. The developed yellow color was read using microplate reader at 405nm (BIORAD plate reader). The amount of p-nitrophenol released was quantified using the pNP standard. The arabinofuranoside activity was expressed as the amount of enzyme required to release 1µl of pNP per min under assay conditions.

6. Endoglucanase activity (EG)

**Substrate**: 1% CMC prepared in sodium citrate buffer.

CMC substrate measures The endoglucanase activity. The reaction was initiated by incubating 500µl of substrate (1% CMC prepared in pH 5.0, 50mM acetate buffer) and diluted enzyme of 500µl. Suitable enzyme blank that contained 500µl of enzyme and 500µl of buffer while substrate blank that contained 500µl of substrate and 500µl of buffer were also included. The reaction was carried out for 10 min at 50°C. 3ml of dinitrosalicylic acid (DNS) is used for stopping the reaction followed by boiling at 100°C for 10 min. The developed colour was read at 540nm using NOVO spec II spectrophotometer .The amount of released sugars was quantified using glucose standard. Endoglucanase (CM Case) activity of one unit defined as 1 µmol of glucose equivalent liberated per unit under assay condition.

7. Fpase activity

**Substrate**: Whatman No. 1 filter paper.

The reaction mixture contained (500µl) buffer (500µl) suitably diluted enzyme and one properly folded filter paper strip. The blank contained buffer 1000µl whereas the substrate blank contained sodium citrate 1000µl buffer and paper strip (1* 6mm) properly folded. Incubated at 50°C for 60 min. 3ml of dinitrosalicylic acid (DNS) was added to stop the reaction followed by boiling at 100°C for 10 min. The developed color was read at 540nm using NOVO spec II spectrophotometer. The amount released sugars was estimated using glucose standard.

8. Xylanase activity

**Substrate**: Sodium citrate buffer is prepared in 1% Birchwood xylan. Mixture of reaction contains (500µl) substrate and (500µl) dilute enzyme. Buffer contains reagent blank 1000µl whereas the substrate blank contained (500µl) xylan and (500µl) buffer. The enzyme blank was constituted by adding (500µl) enzyme and 500µl buffer this was then incubated for 10 min at 50°C temperature. The reaction was stopped by adding 3ml of dinitrosalicylic acid (DNS) followed by boiling at 100°C for 10 min. The developed color was read at 540nm using NOVO spec II spectrophotometer. The amount of released sugars was quantified using glucose standard. One unit of xylanase activity was defined as 1µl mol of glucose equivalent liberated per unit under assay condition.

**Results and Discussion**

3.1 Morphological and microscopic characterization

The culture up on grown on YpSS agar produced sporulating colonies which were velvety in texture and blue-green in color. Light microscopy revealed elliptical conidia produced on smooth-walled conidiophores. Microscopic view also showed branched conidiophores bearing phialides in clusters (brush like structure) which had chains of conidia. The conidiophores showed divaricate branching pattern (Fig. 4).

3.2 Carbon sources effects on cellulase/hemicellulase production

The results in Table 1 shows that cellulbiohydrodrolase activity (CBH) as measured against pNPL was maximally produced (0.34 units/ml) when rice straw,
sorghum straw or wheat straw was used as carbon source (Fig. 5). The cellobiosidase activity (Fig. 6) against pNPC which corresponds to sum total of CBH I, CBH II and EG 7 was again maximally supported by rice straw (1.4 units/ml) followed by bagasse (1.17 units/ml). Recombinant DNA technology can be used to understand molecular mechanism which can help in biodegradation of lignocelluloses.

Table 1: carbon sources effects at cellulase production by *P. oxalicum* (Dal5)

<table>
<thead>
<tr>
<th>Carbon Source</th>
<th>CBH (units/ml)</th>
<th>Cellobiosidase (units/ml)</th>
<th>β-glucosidase (units/ml)</th>
<th>CMCase (units/ml)</th>
<th>FPase (units/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rice straw</td>
<td>0.34</td>
<td>1.41</td>
<td>2.74</td>
<td>17.17</td>
<td>0.81</td>
</tr>
<tr>
<td>Bagasse</td>
<td>0.23</td>
<td>1.17</td>
<td>2.76</td>
<td>17.45</td>
<td>0.60</td>
</tr>
<tr>
<td>Corn –cob</td>
<td>0.24</td>
<td>0.80</td>
<td>1.62</td>
<td>18.50</td>
<td>0.77</td>
</tr>
<tr>
<td>Sorghum straw</td>
<td>0.34</td>
<td>0.92</td>
<td>1.37</td>
<td>19.81</td>
<td>0.73</td>
</tr>
<tr>
<td>Wheat straw</td>
<td>0.34</td>
<td>0.85</td>
<td>1.74</td>
<td>19.44</td>
<td>0.60</td>
</tr>
</tbody>
</table>

The β-glucosidase activity measured against pNPG as substrate was maximal (2.76 units/ml) when bagasse was used as carbon. Rice straw also supported high β glucosidase activity (2.74 units/ml) (Fig. 7). The Endoglucanase activity as measured against 1% CMC was produced maximally (19.8 units/ml) in presence of sorghum straw as carbon source followed by wheat straw (19.44 units/ml) (Fig. 8). The total cellulase activity as measured against Whatman No. 1 filter paper was highest (0.80 units/ml) in presence of rice straw.

The results in Table 2, shows that the xylanase activity as measured against 1% Birch wood xylan was maximal (92.50 units/ml) when wheat straw was used as carbon source whereas, in presence of corn cob the culture produced (91.21 units/ml) of xylanases (Fig. 10).

Table 2: Effect of Carbon sources on hemicellulase production

<table>
<thead>
<tr>
<th>Carbon Source</th>
<th>Xylanase (units/ml)</th>
<th>β-Xylosidase (units/ml)</th>
<th>Arabinofuranosidase (units/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Rice-straw (control)</td>
<td>61.05</td>
<td>0.07</td>
<td>0.09</td>
</tr>
<tr>
<td>2. Bagasse</td>
<td>56.76</td>
<td>0.06</td>
<td>0.20</td>
</tr>
<tr>
<td>3. Corn –cob</td>
<td>91.21</td>
<td>0.07</td>
<td>0.11</td>
</tr>
<tr>
<td>4. Sorghum straw</td>
<td>88.19</td>
<td>0.10</td>
<td>0.08</td>
</tr>
<tr>
<td>5. Wheat straw</td>
<td>92.50</td>
<td>0.08</td>
<td>0.12</td>
</tr>
</tbody>
</table>

The β-Xylosidase activity as measured against pNP X was maximally produced (0.11 units/ml) when Sorghum straw was used as carbon sources followed by (0.08 units/ml) in presence of wheat straw as carbon source in (Fig. 11). The arabinofuranosidase activity was measured against α-L arabinofuranoside was maximal (0.19 units/ml) when wheat straw was used as carbon source in (Fig. 12) when bagasse was used as carbon source followed by (0.12 units/ml).

Fig. 2: Petriplate containing *P. oxalicum* (Dal 5) culture
Fig. 3: a: Well Microtitreplate; b: P. oxalicum (Dal 5) slide culture stained with lactophenol observed under microscope (40X)

Fig. 4: a: Cellobiohydrolase activity produced by P. oxalicum (Dal 5) in presence of different carbon sources; b: Cellobiosidase activity produced by P. oxalicum (Dal 5) in presence of different carbon sources
Fig. 5: a: β-Glucosidase activity produced by P. oxalicum (Dal5) in presence of different carbon sources; b: Endoglucanase activity by P. oxalicum (Dal5) in presence of different carbon sources

Fig. 6: a: Filter paper activity by P. oxalicum (Dal5) in presence of different carbon sources; b: Xylanase activity by P. oxalicum (Dal5) in presence of different carbon sources
Conclusion

The present study was focused to determine the potential of fungus *Penicillium oxalicum* (Dal 5) for cellulase production on lignocellulosic substrates (Corn cob, Wheat straw, Rice straw, Sorghum straw, Baggase), organic nitrogen source (Corn steep liquor).

References


How to cite this article: S. Ankita, Chadha B. S. Mutagenesis and screening of *Penicillium oxalicum* (Dal 5) for enhancing cellulase production. Indian J Microbiol Res 2018;5(1):7-13