Purification and Characterization of Cellulase from Aspergillus niger Causing Soft Rot of White Yam in Three Environments in Nigeria

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Abstract: Aspergillus niger is abundant in most tropical soils and invariably, on the surface of yam tubers while still attached to the plant and on the root hairs during harvesting or storage. Naturally, the peridermic surface of the tubers function to exclude pathogen but damage caused by accidental incision or cut surface during weeding, insect attack and harvesting provide avenue for the infection. Aspergillus niger isolated from diseased yam in three zones in Nigeria synthesized cellulase which caused soft rot of the yam within nine days of inoculation. Microscopic and molecular analyses revealed two isolates of A. niger, P1 and P2, from different environment produced cellulase enzymes in significantly different proportions. When protein extracts from the infection were subjected to molecular exclusion chromatography, three peaks of absorption (A, B and C) were produced with only the components of peak A showing cellulase activity. Further fractionation of the components of peak A produced two absorption peaks (Aa and Ab) with only component Aa showing Cellulase activity. A. niger isolates, P1 and P2 showed considerable differences in the intensity of cellulase production suggesting that multiple strains of A. niger in the soil of yam-growing environments synthesized cellulase as transcriptional products in different manner underscoring the effect of physico – chemical properties of the soil on infectivity and virulence of the organism during yam rot.

Keywords: Aspergillus niger, Cellulase, Soft Rot

1. Introduction

The black mould rot of yam is caused by Aspergillus niger. The infection is characterized by decay of entire tuber during which the cells collapse with loss of integrity as a functional unit. In host-pathogen interaction, the ability of the pathogen to produce extracellular enzymes capable of degrading the host tissues is one of the more obvious properties influencing virulence [1]. A. niger is abundant in most tropical soils and, invariably, on the surface of yam tubers while still attached to the plant and on the root hairs during harvesting or storage. Naturally, the peridermic surface of the tubers function to exclude pathogen but damage caused by accidental incision or cut surface during weeding, insect attack and harvesting provide avenue for the infection. According to the study [2], chemotaxonomic characteristic is a virulence factor with variation arising from a multiplicity of factors including the nature of soil ecology and production of primary and secondary metabolites such as enzymes, aflatoxins and phytoalexins [1]. A. niger is able to synthesize cellulase enzyme complex that breaks down cellulose of the cell wall components into glucose [2] giving the organism access to utilize the tissues for its metabolism. According to these studies [1, 10], Cellulase {EC 3.2.1.4} secreted by pathogens play a critical role in softening and disintegrating the cell wall material. Cellulase facilitates the penetration and spread of the pathogen in the host and causes the collapse and
disintegration of the cellular structure thereby aiding the pathogen in the propagation of the disease [3, 9].

The environment in which a pathogen grows dictates to a large extent, the quality and nature of enzymes involved in the infection and degradation of the host tissues. This study examined the pattern of synthesis of cellulase as transcriptional product of A. niger causing soft rot of yam from three different environments in Nigeria.

2. Materials and Methods

2.1. Organism and Culture Condition

A. niger was obtained from diseased yam from three yam growing zones in Nigeria; Abuja, Iwo and Owerri. It was grown in Petri dishes containing Potato Dextrose Agar (PDA) medium in aseptic conditions. The plates were incubated at ambient temperature and then subcultured on the same media plates and the fungi spores from 72hr – old culture was used to inoculate healthy yam tubers.

2.2. Extraction and Purification of Enzymes

Inoculated and uninoculated tissues of yam in the three environment were extracted for enzyme activity every 24hr. The extractant was 0.5N NaCl in 0.02M citrate phosphate buffer (pH 5.0).

2.3. Precipitation and Dialysis of Enzyme

The enzyme was partially purified by ultracentrifugation followed by ammonium sulphate precipitation and dialysis for 24 hours at 4°C against the buffer (pH 6.0) using acetylated cellophane tubing [4, 5]. The protein content was determined by the method of these studies [5, 6].

2.4. Cellulase Assay

The assay for Cellulase was determined by the modified Dinitro-Salicylic Acid (DNSA) method of the study [4]. Glucose was used as standard and one unit of cellulase activity was defined as the amount of enzyme in 1ml of the reaction mixture required to liberate reducing sugar equivalent to 10mg glucose in one minute under the specific condition of the reaction on application of enzyme.

2.5. G-75 Column Calibration

A vertical glass tube chromatography column (640 x 25mm) was calibrated with proteins of known molecular weights according to the study [4], and employed for the fractionation of the enzyme. 10ml of dialysed enzyme concentrate was applied to the column (280 x 250mm) of Sephadex CM G-50 and the fraction were collected and measured for protein content and cellulase assay. The effect of physico-chemical factors of soil on purified enzyme (G-75 factor) was determined. The effects of temperature, pH and cations (k+, Na+ and Ca2+) on enzyme activity in the three yam growing zones were determined by the method of the study [6].

2.7. Morphological Identification

Each of the A. niger isolated from decayed yam in the three environments were identified using the manuals about the genus Aspergillus [6]. Slides were stained with cotton blue and mounted in Lactophenol. Photomicrographs were taken with digital canon camera (A550, 7.1 megapixels).

2.8. Preparation of Genomic DNA

The method [7] was used for DNA isolation. Dissolved generic DNA samples from each of the A. niger isolates P1 - P6 were stored at – 200°C.

2.9. DNA Amplification and Electrophoresis

Polymerase Chain Reaction (PCR) was carried out on the isolated DNA of A. niger from the different environments using the amplification kit and automated programmable PCR thermal cylinder with ITS 1 and ITS 4 as primers at the DNA laboratory, Ungwan Sarki, Kaduna, Nigeria. Amplified fragments P1, P2, P3, P4, P5 and P6 were each separated in agarose gel in the electrophoretic chamber and Chemidoc was used to produce camera snaps of DNA band pattern according to the method of the study [8].

3. Results

3.1. Infection of Yam Tissues by A. niger

A. niger caused extensive degradation of healthy yam tissues at the point of infection irrespective of the source of yam. Extensive sporulation occurred at the point of inoculation. Tubers aseptically inoculated with sterile water lacked sporulation. Infection of the Yam tissues was very slow in yams placed on bare ground devoid of soil.

When the organisms from different yam-growing zones were sub-cultured on Potato Dextrose Agar (PDA), there were variations in the cultural appearance of conidiospores of A. niger appearing on agar plates. Structural morphologies of the isolates also showed some differential characteristics.

3.2. Microscopic Examination of A. niger

Microscopy of P1 and P2 isolates showed a large black conidial that are arranged in a globose biseriate head arising from a spherical conidiophore. The black coloured colonies were identified as A. niger based on the structural morphologies as observed under the light microscope. It was observed that the isolates possessed distinct conidiophores terminated by a swollen vesicle bearing flask-shaped
phialides. The black-coloured colonies consist of a compact white or yellow basal felt covered by a dense layer of dark-brown to black conidial heads similar to the observation of the study [8]. The ones with the yellow basal felt (marked P1 isolates) were isolated from Zone 1 (Abuja) while the ones with yellow and white basal felt (marked P1 and P2) were both isolated from Zones 2 and 3 (Iwo and Owerri).

### 3.2.1. Macroscopic Characterization of P1

Colonies on PDA plates attained 31mm in diameter in 7 days at 27°C colony coloured brown to dark, reverse is light yellow and transparent (Figure 1).

![Figure 1. Growth of A. niger Strain Isolate P1 from Infected Tubers on PDA Plates.](image1)

**Figure 1.** Growth of A. niger Strain Isolate P1 from Infected Tubers on PDA Plates.

### 3.2.2. Microscopic Characterization of P1

The conidia head on PDA plates radiate, 80 – 100μ in diameter, conidiophores hyaline, smooth, 100 – 200μ long and 4-5μ wide with 1μ thick wall. Vesicle was globose and 10 – 18μ in diameter. Matulae were ampuliform 5 – 8μ by 1.5 – 2μ wide. Phialides ampuliform, 5-7μ by 1.5-2μ wide. Conidia were also globose, 1.6 – 2.5μ in diameter (Figure 2).

![Figure 2. Growth of A. niger Isolate P2 from Infected Yam Tubers.](image2)

**Figure 2.** Growth of A. niger Isolate P2 from Infected Yam Tubers.

### 3.2.3. Macroscopic Characterization of P2

The conidia based on PDA plates attained 55mm after 7 days at 30°C colony colour was black (Figure 2 and 3). The reverse side was mostly hyaline to light yellow.

![Figure 3. Microscopic Features of A. niger Isolate P2 Stained with Lactophenol in Cotton Blue (Mag.40x).](image3)

**Figure 3.** Microscopic Features of A. niger Isolate P2 Stained with Lactophenol in Cotton Blue (Mag.40x).

**Figure 4.** Microscopic Features of A. niger Isolate P2 Stained with Lactophenol in Cotton Blue (Mag.40x).

### 3.2.4. Microscopic Characterization of P2

Conidial heads on PDA plates radiate conidiophores that are 300-400μ long, 8-12μ wide with 1.5-2.5μ thick wall. Vesicles are globose and 3.5 -4μ in diameter. Conidial heads are also globose 3.5 – 4μ in diameter. Phialides are ampuliform and metulae club shaped (Figure 4).

### 3.3. Molecular Characterization

The two different isolates of *A. niger* identified through their cultural, and morphological characteristics, were further characterized molecularly. When the DNA of the two isolates (P1 and P2) were prepared, quantified and amplified for agarose-gel electrophoresis, six distinct bands were observed suggesting that there are differences in the nucleotide base sequences coding in the two *A. niger* isolates for enzyme production in different isolates. The molecular detection and amplification of the gene coding for *A. niger* cellulase enzymes associated with yam rot was shown in the gel electrophoresis of the DNA fragments of *A.niger* isolates from different yam-growing environments. This is evident with the appearance of bands on the extracted DNA of *A. niger* as shown in Figure 5.

The extent of yam rot may depend on the sample of the *A. niger* associated with the different soil or environment. The strains of *A. niger* may vary depending on their location and the arrangement of the nucleotide sequences in the genome of the organism. This results suggest that cellulase, is associated with yam soft rot as observed from different environmental conditions and that the extent of rot is also dependent on the strain of *A. niger* that confers differences in pathology.
3.4. Infection of Yam by *A. niger* and Enzyme Production

Yam tissues infected with *A. niger* isolates exhibited cellulase activities (Figures 6, 7, 8, 9, 10 and 11).

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**Figure 5.** Agarose Gel Electrophoresis: Molecular Detection and Amplification of DNA of *A. niger* Isolates from Rotten Yam.

**Figure 6.** Enzyme Activities of Yam Tissues Infected by *A. niger* (P1 Isolate) in Env. 1 (Abuja) by Incubation Period.

**Figure 7.** Enzyme Activities of Yam Tissues Infected by *A. niger* (P1 Isolate) in Env. 2 (Iwo) by Incubation Period.

**Figure 8.** Enzyme Activities of Yam Tissues Infected by *A. niger* (P1 Isolate) in Env. 3 (Owerri) by Incubation Period.

**Figure 9.** Enzyme Activities of Yam Tissues Infected by *A. niger* (P2 Isolate) in Env. 3 (Iwo) by Incubation Period.

**Figure 10.** Enzyme Activities of Yam Tubers Infected by *A. niger* (P2 Isolate) in Env. 3 (OWERRI) by Incubation Period.

**Figure 11.** Elution Volume against log (M.WT) for proteins in Peaks A, B and C Calibrated by a Gel Filtration with Sephadex G-75.
Generally, the activity of each of the enzymes increased with the intensity of infection. Uninfected yam tissues lacked any appreciable decline in the infection of yam. In Zone 1 (Figure 7) the activity of cellulase increased progressively and continued until the eighth day where it reached a maximum and in the ninth day when it started to decline. However, enzyme activity of \textit{A. niger} from zones 2 and 3 could be noticed as from the second day of inoculation. Cellulase was the first to be detected in appreciable quantity. The activity of enzymes in yam Zone 2 continuously increased to the tenth day when it started to decline (Figure 7). An almost similar pattern of increase and decline was observed in yams infected with \textit{A. niger} in yam Zone 3 (Figure 9).

The nature of the soil around which the tuber is grown or harvested and their physico-chemical properties was observed to contribute to the intensity of enzyme production, infectivity and tuber damage. Sporulation of \textit{A. niger} and yam rot was very slow when the yam was placed on bare ground. In some cases, sporulation was restricted to the zone of inoculation and rotting took a longer time than when the yam was damaged while attached to the plant or placed on the soil in ambient conditions after harvest underlining the influence of soil ecology on infectivity.

3.5. Purification of Enzymes; Enzyme Separation on Sephadex; G-75 and CM G-50

Fractionation of the enzyme concentrate on Sephadex G-75 produced three absorption peaks marked A, B and C. (Figure 12). The molecular weights of the components estimated from the calibration using their respective elution volumes are about 44,670 (Peak A), 17,780 (Peak B) and 12,590 (Peak C). The activities of cellulase were detected only in Peak A. Further fractionation of Component A on CM-Sephadex CM-C50 gave two new absorption peaks marked (Aa) and (Ab) respectively (Figure 13). Similarly, the activity of cellulase was detected only in Peak (Aa).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image12.png}
\caption{Separation of Protein in the Concentrated Extract of \textit{A.niger} Infected Yam Tissues by Molecular Exclusion Chromatography and the Enzyme Activity of the Fraction towards Pectin and CM-cellulase.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image13.png}
\caption{Separation by Ion-exchange Chromatography of High Molecular Weight Proteins (Fractions 12-50) Produced from \textit{A. niger} Infected Yam Tissue Extracts by Gel Filtration and the Enzyme Activity of the Fractions towards Cellulase and Pectin.}
\end{figure}
Table 1 shows the purification levels of cellulase. The total activity of the crude enzyme at maximum yield was 29,400 units and specific activity of 73.3 units/mg protein. These mean activity values were derived from the replicates with the limit of standard error. When the proteins in the crude enzyme were subjected to ammonium sulphate precipitation, the specific activity increased to 209.3 units/mg proteins and a yield of 85.4% was obtained with 2.9 fold purification. Further fractionation and purification by molecular exclusion chromatography (Sephadex G-75) and ion-exchange chromatography (CM-Sephadex G-50) yielded much more purified enzyme up to 30.7 fold by gel filtration and 77.3 fold by ion-exchange chromatography. The specific enzyme activity increased to 2,215 units/mg protein and 5,581 units/mg protein by gel filtration and ion-exchange chromatography respectively. The partially-purified cellulase enzyme, G-75 fraction was used for further analysis.

Table 1. Partial Purification of cellulase Obtained from Yam Infected with A. niger.

<table>
<thead>
<tr>
<th>Purification Step</th>
<th>Total Protein (mg)</th>
<th>Total Activity (Units)</th>
<th>Specific Activity (units/mg Protein)</th>
<th>Yield %</th>
<th>Purification (Fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude Enzyme</td>
<td>402 ± 1.0</td>
<td>29400 ± 1.7</td>
<td>73.3 ± 0.9</td>
<td>100</td>
<td>1.</td>
</tr>
<tr>
<td>(NH4)2SO4 Precipitate</td>
<td>120 ± 0.7</td>
<td>25120 ± 2.0</td>
<td>209.3 ± 1.0</td>
<td>85.4</td>
<td>2.9</td>
</tr>
<tr>
<td>Sephadex G-75 (Peak Aa)</td>
<td>8.4 ± 0.2</td>
<td>18610 ± 1.0</td>
<td>2215.5 ± 1.2</td>
<td>63.3</td>
<td>30.7</td>
</tr>
<tr>
<td>CM-Sephadex G-50</td>
<td>2.9 ± 0.1</td>
<td>16185 ± 0.8</td>
<td>5581.0 ± 0.8</td>
<td>55.1</td>
<td>77.3</td>
</tr>
</tbody>
</table>

3.6. Effect of Soil Temperature

As shown in Figures 14, 15 and 16, optimum enzyme activity was observed at about 40°C in yam rot caused by A. niger in all the three zones (Zones 1, 2 and 3). Temperatures lower than 25°C and higher than 40°C resulted in diminished enzymes activity and the enzyme was almost completely inactivated at 50°C. This indicated that the quantity of cellulase activity was completely influenced by temperature. Aspergillus niger growing on yam tubers in the three ecozones achieved maximum sporulation and enzyme activity at ambient temperature of 40°C.

In Zone 1, the enzyme activity of cellulase was very slow at temperature of 20°C and 30°C. This might have contributed to the slow sporulation at the temperatures lower than 35°C.

In Environment 2 (Iwo), there was a steady increase in the activity of cellulase as from 20°C until the activity of enzyme reached a maximum at 40°C. Enzyme activity of A. niger cellulase was affected by increasing soil temperature in Environment 3 (Owerri). The enzyme activity was optimum at 4°C after which increasing the temperature above resulted in a sharp decline of activity.

3.7. Effect of Soil pH

Figures 17, 18 and 19 showed that the intensity of enzyme production is greatly influenced by changes in the hydrogen ion concentration (pH) of the soil. Optimum production occurred in slightly acidic medium of between pH 4.0 and 6.0 in the three environments.
Cellulase enzyme can tolerate high concentration of cations of up to 100mM for Ca²⁺, Na²⁺ and K⁺ with simulating effects before enzyme activity began to slow down gradually.

4. Discussion and Conclusion

The results showed that A. niger caused extensive degradation of yam tissues after nine days of infection. The statistical significance is in agreement with the practical significance buttressed in the analysis of variance (ANOVA) of the mean of replicate values across the three environments which showed significant variations in the extent of cellulase produced. The sum of squares of between group was moderately high (23280.150 at 2,38, df respectively). The value of p = 0.48 is less than 0.05 indicating statistical significance in the pattern of cellulase production in the infection of yam by A. niger. The results of multiple comparisons also indicated that the mean difference in the intensity of enzyme production in the three yam-growing environments studied is significant at 0.05 level.

Soil ecology has also been observed to play a prominent role in the infection process as washed yams that were placed on bare concrete floor without soil on them took a much longer time to register decay of the yam tissues. The increase in the rate of enzyme production and infectivity might be attributed to the virulence factor of the isolate of Aspergillus niger involved in the infection which has contributed to its pathogenicity.

Identified through their cultural, microscopic and morphological characteristics, the isolates of A. niger labelled P1 and P2 (isolated from infected yam) were the same organisms isolated from the soil of the yam-growing areas investigated.

When the DNA of the two isolates (P1 and P2) were prepared, quantified and amplified for agarose gel electrophoresis, six distinct bands were observed suggesting that there are possibly differences in the nucleotide bases sequences in the two A. niger isolates in the three environments. This may also be responsible for the variations in the production of cellulase as an aid to infection of yam in the three yam-growing environments studied.

The mere fact that yam tubers placed on bare soil without the effect of the physicochemical properties of the soil ecology playing a facilitative role failed to develop appreciable sporulation and decay at the point of infection emphasized the importance of the soil ecology on infection. Yam tubers grow in the soil and when harvested usually have soil particles attached to the root hairs. The knowledge of the understanding of the physicochemical properties of the soil ecology is important to quantify the conditions that are favourable for infection and disease development.

This may provide insights into the development of measures that can be used to prevent or slow down the infection. This may also lead to the devising of mechanisms for preservation of the tubers while still attached to the plant in the soil and inhibit or slow down deterioration during storage. In a similar vein, having proper knowledge of the genetic blueprint for enzyme production by strains of A. niger will increase the understanding of the organisms metabolic pathway that can be used to slow down or prevent the infection process and increase the shelf life of yam during storage. This of course, can be applied to other crops facing soft rot during storage.

References


