In Vitro and In Silico Study of Immobilized Xylanase from Trichoderma viride using Bentonite Matrices Activated with HCl

Nur Lailah, Sasangka Prasetyawan, and Arie Srihardyastutie*

Chemistry Department, Faculty of Mathematics and Natural Sciences, University of Brawijaya, Jl. Veteran 65145 Malang, Indonesia

*Corresponding author : arie_s@ub.ac.id

Received 28 March 2017; Revised 1 May 2017; Accepted 2 May 2017

ABSTRACT

The purpose of this study was aimed to determine the optimum conditions of agitation time and concentration of immobilized xylanase by in vitro and in silico and efficient reuse of immobilized xylanase. The results of in silico test showed that xylan bounded on the residues of xylanase active site through hydrogen bonding with the amino acid glutamic 177A, glutamate 86A, tyrosine 88A, arginine 122A, and glutamine 136A, whereas bentonite and the residues of xylanase active site form hydrogen bonding with the amino acid glutamine 52, arginine 81, isoleucine 128 and glycine 130. The results of in vitro study showed that the optimum condition of xylanase immobilization was achieved at agitation time of 3 hours with the amount of xylanase adsorbed was 12.593 mg/g bentonites and activity of 50.328 units and the concentration of immobilized xylanase of 4.259 ppm and the amount of xylanase adsorbed of 16.162 mg/g bentonites and activity of 56.362 units. The immobilized xylanase can be used as many as five repetitions with the residual enzyme activity of 60%.

Key word: xylanase, Trichoderma viride, immobilization, bentonite, molecular docking

INTRODUCTION

One of the corn waste which can be used is corn husk that containing lignocellulose. The chemical composition of the corn husk consisting of cellulose 42.3%, ash 4.16%, lignin 12.58%, and others 40.95% [1]. The waste containing lignocellulose with the aid of microorganisms can produce extracellular enzymes which capable of degrading materials containing lignocellulose into its constituent fractions. One of the extracellular enzymes produced is xylanase which can hydrolyze xylan (hemicellulose) into xylose. The use of xylanase enzyme in the hydrolysis process in the industry is a new prospect in handling the hemicellulose waste. Microorganisms such as fungi, bacteria, and yeast can produce xylanase enzyme. One of the microorganisms that produces xylanase enzyme is Trichoderma viride [2].

Xylanase has biotechnological potential in many industrial processes, such as in ethanol production [3], in paper industry [4], food industry [5], textile processes [6], and organic waste treatment [7]. The utilization of enzymes in industry has many obstacles because the enzyme is easily denatured by several factors, including concentration of the substrate, an activator, a specific inhibitor of certain enzymes, salt, buffer, pH, ionic strength, temperature and alkali [8]. Therefore, the stability of the enzyme needs to be improved using the immobilization techniques [9]. The immobilized enzyme has several advantages, i.e. the
enzyme can be reused which reduce the operating cost and increase the stability of the immobilized enzyme. The selection of immobilization technique is based on the matrix used.

Bentonite matrix has the high ion exchange capacity, the ability to expand with a layered structure and has cations which can be exchanged. The adsorption method of binding on the carrier physically has the advantage that the enzyme is not denatured and media can be reused. The capacity of bentonite adsorption is limited. These limitations can be overcome through the activation process by acid including HNO₃, H₂SO₄ and HCl [10]. The activation of bentonite with acid will produce active sites and has a larger surface acidity resulting bentonite with higher adsorption capacity than bentonite which not activated [9]. In the current research, xylanase enzyme will be immobilized on bentonite matrices that have been activated using HCl, in order to reduce its alumina content.

Sari, Sutrisno, and Prasetyawan (2014) reported the results of the optimization of xylanase immobilization from Trichoderma viride using zeolite matrix activated with HCl with physical adsorption method. The optimum time was 3 hours with the amount of xylanase adsorbed was 7.370 mg and the activity of xylanase immobilized of 22.67 units and the optimum concentrations of immobilized xylanase of 3.366 mg/mL with amount of xylanase adsorbed of 15.65 mg/g with xylanase activity of 25.74 units. The immobilized enzyme can be used as many as five times with efficiency of 51.58 % [11].

In physical adsorption, the agitation is very influential because agitation times and enzyme concentrations affect the adsorption rate and the mass of enzyme adsorbed therefore will affect the activity of the immobilized enzyme. Furthermore, in silico study is necessary to perform in order to determine the level of the complex structure stability and strength of the bonding between enzymes, substrate and matrix (bentonite) by simulating the interaction between the active site of the receptor (enzyme), the ligand (substrate) and the matrix. The simulation of molecular interaction using in silico test (Molecular Docking) is performed optimally which can be used to determine the strength level of the complex bonding formed which can determine the enzyme activity. Therefore, it is important to do research on study of in vitro and in silico on xylanase immobilization of Trichoderma viride using bentonite matrices activated with HCl.

EXPERIMENT
Chemicals and instrumentation

All reagents were purchased from Merck, Sigma, or Smart Lab. The materials used in this study include: Trichoderma viride, dextrose, oleic acid, iron(III) chloride (FeCl₃), calcium chloride (CaCl₂), boric acid (H₃BO₃), zinc sulfate (ZnSO₄), copper(II) sulfate pentahydrate (CuSO₄·5H₂O), potassium sodium tartrate (KNA-tartrate), barium chloride (BaCl₂), ammonium sulfate ((NH₄)₂SO₄), magnesium sulfate heptahydrate (MgSO₄·7H₂O), potassium dihydrogen phosphate (KH₂PO₄), sodium hydrogen phosphate (Na₂HPO₄), glucose anhydrous, hydrochloric acid (HCl), sodium chloride (NaCl), potassium hydroxide (KOH), nitric acid (HNO₃), ammonium chloride (NH₄Cl), potassium hydrogen phosphate (K₂HPO₄), dinitrosalilat acid (DNS), acetic acid (CH₃COOH) 1.05 g/mL, sodium acetate (CH₃COONa), and natural bentonite.

The instrumentation used are refrigerator, laminar flow, autoclave (All American Model 20X), shaker (Edmund Buhler SM 25, 24B), incubator, cold centrifuge, pH meter (WTW inoLab), pH paper, spectrophotometer UV-Vis (Shimadzu Model 160A double-beam), HyperChem, Pyrex-Vinawizard8.0, PyMOL, and patchdock software.
Production and isolation of xylanase

Liquid media (65 mL) was taken with pipette and transferred into erlenmeyer and was added using 25 g of substrate and was sterilized by autoclaving at pressure of 15 psi and temperature of 121 °C for 15 minutes. Furthermore, the inoculum (10 mL) was added into solution and was performed in laminar air flow. The mixture solution was incubated for 60 hours at room temperature and was shaken using shaker at 100 rpm. After that, solution of acetate buffer pH 5 (30 mL) was added into mixture solution and then was centrifuged in cold conditions at speed of 3000 rpm for 30 minutes at 4 °C. Supernatant containing xylanase crude extract was purified by 40-80% (NH4)2SO4 and was dialyzed using a semi-permeable membrane pouch (cellophane). The crude extract of purification was tested for its enzyme activities and protein contents.

Determination of optimum agitation time

The xylanase purification process (2 mL) was taken with pipette and then was added with acetate buffer of pH 5 until volume of 5 mL and was transferred in erlenmeyer containing bentonite (0.1 g) which had been activated. Each mixture was incubated in a shaker at 100 rpm at room temperature for 1, 2, 3, 4, and 5 hours and was filtered with Whatman paper no. 40 to separate between filtrate and immobilized xylanase. The sediment was tested for its enzyme activity, and filtrate was tested for its protein content.

Determination of optimum xylanase concentration

The immobilization procedures at various xylanase concentrations were similar to the immobilization procedures at various agitation times. The difference was on the additional of xylanase volume used (2.5, 3, 3.5, 4 and 4.5 mL) to obtain 5 enzyme concentrations (3.549, 4.259, 4.969, 5.679, and 6.389 mg/mL).

Determination of activity of immobilized xylanase

Determination of activity of immobilized xylanase was performed with 1% xylan substrate (0.2 mL) which heated at temperature of 60 °C for 15 minutes and was added using immobilized xylanase (0.1 grams), water-free reductant (0.2 mL), and acetate buffer pH 5(0.2 mL). The solution mixture was incubated at 50 °C for 55 minutes, and then was heated for 15 minutes in boiling water. The mixture was filtered and the precipitate obtained was added by DNS reagent (0.4 mL) and was heated in boiling water for 15 minutes. After that, the solution mixture was diluted using distilled water until volume 10 mL and was measured the absorbance at the maximum wavelength.

Determination of reuse efficiency of xylanase

The efficiency test was conducted on immobilized enzymes resulted in the optimum conditions of agitation times and xylanase concentrations. The immobilized xylanase (0.1 gram) was inserted into 1% (w/v) of xylan substrate solution (1 mL) which incubated at 50 °C for 15 minutes. The mixture solution was added acetate buffer pH 5 (1 mL) and distilled water (1 mL) and was incubated at 50 °C for 55 minutes. The mixture solution was filtered using Whatman filter paper no. 40 and the filtrate obtained was added by DNS reagent and was heated in boiling water bath for 5 minutes and was cooled at room temperature and was measured the absorbance as enzyme activity. The precipitate of immobilized xylanase obtained was added on xylan substrate 1% (w/v) which has been carried incubation at 50 °C for 15 minutes and was continued as in the previous procedure.
Molecular Docking

*PyRex* and *PatchDock* program were used to visualize interaction among xylanase, bentonite and xylan. Protein and ligand were downloaded from the website protein data banks (pdb) and were inserted into *Pyrex-Autodock-Vina* 8.0 program. In this study, a rigid docking protocol was considered in order to reduce the computational cost and time in which the receptor was kept rigid and ligands were allowed to rotate. The size of the grid box was set to 25 Å x 25 Å x 25 Å (x, y and z). The docking results were saved after the table has emerged which consists of bond energy or binding affinity and RMSD values. PyMOL program was used to view complex in 3D. Docking begins with uploading the receptor and ligand molecule.

RESULT AND DISCUSSION

Isolation and purification of xylanase from *Trichoderma viride*

Isolation result of pure extract of *Trichoderma viride* was dark yellow. Table 1 and 2 show the results of enzyme activity and protein content tests of free xylanase.

**Table 1. Activity of free xylanase**

<table>
<thead>
<tr>
<th>Type</th>
<th>Absorbance (nm)</th>
<th>[Reducing sugar] (mg/mL)</th>
<th>AE(µg/mL.mnt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude extract</td>
<td>0.307 0.307 0.307 0.307</td>
<td>19.188</td>
<td>7.274</td>
</tr>
<tr>
<td>Pure extract</td>
<td>0.698 0.714 0.712 0.708</td>
<td>44.250</td>
<td>16.776</td>
</tr>
</tbody>
</table>

**Table 2. Protein content of free xylanase**

<table>
<thead>
<tr>
<th>Type</th>
<th>Absorbance (nm)</th>
<th>Protein concentration (mg/mL)</th>
<th>Protein content (mg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure extract</td>
<td>0.34 0.343 0.342 0.342</td>
<td>11.389</td>
<td>6.389</td>
</tr>
</tbody>
</table>

Based on Table 1, the xylanase of purification result had specific activity of two times higher than the crude extracts. Based on Table 2, it can be seen that the protein content of pure extract is equal to 6.389 mg/mL. By in vitro, xylan will be bounded on xylanase active side of aspartic acid and glutamic acid. Then, xilan will be converted into xylose. The type of bond that happened between xylan and xylanase can by tested by in silico (molecular docking) using *Autodock-Vinawizard* 8.0 program. The xylanase protein receptor is downloaded from the bank data protein (PDB) with 3AKT code and xylan ligand with 1KNM code.

The results of docking calculation between the protein and ligand provided the valid result because it had value of ≤ 2 Å [12]. Based on the data obtained, xylan bounded to the active site of xylanase amino acid glutamate 177A, glutamate 86A, tyrosine 88, arginine 122A, and glutamine 136A, asparagine 44A which can improve the structure of the xylanase for inhibition constants. Results of docking between the xylanase and xylan can be seen in Figure 1. Hydrogen bond formed through O atoms derived from xylan and H atoms derived from glutamic amino acids. The hydrogen bond was showed by dashed lines and interaction can be seen in Figure 2.
**Figure 1.** Interactions between xylanase with xylan
Note: xylanase (■), amino acids ( ), xylan ( ■ ), hydrogen bonding ( --- )

RMSD 0,0 Å; $\Delta G^\circ$-7 kcal/mol

**Figure 2.** The interaction model of complex xylanase enzyme molecules with xylan ligands
Note: Glu, Tyr, Arg, Gln are abbreviation of amino acids

**Determination of optimum condition of time agitation and enzyme concentration on immobilization**

In this study was investigated the effect of time agitation to the amount of adsorbed enzyme and enzyme activity. The time agitation was performed on the variation of 1, 2, 3, 4, 5 hours. Based on this research, it was known that the optimum agitation time was 3 hours and amount of xylanase was 12.593 mg.
Figure 3 (A) show that the amount of adsorbed enzyme was affected by time agitation. On the addition of time agitation from 1 to 3 hours, the amount of adsorbed enzyme increased significantly. Increase the agitation time cause higher interaction between the bentonite and xylanase. This is because xylanase can be attached to bentonite surface longer. In contrast, the addition of 4 to 5 hours caused decline in the amount of adsorbed xylanase significantly. This is possible due to the desorption process, therefore, the amount of xylanase adsorbed decreased.

Based on Figure 3 (B), enzyme activity increased on agitation time 1 to 3 hours. The optimum enzyme activity was at time of 3 hours, which amounted to 50.328 units. The increase in the activity of enzymes associated with the amount of more substrate binding to xylan xylanase. This was because the amount of adsorbed xylanase on bentonite increased so that its activity was also higher.

The optimum of agitation time of 3 hours was used to study the effect of enzyme concentration on the amount of absorbed enzyme and enzyme activity. Determination of optimum concentrations was performed in five variations of the enzyme with a mass concentration of 0.1 g bentonite. Figure 3 shows the results of determination of the optimum xylanase concentrations.
Figure 4. Influence of enzyme concentration on the amount of adsorbed enzyme (A) and enzyme activity (B)

Based on the Figure 4 (A), it can be seen that increasing the concentration of xylanase caused increasing the amount of adsorbed xylanase. The optimum xylanase concentration was 4.259 ppm (3 mL) with amount adsorbed xylanase of 16.162 mg/g of bentonite. The increase was due to the difference in chemical potential on the surface of the bentonite matrix which caused high diffusion rate therefore the mass of adsorbed xylanase increased. As a result, decrease in the amount of adsorbed xylanase on the surface of the matrices because bentonite has been saturated with xylanase, thus bentonite is no longer adsorbing xylanase.

Based on the Figure 4 (B), it can be seen that increasing of enzyme concentration from 3.549 to 4.259 ppm, and decreasing was occurred at concentration of 4.969, 5.679, and 6.389 ppm. The xylanase activity was shown on the optimum enzyme concentration of 4.259 ppm, with activity of 56.362 units. The decreasing that occurred was due to the mass desorption of absorbed xylanase. This is due to the greater amount of adsorbed xylanase so that the substrate is able to bind to the active site of the xylanase greater.

Based on the value of enzyme activity, free xylanase had enzyme activity which was much smaller when compared to the immobilized enzyme. Basically, the activity of immobilized enzyme should have lower than free enzyme. These activities can be learned through the study of using in silico molecular docking to understand the bond between xylan-xylanase and bentonite matrix, with regards of the effect of agitation times and the enzyme concentrations. Based on in silico test, the interaction between the active site of xylanase and bentonite can be illustrated in Figure 5.
Figure 5. Interaction between the bentonite and the side active of xylanase

Figure 6. Interactions between xylanase, bentonite and xylan

Figure 5 showed that bentonite form hydrogen bonding with the active site residues included amino acid glutamine 52, arginine 81, glycine isoleucine 128 and 130. Values of $\Delta G^\circ$ was -35.89 kcal/mol and Van Der Waals style was -16.42 kcal / mol. Very low $\Delta G^\circ$ value indicated that the xylanase-bentonite complex was very strong. Value of $\Delta G^\circ$ was minus, therefore the complex run spontaneously. The binding between xylanase and bentonite occur by physical adsorption based on the Van Der Waals forces. The physically adsorbed molecules are not strong bonded on the surface, and usually there was a reversible process. Therefore, it was easy to replace with other molecules [13].

Figure 6 showed that bentonite and xylan was interconnected to form hydrogen bonding through the O atom derived from xylan and O atom derived from SiO$_2$ and bentonite AlO$_2$ from bentonite. Overall, xylan and bentonite bounded to the different active siteS so can improve the structure of the xylanase, and can increase the activity of the immobilized enzyme. The activity of immobilized enzyme was much greater than free enzyme because the activation of matrix with HCl the ratio of Si/Al increased. Therefore, more xylanase enzymes
were adsorbed and the formation of enzyme-substrate complexes also increased. The docking results show that the bond between bentonite and xylanase via the O atoms of SiO$_2$ which form hydrogen bond with several active sides of amino acids on the xylanase and via the O atoms of Al$_2$O$_3$. The interaction of hydrogen bonding between molecules was strong enough.

**Determination of reusage efficiency of xylanase immobilized**

Determination of reuse of immobilized xylanase was performed at the optimum condition which obtained from previous treatment by using the optimum time agitation for 3 hours and the optimum concentration of 4.259 ppm. This treatment was conducted to determine how much the immobilized enzyme that can get known to test the feasibility of xylanase reuse. The results of xylanase re-used were shown in Table 3. Based on Table 3, it can be seen that the enzyme activity on the reuse efficiency decreased after the first use. The reason was possible due to the strength of the bentonite matrix reduced which caused the enzyme easily detaching and can caused by the Van der Waals forces and ionic bonds between hydrogen ion of bentonite and RNH$_3^+$ of the enzyme.

**Table 3.** Effect of reuse to the activity of enzyme immobilized in optimum condition

<table>
<thead>
<tr>
<th>Repetition</th>
<th>Enzyme activity</th>
<th>Efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>46.063</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>44.704</td>
<td>97</td>
</tr>
<tr>
<td>3</td>
<td>42.809</td>
<td>96</td>
</tr>
<tr>
<td>4</td>
<td>35.542</td>
<td>83</td>
</tr>
<tr>
<td>5</td>
<td>21.294</td>
<td>60</td>
</tr>
<tr>
<td>6</td>
<td>6.761</td>
<td>32</td>
</tr>
</tbody>
</table>

The result in the initial protein content measurement which re-use efficiency was 0.656 mg/mL and residual protein content from the second to the last repetition was 5.167 mg/mL. This indicates that the more detached protein contents caused less bonding between the enzyme and the substrate contained in the surface of the bentonite matrix and caused the enzyme activity in reuse decreases. Based on the results of study, the immobilized enzyme using bentonite matrix activated HCl can be used as many as 5 times with efficiency of 60%. The immobilized enzyme was effectively used when its activity was still above 50%.

**CONCLUSION**

The optimum agitation time in the process of xylanase immobilization using the bentonite matrix which activated using 0.4 M HCl was obtained for 3 h, with amount of xylanase adsorbed was 12.593 mg/g of bentonite and enzyme activity of 50.328 units. The optimum concentration of enzyme immobilization of xylanase was obtained at concentration of 4.259 ppm and the amount of adsorbed xylanase of 16.162 mg/g of bentonite and xylanase activity of 56.362 units. Based on **in silico** test, it can be concluded that the bond between the bentonite and the residues of the active sides of xylanase can improve the xylanase structure which caused the activity of immobilized xylanase was much higher than pure free enzyme. The results of reuse efficiency of xylanase was performed on the optimum condition of time agitations and enzyme concentrations were 5 times repetitions with 60% and the percentage of residual protein content of 5.167 mg/L.
REFERENCES