MOLECULAR CONFIRMATION, IDENTIFICATION AND INFLUENCE OF CARBON SOURCE FOR THE PRODUCTION OF XYLANASE FROM *PENICILLIUM CITRINUM*

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INTRODUCTION

Until recently, in the pulp and paper industry, the use of enzymes was not considered technically or economically feasible. Quite simply, suitable enzymes were not readily available, except for limited use in the modification of starch for paper coatings. However, research by scientific institutions and enzyme producers has led to the development of enzymes that offer significant benefits for the industry. Several commercial products have been launched successfully in the past few years.

The enzyme, xylanase are the upcoming enzyme of commercial sector and are widely used in paper and pulp industry, animal feed, textile industry, coffee and tea fermentation, oil extraction, waste paper recycling and in the fruit juice industries. This enzyme are the tools of nature that help us in providing everyday products in an environmentally conscious manner.

Xylanases are hydrodrolases depolymerising the plant cell wall component-xylan, the second most abundant polysaccharide. The Molecular structure and hydrolytic pattern of the xylanases have been reported extensively and mechanism of hydrolysis has also been proposed.

Many different microbial genera, ranging from bacteria to fungi, have been used for more than 50 years in the production of industrial enzymes. They are particularly interesting producers of xylanases and excrete much higher xylanolytic enzymes into the medium than bacteria or yeast.

Due to importance of xylanase production, we made an effort to produce xylanase from *Penicillium citrinum* through submerged fermentation, achieved an enhanced level production of xylanase by supplementation carbon source were. The molecular identification of fungal strain was added.

MATERIALS AND METHODS

Fungal Strain

The *Penicillium citrinum* strains were isolated from different soils. Soils are taken from different regions from in and around Bangalore and tentatively identified in the laboratory.

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Screening of Xylanase Producers

*Penicillium citrinum* strains were screened for their xylanase activity by plate assay. (Plate-1) and among the thirty isolates, *Penicillium citrinum* KGSN 05 were used for further studies. The selected *Penicillium citrinum* KGSN 05 were confirmed at molecular level in next steps.

**IDENTIFICATION AND CONFIRMATION OF FUNGAL STRAIN BY MOLECULAR LEVEL**

i. **DNA extraction:**

The genomic DNA was isolated from the given fungus using fungus genomic DNA extraction Kit.

ii. **PCR Amplification:**

a. **Amplification of the ITS region (ITS1, ITS2, and 5.8 S rRNA Gene)**

Amplification of the ITS region (ITS1, ITS2, and 5.8 S rRNA Gene) was performed using the Universal primers ITS1 and ITS4

Primer ITS1: 5’- TCCGTAGGTAACCTGCGG-3’

Primer ITS4: 5’- TCCTCCGCTTATGATGC-3’

PCR was performed as follows in a total volume of 50 µl in a 0.2 ml thin walled PCR tube.

<table>
<thead>
<tr>
<th>Components</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nuclease free water</td>
<td>37 µl</td>
</tr>
<tr>
<td>Genomic DNA (0.1µg/µl)</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>Forward Primer (10µM)</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>Reverse Primer (10µM)</td>
<td>2.0 µl</td>
</tr>
<tr>
<td>10X Reaction Buffer</td>
<td>5.0 µl</td>
</tr>
<tr>
<td>dNTP Mix (10mM)</td>
<td>1.5 µl</td>
</tr>
<tr>
<td>Taq DNA polymerase (5 U/µl)</td>
<td>0.5 µl</td>
</tr>
<tr>
<td>Total volume</td>
<td>50 µl</td>
</tr>
</tbody>
</table>

The amplification was carried out in a Master cycler® Thermocycler (Eppendorf, Germany) using the following program.

Initial denaturation of 94°C for 2 minutes followed by 40 cycles of denaturation at 94°C for 1 min, annealing at 55.5°C for 1 min and extension at 72°C for 30 sec. Final extension was carried out at 72°C for 10 min. The ~ 600 bp PCR products were purified to remove unincorporated dNTPs and Primers before sequencing using PCR purification kit (Norgen Biotek, Canada).

iii. **Sequencing:**

Both strands of the rDNA region amplified by PCR were sequenced by automated DNA sequencer -3037x/ DNA analyzer from Applied Biosystems using BigDye® Terminator v3.1 cycle sequencing Kit (Applied Biosystems). Sequence data were aligned and dendrograms were generated using Sequence analysis software version 5.2 from applied biosystems. The sequences obtained for plus and minus strands were aligned using DNA baser program before performing the bioinformatics analysis.

iv. **Bioinformatics analysis:**

Sequences were compared to the non-redundant NCBI database by using BLASTN, with the default settings used to find the most similar sequence and were sorted by the E score. A representative sequence of 10 most similar neighbors was aligned using CLUSTAL W2 for multiple alignment with the default settings. The multiple-alignment file was then used to create phylogram using MEGA5 software.

**INFLUENCE OF CARBON SOURCE FOR THE BIOSYNTHESIS OF Xylanase**

A set of conical flasks with 100 ml of production medium supplemented with a particular carbon source with concentrations ranging from 0.25% to 1.0% with increments of 0.25%. The different carbon sources like, monosaccharides (glucose) and disaccharides (Maltose and sucrose) were used under the present study.

The production medium consists (mg/100 ml) of sucrose 3, di potassium hydrogen phosphate 0.1, MgSO4,0.05g, KCl 0.05g, NaCl, 0.01%, FeSO4. The condition of the fermentation medium is as follows .pH 6 temperature 30°C and inoculums size is of 0.5 ml.

**EXTRACTION OF Xylanase**

The samples were withdrawn periodically at 24 hrs in aseptic condition. The extract was filtered through Whatman filter No.1. The clear extract was centrifuged at 2000-3000 rpm for 15 min, supernatant were used as enzyme preparation. Thus prepared crude enzyme was used for assay of xylanase.

**ASSAY OF Xylanase**

The xylanase activity was determined by measuring the release of reduced sugars from oat spelt Xylan (1% w/v) by dinitrosalicylic acid method. The enzyme solution (0.5 ml) and 0.5 substrate (xylan 1% w/v) along with 1 ml of buffer were taken in a test tube. The tubes were then allowed to stand at room temperature for 10 mins, and 3ml of dinitrosalicylic acid was added to arrest the reaction. After the addition of dinitrosalicylic acid, the tubes were placed in boiling water bath for 10 min. The color which had developed was read at 540nm. A blank test tube was prepared by adding dinitrosalicylic acid prior to the addition of enzyme to the test tubes.

**INTERNATIONAL UNIT (IU)**

One unit of xylanase was defined as the amount of enzyme required to release 1μmol of xylose from oat spelt xylan in one minute under standard assay conditions.

**RESULTS AND DISCUSSION**

Thirty *Penicillium citrinum* isolates were isolated from different soil samples from Bangalore. All thirty isolates
were named serially *Aspergillus* KSN1-KSN30 and used for screening of xylanase production by plate assay method. Out of thirty isolates *Penicillium citrinum* KGSN 05 were showed maximum enzyme hydrolytic zone were observed.

Fungal isolates were identified as *Penicillium citrinum* KGSN 05 identified in the laboratory. The extracted genomic DNA from *Penicillium citrinum* was represented in plate-1. Amplification of the ITS region (ITS1, ITS2, and 5.8 S rRNA Gene) were run on 1.2 % Agarose gel electrophoresis showed PCR product of ~600bp and represented plate-2. The aligned sequence of ITS region were represented in plate-3.

The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model. The tree with the highest log likelihood (-759.4784) is shown. Initial tree(s) for the heuristic search were obtained automatically as follows. When the number of common sites was < 100 or less than one fourth of the total number of sites, the maximum parsimony method was used; otherwise BIONJ method with MCL distance matrix was used. The analysis involved 11 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There were a total of 489 positions in the final dataset. Evolutionary analyses were conducted in Molecular Evolutionary Genetics Analysis (MEGA5) Plate-4.

Plate-1: 0.8% Agarose gel electrophoresis of Fungal Genomic DNA. (Lane 1)

Plate-2: Agarose gel electrophoresis SM- standard marker. Lane 1-sample PCR product of ~600bp

Plate-3: Aligned sequence of ITS region
The results on the studies pertaining to the production of xylanase by *Penicillium citrinum* KGSN 05 on synthetic medium supplemented with different concentrations of various carbon sources like glucose (monosaccharide), maltose (disaccharide) and sucrose (disaccharide) are presented in Fig-1, Fig-2 and Fig-3 respectively.

The process economization for xylanase production with carbon sources supplemented to the production medium were carried out with concentration of 0.25%, 0.5%, 0.75% and 1.0%. The results revealed that all the carbon sources employed under the present study have enhanced the production of xylanase up to 0.5% of monosaccharide and disaccharides (glucose, maltose and sucrose) at 72 hrs of fermentation, thereafter no significant production of xylanase was observed on all the days of fermentation period. In case of monosaccharides (glucose) the maximum xylanase production of 5.87 IU was observed at 0.5% and where as disaccharides like maltose yielded maximum xylanase of 4.65 IU.

Ana Cláudia Elias Pião Benedetti et al. suggest that extracellular xylanase activity in this fungus is induced by xylan and xylose (up to 1%). However, at 2%, xylose partially represses the enzyme. Pang Pei Kheng and Ibrahim reported xylose has been described as an effective inducer and carbon source for xylanase production. Our results are coincides with the Ana Cláudia Elias Pião Benedetti et al.

Figure 1: Effect of Glucose on Xylanase production
Figure 2: Effect of Maltose on Xylanase production

Figure 3: Effect of Sucrose on Xylanase production

CONCLUSION:
Among the thirty isolates Penicillium citrinum KGSN 05 were showed better xylanase production. The enhanced level of biosynthesis of xylanase observed by supplementation of carbon source such as glucose, maltose and sucrose have showed better yield under submerged fermentation by using synthetic medium.

REFERENCES