A Comparison of Methods to Determine Tannin Acyl Hydrolase Activity

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ABSTRACT

Six methods to determine the activity of tannase produced by Aspergillus niger Aa-20 on polyurethane foam by solid state fermentation, which included two titrimetric techniques, three spectrophotometric methods and one HPLC assay were tested and compared. All methods assayed enabled the measurement of extracellular tannase activity. However, only five were useful to evaluate intracellular tannase activity. Studies on the effect of pH on tannase extraction demonstrated that tannase activity was considerably under-estimated when its extraction was carried out at pH values below 5.5 and above 6.0. Results showed that the HPLC technique and the modified Bajpai and Patil methods presented several advantages in comparison to the other methods tested.

Key words: Tannin acyl hydrolase, assay, polyurethane foam, solid state fermentation, tannase

INTRODUCTION

Tannin acyl hydrolase (EC 3.1.1.20) or tannase is an enzyme which hydrolyses the ester and depside bonds of hydrolysable tannins and gallic acid esters (Lekha and Losane, 1997). It breaks down tannic acid, releasing gallic acid and glucose (Haworth \textit{et al}., 1958). Tannase is produced by fungi, mainly by species of Aspergillus, Penicillium, Fusarium and Trichoderma (Iibuchi \textit{et al}., 1967; Rajakumar and Nandy, 1983; Kawakubo \textit{et al}., 1991; Lekha and Lonsane, 1994; García-Peña, 1996; Bajpai and Patil, 1996), but also by bacteria (Deschamp \textit{et al}., 1983) and yeast (Aoki \textit{et al}., 1976). Tannase activity has also been detected in the fruit of several plants (Sourlangas, 1947; Madhavakrishna \textit{et al}., 1960).

The major commercial applications of tannase are in the manufacture of instant tea or acorn wine and the production of gallic acid (Coggon \textit{et al}., 1975; Chae and Yu, 1983; Pourrat \textit{et al}., 1985). The latter is a key intermediate required for the synthesis of an antifolic antibacterial drug, trimethoprim, used in the pharmaceutical industry (Sittig, 1988). Gallic acid is also a substrate for the chemical or enzymatic synthesis of propyl gallate in the food industry. In addition, tannase is used as a clarifying agent in wine, fruit juices and coffee-flavoured soft drinks (Lekha \textit{et al}., 1993).

Several titrimetric (Nishira, 1961; Haslam and Stangroom, 1966; Yamada \textit{et al}., 1967), photometric (Chen, 1969), colorimetric (Haslam and Tanner, 1970), UV-spectrophotometric (Iibuchi \textit{et al}., 1967; Aoki \textit{et al}., 1976; Rajakumar and Nandy, 1983; Sanderson \textit{et al}., 1974; Bajpai and Patil, 1996) and chromatographic (Jean \textit{et al}., 1981; Beverini and Metche, 1990) methods to quantify enzyme-released gallic acid have been proposed. Some methods are non specific or present problems in determining an endpoint accurately. In addition, conditions of enzymatic reaction are different. Some others are long and tedious, requiring an extended reaction time. Often, non readily available substrates are used. In the spectrophotometric method described by Sanderson \textit{et al}. (1974), tannase activity was not expressed in standard international units. Such methods have been previously reviewed by Lekha and Lonsane (1997), Madhavakrishna \textit{et al}. (1960), Jean \textit{et al}. (1981) and Bajpai and Patil (1996).

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In order to classify, evaluate and select an adequate method for tannase activity and to attempt to homogenize the data presented in the literature concerning tannase units, six methods for the determination of tannase activity were compared. Additionally, the effect of sample and extractant solution pH on tannase activity was considered. Along with the results, the advantages and disadvantages of each method are discussed.

MATERIALS AND METHODS

Microorganism and culture medium. Aspergillus niger Aa-20 (IRD-UAMI collection) was used in all experiments. Medium composition was the same as reported previously by Lekha and Lonsane (1994) using \((\text{NH}_4)_2\text{SO}_4\) as nitrogen source. Salt-containing medium was autoclaved at 121°C for 15 min. Tannic acid (Sigma, U.S.A) solution was filter-sterilized and added to a final concentration of 25 gL\(^{-1}\).

Inoculum preparation. Spores from A. niger Aa-20, stored at -20°C in protect-crioblocks (bead storage system, Technical Service Consultants Limited) were used to inoculate 50 mL of PDA medium contained in two 125 mL flasks. The flasks were incubated at 30°C for 5 days, and the spores were scraped into 0.01% Tween 80 solution and counted in a Neubauer chamber.

Enzyme production in solid state fermentation (SSF). The SSF process involved the use of polyurethane foam (PUF) (Expomex, Mexico) as support to absorb the liquid medium. PUF was washed using the technique described by Zhu et al. (1994) and then pulverized in a plastic-mill. Glass columns (25mm x 180 mm) were packed with 10 g of inoculated PUF(2 x \(10^7\) spores g\(^{-1}\) of dry inert carrier)impregnated with the medium (1.94 mL of inoculated medium per gram of dry inert carrier). Culture conditions were: 30°C; aeration rate, 20 mL of air per gram of inert carrier per min; initial pH, 5.5; initial moisture content, 65%, and incubation time, 48 h. For enzyme leaching, the content of each column was mixed with distilled water (10:1 vw\(^{-1}\)) and vortexed for 1 min. Solid particles were filtered (Whatman 41) and the clear filtrate was assayed for extracellular tannase activity. The remaining solids were washed three times with 50 mL of distilled water. Intracellular enzymes were recovered by freezing the cells in liquid nitrogen and by macerating in a chilled mortar. The macerated material was recovered in an equal amount of 0.02 M acetate buffer (pH, 5.0) and centrifuged at 5000g for 20 min. The resulting supernatant was assayed for intracellular tannase activity. Each data point was obtained from triplicate assays and the average values were reported as mean values \(\pm\) standard deviations.

Tannase activity assays. Six tannase activity methods were assayed: A) HPLC assay as reference method (Beverini and Metche, 1990), B) unbuffered titrimetric assay (Haslam et al. 1961), C) buffered titrimetric assay (Yamada et al. 1967), D) single wavelength spectrophotometric assay (Skene and Brooker, 1995), E) double length spectrophotometric assay (Bajpai and Patil, 1996), and F) a modified double length spectrophotometric assay (present work). In the latter, the modification consisted in the addition of the enzymic reaction described by Iibuchi et al. (1967) to the spectrophotometric assay described by Bajpai and Patil method (1996). An equation using specific extinction coefficients was established to estimate the concentration of gallic acid released during the reaction. Tannic acid was the sole substrate used in the enzymatic reaction.

Effect of pH. Fermented material was extracted with a sodium acetate buffer (200 mM) at varying pH (3.0 - 7.5). Each data point was obtained in triplicate and the average values were reported as mean values. Tannase activity was assayed by HPLC.

Statistical Analysis. Analysis of variance and t-test were used to evaluate all the obtained results. Statistical analysis was made using InStat for Macintosh, version 2.03, 1994 Graphpad Software.
RESULTS AND DISCUSSION

Table 1 shows the equations applied to calculate enzyme activity in all the six cases. In addition, the reaction mixture and conditions used are presented. For the spectrophotometric and HPLC methods, a unit of enzyme activity was defined as the amount of enzyme liberating 1 µmol gallic acid per mL per min; while for the titrimetric methods, a tannase unit was the amount of enzyme able to release 1 µmol carboxyl group per mL per min. Features of method F were as follows: the two wavelengths for maximum difference in the absorbance of gallic and tannic acids were 254 nm and 290 nm, respectively. The specific extinction coefficients at these wavelengths were found to be 30.77 and 11.39 for gallic acid and 26.34 and 41.45 for tannic acid respectively. Using these values, equation of method F for the estimation of gallic acid released after the enzymic reaction was deduced (Table 1).

All the six methods tested enabled the evaluation of extracellular tannase activity. However, for intracellular tannase activity, the original double length spectro-photometric assay (E) of Bajpai and Patil (1996) could not be applied because the sample used to assay for activity was the culture medium itself, after filtration and proper dilution without carrying out the enzymatic reaction. This aspect was its major drawback. Moreover, the results obtained for extracellular activity showed the highest standard deviation. Except for this method, all results for extracellular enzyme activities yielded values of tannase activity within the same range (1.5 - 5.0 I.U.), while the values obtained with the spectrophotometric method (E) were 9 times higher. This could be due to the fact that the values obtained through this method represented the amount of gallic acid liberated in the fermentation process, which was constantly consumed by the organism thereby showing point to point variation.

Statistical analysis of measurements of extracellular tannase indicated that all were different (p < 0.01) except for technique A (HPLC) and F (modified spectro-photometric) which were quite similar (p > 0.05). Measurements of intracellular activities had high coefficients of variation (around 23%) as compared with the measurements of extracellular activities (around 14%) suggesting that there were significant experimental variations in the extraction of tannase. However, the smallest value of the coefficient of variation was related to HPLC technique indicating a higher reliability than others.

Table 1: Conditions and equations used to calculate tannase activity produced by A. niger Aa-20 on SSF.

<table>
<thead>
<tr>
<th>Method</th>
<th>Reaction mixture</th>
<th>Equation</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A)</td>
<td>Tannic acid (0.51 g/l)</td>
<td>1.0 mL</td>
<td>Standard sample read at λ = 280 nm</td>
</tr>
<tr>
<td></td>
<td>In acetate buffer (pH 5.0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Enzymic extract</td>
<td>0.05 mL</td>
<td></td>
</tr>
<tr>
<td>(B)</td>
<td>Tannic acid (1 g/l)</td>
<td>20.0 mL</td>
<td>U = Vol. NaOH x N NaOH x 10^3 R x min</td>
</tr>
<tr>
<td></td>
<td>Enzymic extract</td>
<td>2.0 mL</td>
<td>R = mL extract/mL substrate</td>
</tr>
<tr>
<td></td>
<td>PH</td>
<td>5.5</td>
<td></td>
</tr>
<tr>
<td>(C)</td>
<td>0.05 M citrate buffer</td>
<td>40.0 mL</td>
<td>U = Vol. NaOH x N NaOH x F x 10^3</td>
</tr>
<tr>
<td></td>
<td>enzymic extract</td>
<td>10.0 mL</td>
<td>F = mL extract/mL substrate</td>
</tr>
<tr>
<td></td>
<td>tannic acid (1 g/l)</td>
<td>10.0 mL</td>
<td></td>
</tr>
<tr>
<td></td>
<td>pH</td>
<td>6.0</td>
<td></td>
</tr>
<tr>
<td>(D)</td>
<td>Substrate (5.0 mg/mL)</td>
<td>1.0 mL</td>
<td>Standard sample read at λ = 520 nm</td>
</tr>
<tr>
<td></td>
<td>In phosphate buffer (pH 6.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Enzymic extract</td>
<td>1.0 mL</td>
<td></td>
</tr>
<tr>
<td>(E)</td>
<td>0.1 mL of culture medium diluted 1:100 in acetate buffer (pH 6.0)</td>
<td>Gallic Acid (GA) in µg/mL</td>
<td>Bajpai and Patil, 1996</td>
</tr>
<tr>
<td></td>
<td>GA = 21.77 (abs254.6) - 17.17 (abs293.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(F)</td>
<td>Tannic acid (0.35%)</td>
<td>4.0 mL</td>
<td>GA in µg/mL</td>
</tr>
<tr>
<td></td>
<td>In citrate buffer (pH 5.5)</td>
<td>GA = 42.5 (abs254) - 27.0 (abs290)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Enzymic extract</td>
<td>1.0 mL</td>
<td></td>
</tr>
</tbody>
</table>

* (A) HPLC assay, (B) and (C) are unbuffered and buffered titrimetric assays respectively, (D), (E) and (F) are single wavelength, double wavelength, modified double wavelength spectrophotometric methods.
Tannase activity of *A. niger* Aa-20 on SSF after 48 h incubation period is shown in Table 2. From the obtained data, it can be observed that HPLC (A), buffered titrimetric (C), and modified spectrophotometric (F) methods showed the highest values of tannase activity. Among these three methods, B and F presented higher variability. In addition, some other aspects can be considered to select a method of choice (Table 2).

The data obtained from this comparative study indicated that the values of tannase activity could be compared, mainly due to the fact that values were in the same range of magnitude, except for the original Bajpai and Patil assay. Extracellular/intracellular ratios were obtained for each of the methods described. The lowest values were for the HPLC and the titrimetric assay proposed by Yamada et al. (1997). The latter, however, has a disadvantage since large volume of extract was needed (10 mL). In contrast, the HPLC assay required only 50 µL of enzymatic extract and could be the method of choice for purification purposes. However, selection of the method of choice would be a function of precision, sensitivity, time required and equipment available.

<table>
<thead>
<tr>
<th>Method</th>
<th>Extracellular tannase activity (U mL⁻¹ min⁻¹)</th>
<th>Intracellular tannase activity (U mL⁻¹ min⁻¹)</th>
<th>Ratio extracellular: Intracellular</th>
<th>Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>5.30 ± 0.174</td>
<td>1.85 ± 0.173</td>
<td>3:1</td>
<td>Expensive, rapid (40 min), laborious, highly sensitive, requires small amounts of extract.</td>
</tr>
<tr>
<td>B</td>
<td>2.68 ± 0.33</td>
<td>0.38 ± 0.13</td>
<td>7:1</td>
<td>Economical, rapid (30 min), easy, sensitive to changes in pH by other compounds present in the 2 mL of extract.</td>
</tr>
<tr>
<td>C</td>
<td>4.75 ± 0.25</td>
<td>1.66 ± 0.34</td>
<td>3:1</td>
<td>Economical, rapid (30 min), easy, requires great amounts of extract.</td>
</tr>
<tr>
<td>D</td>
<td>1.64 ± 0.55</td>
<td>0.43 ± 0.10</td>
<td>4:1</td>
<td>Relatively expensive, rapid (30 min), laborious, requires rhodamine, not easily available. Protein determination is needed.</td>
</tr>
<tr>
<td>E</td>
<td>44.70 ± 1.73</td>
<td>-</td>
<td>-</td>
<td>It is not an enzyme assay, rather is a method to estimate gallic acid present in the culture medium, very fast but criticizable.</td>
</tr>
<tr>
<td>F</td>
<td>3.70 ± 0.87</td>
<td>0.12 ± 0.03</td>
<td>34:1</td>
<td>Is rapid, easy, relatively expensive and sensitive.</td>
</tr>
</tbody>
</table>

From the results obtained, methods A and F showed major advantages in the evaluation of tannase activity. A comparison of these methods showed that the HPLC method had more precision, sensitivity and presented lower variability (Figure 1a). However, the correlation coefficient was high between both methods (0.965) as shown in Figure 1b. It is important to state, that in both methods the activity is expressed as gallic acid released. When the results obtained in both methods are compared with those obtained by the titrimetric assay (B) where the tannase activity is expressed as amount of carboxyl groups released, it is clear that the discrepancies in the values are low (less than 5%). This corroborates the observations made by Madhavakrishna et al. (1960), Jean et al. (1981), Bajpai and Patil (1996) and Lekha and Lonsane (1997), where differences in the enzymatic activity values depend on the substrate used for the assay. In this study, the use of tannic acid as sole substrate and the measurement of several reaction products resulted in low discrepancies in observed activity values.
Only five of the six methods evaluated, enabled measurement of intracellular tannase activity (with the exception of the Bajpai and Patil method). The difference between values for each was due to sensitivity, precision and mainly to the product of the enzymatic reaction followed. However, about the latter aspect, it is important to consider that the amount of carboxyl groups released is proportional to the amount of gallic acid produced. This is demonstrated by the obtained values between the five methods mentioned. This point is important, as it is clear that the activity is similar in several methods when a same substrate is used. In addition, some of the available methods have at least one drawback, for example, the method reported by Jean et al., (1981) is limited to one substrate (methyl gallate) which measures methanol produced and therefore the compound cannot be substituted; the Skene and Brooker method requires rhodanine which is a non readily available chemical.

Taking into account the different pH values reported to carry out tannase extraction and enzyme assay, a final experiment was conducted determining the effect of sample pH and extraction solution pH on tannase activity assayed by HPLC. The internal control was the same extract after dialysis against a buffer at pH 6.0 and considered as reference (2.7 U/mL).

Figure 2A shows the pattern obtained when the pH of solution was modified. It was clear that at low pH values of the extraction solution (3.0 - 5.5) tannase activity was under-estimated (7% for pH values of 4.5 - 5.0 and >13% for pH values of 3.0 - 4.0). At a pH range from 5.5 to 6.0, 100% of the activity was recovered; pH change due to sample addition was not significant (p < 0.0001), independently of the sample pH at a range from 3.0 to 7.5 (Figure 2B). If an extraction solution at pH 6.0 was used, followed by a variation in the pH of the assay solution, no significant effect was seen on tannase activity (Figure 2B). However, if the pH of the extraction solution was varied, the extracted tannase activity varied greatly.

**Figure 1:** Tannase activity from *A. niger* Aa.20 on SSF using the HPLC and modified double length spectrophotometric methods (A), and relationship between tannase activity values obtained with both methods (B).

**Figure 2.** Effect of the pH of extractant solution on tannase activity (A) and effect of the pH of the sample on tannase activity (B). Enzyme activity was assayed by the HPLC method.
The HPLC technique and the modified Bajpai and Patil method were the fastest, most specific, reliable and reproducible assays under conditions used in the present study. The original Bajpai and Patil method could not be considered as the present study evaluated the gallic acid released in the fermentation process and did not involve an enzymatic reaction. However, we recommend that before selecting a method, the advantages and disadvantages of each method need to be considered. In addition, the enzyme extraction process should be carried out with a buffering solution at a pH of 6.0.

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