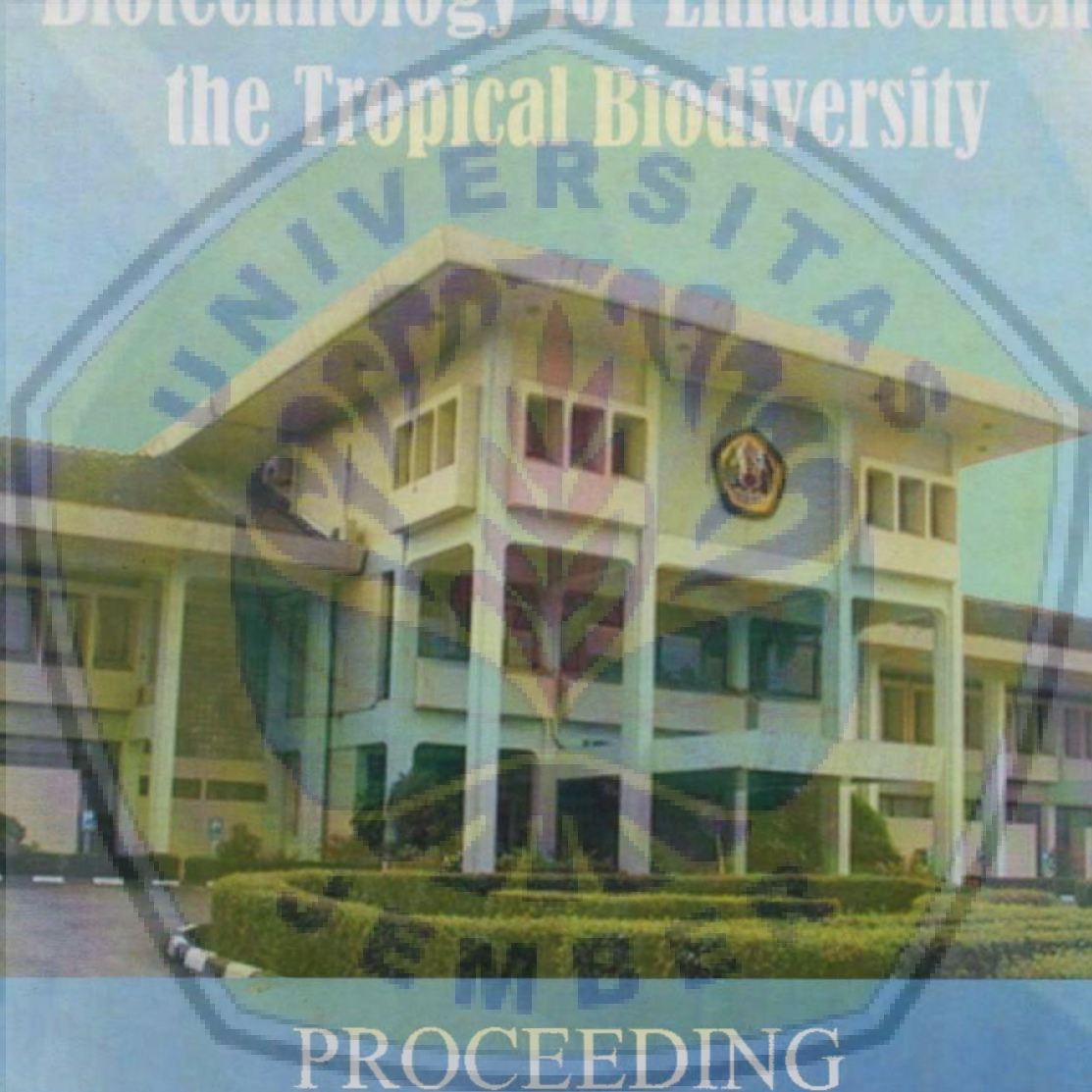




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INTERNATIONAL SEMINAR

# Biotechnology for Enhancement the Tropical Biodiversity



PROCEEDING

UNIVERSITAS PADJADJARAN  
GRAHA SANUSI HARDJADINATA  
BANDUNG, INDONESIA  
OCTOBER 18-20, 2010

Sponsorship :



## PREFACE FROM THE CHAIRMAN OF COMMITTEE

Assalamu'alaikum Wr. Wb

The progress in the subject of life sciences such as molecular biology, molecular genetics and genetic engineering in this century have been packed into a sophisticated technology which is called biotechnology. One of the benefits of biotechnology is its ability to change the nature of the organism into new properties as desired. Adoption to this a new technology seems very necessary to improve production efficiency, increase product quality and minimize environmental pollution. This new technology not only help increase the agricultural and bio-based industries in the present but also for the future.

On the other hand, human life problems of such as food shortages, health problems, environmental degradation both land and water ecosystems, as well as increased air pollution causes reduction in biodiversity. Especially for Indonesia, biodiversity is still constrained problems in optimal utilization to support human life. To solve these problems, required the involvement of biotechnology that have the potential to improve sustainability in the use of the potential of biodiversity in order to maintain the existing biodiversity, so that the problem of human life can be resolved.

Researchers at various universities and research institutions in Indonesia has been conducting research aimed at improving the potential and value of biodiversity for the welfare of the community, that is why we need a scientific communication.

Seminar and Workshop is a scientific interaction between academia, industrial communities, and governments on an international scale to develop the potential of tropical biodiversity by utilizing biotechnology devices that have been growing rapidly. This seminar and all supporting activities will be focused on tropical biotechnology and biodiversity with theme "Biotechnology Enhancement For The Tropical Biodiversity", as part of a series of scientific event of Universitas Padjadjaran to commemorate the 53rd anniversary.

The organizing committee would like to appreciate the Rector, Head of LPPM who has provide great supports for biotechnology research and development. High appreciation and thanks we give to the Minister of Environment for attending on this seminar. We would like to give the highest appreciation for all key note speakers, invited speakers, contributors, and all the participants. High appreciation is also given to the sponsors that make the conference successful

Bandung, October, 19, 2010

The Chairman of Organizing Comitee

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**SCHEDULE**  
**OF INTERNATIONAL SEMINAR**  
**“Biotechnology Enhancement for Tropical Biodiversity”**  
**October 19 – 20, 2010**

Tuesday, October 19, 2010

Time	Activity	Speaker	Remarks
07.30 – 08.30	Registration		
08.30 - 08.35	Opening		MC
08.35 – 08.45	Speech from Head of LPPM	Prof. Oekan S. Abdoellah, MA., PhD.	
08.45 - 09.00	Welcome speech	Rector of Universitas Padjadjaran Prof. Dr. Ganjar Kurnia, DEA	
09.00-09.30	<i>Keynote speaker</i> : National Policy for Developing and Utilizing the Potential of Tropical Biodiversity	Minister of Environment Republic of Indonesia Prof. Dr. Gusti M. Hatta	Prof. Oekan S. Abdoellah MA, Ph.D
09.30-10.00	Coffee Break	Sundanese Art Performance Unpad	
10.00-10.20	Biotechnology Enhancement for Exploration The Potential of Tropical Biodiversity for Health	Prof. Dr. med. Tri Hanggono Achmad Health Biotechnology Universitas Padjadjaran	Prof. Dr. Johan S. Masjhur, dr., SpPD-KEMD, SpKN
10.20-10.40	In vitro Technology for Biodiversity	Dr Kit Sum Canada Genetic Livestock assoc. IN D life; International New Development, China	
10.40-11.00	Biotechnology for Commerce in Livestock	Dr. Robert Lang Canadian Livestock Genetic Association	
11.00-11.20	Biotechnology and Genetic in Aquaculture	Prof. Dr. Ir. Ketut Sugama, M.Sc Dirjen Budidaya-Kementerian Kelautan Perikanan RI	

11.20-11.35	Biotechnology Products	PT. Dipa Puspa Labsains	
11.35-12.00	Discussion		
12.00-13.00	Lunch		
13.00-13.20	Transplant of Stem Cells in Regenerative Therapy of Spinal Cord Injuries	Prof. Gheorghe Solcan Facultatea De Medicina Veterinara Romania	Sukma Nuswantara, PhD.
13.20-13.40	Embryo Transfer in Cattle	Siti Darodjah Rasad, PhD Husbandry Faculty-UNPAD	
13.40-14.20	Discussion		
14.30-16.45	Paralel Session	Class 1 A : Biotechnology in animal, fish, food and feed 14.30 -16.45	
		Class 1 B: Biotechnology in animal, fish, food and feed 14.30 - 16.45	
		Class 2 : Health Biotechnology 14.30 -16.45	
		Class 3 : Biotechnology and energy 14.30-16.45	

**XYLANASE FROM *Xylanolytic Bacteria* INDUCED  
BY UTILIZING VINASSE WASTES**

KAHAR MUZAKHAR

Department of Biology, Faculty of Mathematics and Natural Sciences, the University of  
Jember

Address: Jl. Kalimantan 37 Jember 68121 Indonesia, Email:kaharmzk@unej.ac.id

**ABSTRACT**

The production of xylanase by using vinasse waste as carbon source was studied. Isolate MK-20, identified as xylanolytic bacteria can grow aerobically on vinasse without any nutrients added. During the cultivation, this isolate released xylanase and capable produced xylose from hydrolyzation of oat spelt xylan. The optimum for xylanase production was obtained after 3 days cultivation at 30<sup>0</sup>C. The enzyme was purified and characterized to have molecular weight of 88.2 KDa, respectively. The enzyme stable in pH 4-8 and temperature below 55<sup>0</sup>C while optimum activity at pH 5 and temperature 50<sup>0</sup>C.

**Keywords:** xylanase, xylanolytic bacteria, vinasse

**INTRODUCTION**

Xylan residues are widely distributed in the form of hemicellulose such as xylan and arabinoxylan [1]. Xylan also found in the major component of the hemicellulose fraction of plant [2]. For industrial application, this simple sugar can be used as carbon source for fermentation in alcohol production [3, 4]. Some potential applications for xylan enzymes in some industries would require the use of enzymes worked in wide range of pH. The enzymes were mostly found in microorganisms especially in fungus and bacteria [3, 5].

The bacteria and other microbes have therefore attracted considerable attention as sources of xylan degrading enzyme. Up to now, there have been many studies of xylanase but only a few studies of the production by utilizing raw material of agriculture wastes. A huge material wastes vinasse was released during ethanol production from molase. This material rich in organic substance including hemicelluloses fiber. In this works, the production of xylanase by utilizing vinasse is expected to be an effective strategy to minimize environmental problem of wastes.

## MATERIAL AND METHODE

### Sampling and screening of isolates

Ten grams of sample soil from 5 polluted locations by vinasse at PT. PASA Ethanol Industry Jatiroto Indonesia were sampled and suspended to 0.9% of NaCl solution 100 ml in Erlenmeyer flask. The samples were diluted  $10^{-2}$  to  $10^{-8}$  from initial concentration. For screening, one hundred micro liters of each diluted sample was plated to NA medium and incubated for 48 hours at  $30^{\circ}\text{C}$ . This step was repeated until getting the single colony of some different type of microorganisms. Twenty four isolates of bacteria were successfully screened. All isolates were further pre-culture in the same medium for 24 hours at  $30^{\circ}\text{C}$ .

### Optimization of xylanase production

The xylanase production was optimized by cultivation of pre-culture xylanolytic bacteria MK-20 in 100 ml vinasse. Under sterile condition, 1 ml of culture was harvested everyday and the xylanase activity was measured to obtain optimum cultivation.

### Enzyme assay

The xylanase activity was determined by measuring of reducing sugar released using the method of Nelson [6] as modified by Somogy Nelson [7]. For this measurement, 1% xylan on 50 mM buffers was used as substrate.

### Gas chromatograph analysis of xylose production

The quantity of xylose production during xylan hydrolysis is analyzed by using Gas Chromatograph (GC) as alditol acetates [8; Hondmann *et al.*, 1994) with a few

modification. The sample was hydrolyzed with 2N HCL for 6 hours, reduced with equal amount of NaBH<sub>4</sub> at room temperature overnight and then a batch of Dowex resin H type was added to the mixture followed by filtration. The filtrate was evaporated to dryness and residual boric acid was removed by repeated evaporation with methanol. The sugar alcohol was acetylated in 2 ml of acetic acid anhydride:pyridine (1:1) at 100<sup>0</sup>C for 10 min. The mixture was diluted with chloroform:water (1:4), shaken well and the supernatant removed by centrifugation at 2000 rpm for 10 minutes. Remaining pyridine was removed by washing with water by removal of aqueous phase after centrifugation. The alditol acetates of sugar was dried and dissolved in chloroform to an appropriate volume. The GC analysis was performed on stainless column, 2mm I.D. □ 1.83 m, packed with 3%(w/w) ECNSS-M on Gas Chrom Q, 100-120 Mesh. The initial column temperature was 190<sup>0</sup>C for 5 minutes and then using gradient up to 210<sup>0</sup>C at 1<sup>0</sup>C/min.

#### Effects of pH and temperature on the enzyme activity and stability

Analyzing the stability of the xylanase at different pH values, the purified enzyme was incubated at 37<sup>0</sup>C in 30 mM buffers Na-acetate, pH 3 to 5; Na-citrate, pH 6 to 6.5; and Tris/HCl, pH 7 to 9. The remaining activity of each sample was measured after 30 min of incubation. For measuring its thermal stability, the purified xylanase was incubated in 30 mM Na-acetate, pH 5, for 30 min, at 25 to 70<sup>0</sup>C. Remaining activity was measured after 30 min of incubation. The effects of pH and temperature on the optimum activity of enzyme were also measured in series pH and temperature range as mentioned above, using 1% substrate with 10 minutes incubation at 37<sup>0</sup>C.

#### Purification of enzymes

All purification steps were carried out at room temperature using 20 mM acetate buffer pH 5. Detail of this experiment was described under result and discussion.

#### Molecular weight analysis

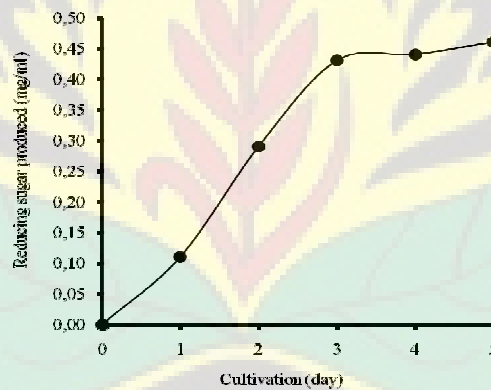
The molecular weight of enzyme was estimated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). Molecular weights of the denatured enzymes were estimated by comparison of their migration rates with those of protein



standards. Proteins Standard (KDa) used were Phosphorylase (97,4), Albumin (66,3), Aldolase (42,4), Carbonic anhydrase (30) and Trypsin inhibitor (20,1). Gel was stained with Coomassie Brilliant Blue.

## RESULTS AND DISCUSSION

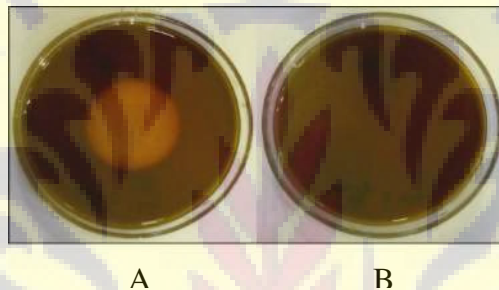
Among 24 isolates, bacteria identified as MK-20 can grow well on vinasse as carbon source although no other substances were added. This bacteria secreted an extracellular xylanase enzyme when aerobically cultivation was done under 30<sup>0</sup>C 120 rpm. The optimum cultivation was achieved in 3-4 days incubation (Figure



**Figure 1.** Aerobic cultivation of MK-20 using vinasse as carbon and nitrogen source at 30<sup>0</sup>C 120rpm. Optimum cultivation was achieved at 3 days cultivation.

1). When they grown 3 days in broth medium containing 1% xylan, 0.5% pepton and 0.5 malt extract, the reducing sugar released almost the same (0.45 mg/ml) comparing with vinasse medium (0.44 mg/ml). It seems that hydrolysis occurred and suggested vinasse medium was suitable enough to provide the carbon and nitrogen sources for MK-20 as well.

Further analysis for xylanolytic activity, MK-20 was tested using a xylan solid medium. Five microlitres of overnight grown culture of MK-20 was spot plated on xylan agar (0.2% NaNO<sub>3</sub>, 0.1% K<sub>2</sub>HPO<sub>4</sub>, 0.05% MgSO<sub>4</sub>, 0.05% KCl, 0.2% oat spelt xylan, 0.02% peptone, 1.7% agar, adjusted to pH 7) and incubated at 30°C. After 72 hours incubation, the medium plate was then flooded with Gram's iodine (2.0 gKI and 1.0 g iodine in 300 ml de-ionized water) within 3 to 5 minutes [10]. It is shown in Figure 2., clear zone within colony indicated that MK-20 released xylanase and actively hydrolysed xylan medium during cultivation, respectively.



**Figure 2.** MK-20 was spot plated on xylan agar (0.2% NaNO<sub>3</sub>, 0.1% K<sub>2</sub>HPO<sub>4</sub>, 0.05% MgSO<sub>4</sub>, 0.05% KCl, 0.2% oat spelt xylan, 0.02% peptone, 1.7% agar, adjusted to pH 7) and incubated at 30°C for 72 hours. The clear zone within colony (A) indicated the hydrolysis of xylan occurred while at control (B) no hydrolysis activity was detected.

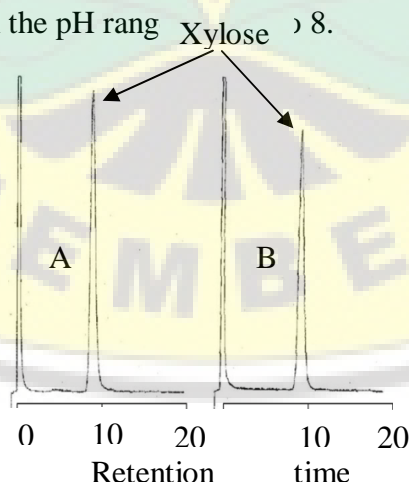
Crude xylanase obtained from optimum cultivation was collected and purified. For the first step, the crude enzyme was brought to 60% of saturated ammonium sulfate, precipitated by centrifugation at 12000 rpm for 20 minutes, and followed by dialysis on cellulose tube (10KDa pores) using 20 mM acetate buffer at pH 5 to remove remaining ammonium sulfate. After dialysis, the concentrated enzyme was loaded onto open column containing DEAE cellulose pre-equilibrated with the same buffer. For the fractionation, the column was eluted using same buffer with the gradient 0-0.5 M NaCl. The active fractions were pooled, reloaded onto DEAE Q-Sepharose and eluted with the same gradient NaCl.

Finally, a size exclusion chromatography Sephadex G-100 was used to obtain purified xylanase, resulting in 11% yield and 898 folds. The result of purification was summarized at **Table 1** as follow.

**Table 1.** Purification of xylanase

Purification step	Total ABS-280	Total Activity (unit)	Specific Activity	Yield (%)	Fold
Ammonium sulfate Precipitation	79,400	2,300	0	100	1
DEAE Cellulose	3,200	2,110	1	92	23
DEAE Q Sepharose	134	1,268	9	55	327
DEAE Sephadex G-100	9	234	26	11	898

The effect of temperature and pH on the optimum activity and stability were examined. The optimum activity for this enzyme was found at temperature 50°C and stability below 55°C. The optimum pH for activity for this enzyme was found at pH 5. The enzyme showed 95% activity in the pH rang Xylose > 8.



**Figure 3.** GC analysis of standard xylose (A) as control and hydrolyzate (B) after 120 min. incubation

To affirm this xylanase attacked the xylan and produced xylose, qualitatively and quantitatively analysis using GC was used. For this purpose, 20 ml of mixtures containing 20 units of xylanase and 1% oat spelt xylan were incubated under optimum temperature and pH for 4 hours. Every 30 minutes of 1 ml of hydrolyzates was sampled and the sugar directly converted to alditol acetate form using method as mentioned at material and method. The acetylated samples were then injected to GC for analysis. Pure xylose was used as standard. We found that the xylan from oat spelt was readily hydrolyzed by xylanase and released xylose (Figure 3). In a period 30 min to 60 min incubation, the hydrolyzation was exponentially occurred and GC analysis revealed of hydrolysis products are 32 and 64mg/ml of xylose produced. After 90 to 120 min incubation, the production of monosaccharide as reducing sugar xylose still increased 72 and 84mg/ml. However, after 180 to 240 min incubation, only 86 and 87mg/ml of xylose were produced. Suggested the hydrolysis was optimum in 120 min incubation where about 85% of xylan was hydrolysed.



**Figure 4.** SDS-PAGE of the purified xylanase. Lane 1 is protein standards (KDa) and lane 2 is purified xylanase.

Molecular weight of the enzyme was estimated by SDS-PAGE. This xylanase was 88.2 KDa approximately

## CONCLUSION

Inducible of xylanase from isolate MK-20 xylanolytic bacteria was done successfully by employing vinasse only as carbon and nitrogen source. The enzyme was purified with 11% yield and 898 folds purification. The enzyme was purified and characterized to have molecular weight of 88.2 KDa, respectively. The enzyme stable in pH 4-8 and temperature below 55<sup>0</sup>C while optimum activity at pH 5 and temperature 50<sup>0</sup>C.

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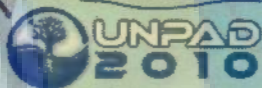
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Phone : +62 22 7279435

Fax : +62 22 7279435

e-mail : [puslitbiotek@yahoo.com](mailto:puslitbiotek@yahoo.com)

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