Expression and characterization of *Trichoderma reesei* endoglucanase II in *Pichia pastoris* under the regulation of the GAP promoter

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ABSTRACT *Trichoderma reesei* is known to be one of the organisms capable for producing various types of cellulase in high concentrations. Among these cellulases, the highest catalytic efficiency of endoglucanases II (EGII, EC 3.2.1.4) are considered important for industrial application. The characterization of the EGII is necessary since it is widely used in high-temperature reactions in the industries. In this study, the recombinant EGII protein was expressed in *Pichia pastoris* and it has a molecular mass of approximately 52 kDa. Recombinant EGII was purified using Ni-NTA affinity chromatography and characterized by SDS-PAGE and western blot analyses. The enzyme activity of recombinant EGII was measured using the Nelson Somogyi method to determine its optimum pH and temperature. The result showed that the maximum EGII expression was achieved after 72 h of culture incubation. The crude enzyme has optimum activity at pH 5.0, resulting in 16.3 U/mL and 14.6 U/mL activity at 40 °C and 50 °C, respectively. While the purified enzyme gave the specific activity of 115.7 U/mg under the optimum condition. Finally, our study demonstrated that recombinant EGII could retain the endoglucanase activity for 89% and 80% at 40 °C and 50 °C, respectively.

KEYWORDS Endoglucanase II; GAP promoter; *Trichoderma reesei*; *Pichia pastoris*; Nelson-Somogyi assay

1. Introduction

*Trichoderma reesei* is a well-studied fungus that is capable of producing large amounts of various cellulases. This fungus secretes at least six types of cellulases that consist of two types of cellobiohydrolases and four endoglucanases (Knott et al. 2014). These enzymes are extensively used in several industries, such as laundry detergent, textile and pulp, paper industry, and potential for bioenergy production. Thereby, this fungus is industrially relevant to meet the target production level of cellulase production. Among cellulases produced by *T. reesei*, endoglucanase II (EGII; EC 3.2.1.4) is predominant and showed the highest catalytic proficiency. The EGs activity of *T. reesei* is known to decrease about 55% when EGH was absent in the secretory complex of EGs (Qin et al. 2008; Boonvithya et al. 2013). This evidence revealed that the presence of EGII is crucial for lignocellulosic biomass hydrolysis and other industrial applications. Several studies reported that EGII production and characteristic improvement were performed by improving the strain of secreting microorganisms, protein engineering, and recombination (Ito et al. 2004; Liang et al. 2011; Charoenrat et al. 2013). Thereby, there is still ample scope for improvement, particularly to produce EGII in a heterologous expression system to facilitate protein engineering work. Several heterologous expression has been carried out to produce endoglucanases in various host microorganisms, including *Escherichia coli*, *Yarrowia lipolytica*, *Saccharomyces cerevisiae*, and *Pichia pastoris* (Nakazawa et al. 2008; Qin et al. 2008; Boonvithya et al. 2013; Akbarzadeh et al. 2014). Yeast is commonly used for its ability to increase protein stability since the glycosylation process has occurred. In consequence, the structural and thermal stability of protein may increase due to the covalent bond formation. The covalent bond formation causes less dynamic fluctuation and reduces protein molecules’ flexibility (Qin and Qu 2014).

*Pichia pastoris* has negligible native protein produc-
tion levels, which helps for easier purification of recombinant protein (Macauley-Patrick et al. 2005). It has also been proven that the expression system promotes economically effective production of recombinant protein as it does not need complex medium and condition (Safder et al. 2018). The expression of a foreign gene in P. pastoris includes three main steps: (a) insertion of the foreign gene into an expression vector; (b) introduction of the expression vector into the expression host, P. pastoris; and (c) the selection of potential strains for foreign gene expression (Macauley-Patrick et al. 2005). The gene encoding EGII (egII gene) from T. reesei was successfully inserted into an expression vector and introduced into P. pastoris genome. In this study, the high-expression transformant with high endoglucanase activity was selected for further steps. The expression of recombinant EGII was regulated under glyceraldehyde-3-phosphate dehydrogenase (GAP) constitutive promoter in the fed-batch fermentation process. Recombinant EGII produced was then characterized for the determination of its optimum pH, temperature, and thermal stability.

2. Materials and Methods

2.1. Strain and plasmid

Pichia pastoris SMD1168H purchased from Invitrogen (USA) was used as an expression host. Constructed plasmid pLIP1-1TrCel5A ordered at ATUM (USA) used as an expression vector for P. pastoris transformant carrying egII gene from T. reesei was available at Research Center for Biotechnology, Indonesian Institute of Sciences.

2.2. Colony selection

Pichia pastoris was streaked from glycerol stock was streaked in YPD agar (1% w/v yeast extract, 2% w/v peptone, 2% w/v dextrose, 1.8% bacteriological agar) medium containing zeocin (500 µg/mL) and ampicillin (100 µg/mL) antibiotics (Invitrogen, USA) and incubated at 28 °C for 36 h. The colonies were selected from the selection medium and characterized by colony PCR. Specific primers, namely MFe-FP1 (5’-ATGAGATTCCCATCTATTTTCACCGCTGTCT-3’) and TrCel5A-RP (5’-GAGCGGGGGATATCCTTGGGAAGTACACAA-3’), were used to detect the inserted gene in the yeast genome. PCR was performed as follows: initial denaturation at 95 °C for 5 min; 30 cycles of denaturation at 95 °C for 30 s, annealing at 55 °C for 40 s and extension at 72 °C for 40 s, then final extension at 72 °C for 5 min. The amplified fragments were then analyzed using 1% agarose gel electrophoresis.

The colonies were then confirmed for their expression of EGII, measured by a plate diffusion assay according to the method proposed by Ratnakomala et al. (2019). Ten microlites of each transformant cultures were added into agar wells containing 0.5% (w/v) carboxyl methylcellulose (CMC) and incubated for 3 d at 30 °C. Finally, plates were stained using 1% Congo Red solution for 15 min for color development, followed by washing the plate with 1 M sodium chloride solution to detect halo zones. The diameter of the halo zones was measured and documented. Transformants with bigger halo zones were selected for further analysis.

Selected transformants were tested for their endoglucanase activity. The activity was measured by Nelson-Somogyi (NS) method using CMC as a substrate, according to the reference by Gusakov et al. (2011). About 160 µL of 6.25 mg/mL CMC in 0.1 M acetate buffer (pH 5.0) and 40 µL culture supernatant were preheated at 50 °C for 5 min. Both were mixed and heated at 50 °C for another 10 min. Zero-point two milliliters of copper tartrate was added into the mixture to stop the reaction, and then the assay mixture was incubated in boiling water for 40 min. The mixture was allowed to decrease the temperature to 25 °C (room temperature), then 0.2 mL of arsenomolybdate was added and incubated at room temperature for 10 min. One point four milliliters of a mixed solution and 0.4 mL of acetone was added into the assay mixture and then centrifuged at 13,000 rpm for 1 min. The optical density was measured using a spectrophotometer at λ=610 nm to estimate the quantity of reducing sugars produced in the assay mixture. One enzyme activity unit was defined as the amount of enzyme-producing one µmol of reducing sugar per min under assay conditions (Jin et al. 2011).

2.3. Culture condition of endoglucanase II expression in Pichia pastoris

The pre-culture of each selected colonies was prepared into 2 mL YPD medium containing 100 µg/mL ampicillin and 100 µg/mL zeocin, then incubated at 28 °C for 2 d. One milliliter of pre-culture was added into 19 mL YPD medium containing 100 µg/mL ampicillin without zeocin in an Erlenmeyer flask and incubated at 28 °C for 4 d at 250 rpm. Fed-batch culture, including sampling, was carried out every 24 h intervals by withdrawing 4 mL of culture supernatant, along with the addition of 4 mL of 5× YPD medium to return the initial medium volume. The batch culture was only involving sampling every 24 h intervals by withdrawing 4 mL culture supernatant without the addition of YPD media.

2.4. SDS-PAGE and Western Blot

Sample preparation for SDS-PAGE analysis was done following the reference by Koontz (2014). One milliliter of culture from sampling was centrifuged in 12,000 rpm to separate its pellet and supernatant. The sample supernatant was added with 150 µL of 100% TCA, then vortex to homogenize. The sample was incubated at 4 °C overnight to allow protein precipitation, then centrifuged at 12,000 rpm for 10 min to allow separation between supernatant and precipitated protein. The supernatant was removed without disrupting the pellet. The pellet was washed using 200 µL of acetone, followed by centrifugation at 12,000 for 10 min. The step of washing the pellet with acetone
was repeated twice to ensure no more TCA residue was left. Pellet was allowed to dry to remove the acetone, then added with 15 µL of 5× SDS-PAGE loading buffer and being heated for 10 min in boiling water. Samples then analyzed by 15% SDS-PAGE gel using 90 V for 80 min to run the samples. The molecular mass was estimated from the migration distance in comparison with the prestained protein molecular weight marker (Thermo Scientific, USA).

Western blot was performed by electrotransfer the bands from acrylamide gels onto a nitrocellulose membrane at 90 V for 2 h. The membrane was blocked using a 10 mL blocking agent containing 1% BSA, then incubated for 1 h with gentle shaking at room temperature. The 5 µL of KPL HisDetector Nickel-HRP (SeraCare, USA) was added directly into the block solution, and incubation was continued for 1 h with gentle shaking. The membrane was washed three times with TBS solution containing 0.05% v/v Tween-20, each time for 5 min. Finally, the detection was done by adding 5 mL KPL TMB to visualize bands of interest (Thermo Scientific, USA).

2.5. Purification of recombinant endoglucanase II

The cell-free medium culture was harvested using centrifugation after 72 h of incubation and purified manually using Ni-NTA sepharose affinity chromatography column (Thermo Fisher Scientific, Massachusetts, U.S.). One milliliter of column material was washed with a five-bed volume of 20% ethanol, then washed with aquadest with the same volume. The column was then equilibrated with a five-bed volume of 50 mM PBS buffer (pH 7.4). A three-bed volume of crude EGII sample was added and incubated at 4 °C for 2 h. The sample was allowed to flow through the column and collected in an Eppendorf tube, labeled as a flow-through fraction. The column was then being washed with a three-bed volume of 50 mM PBS pH 7.4. Finally, a three-bed volume of an elution buffer containing 250 mM of imidazole was added into the column. Each 250 µL was collected as one fraction in a 1.5 mL centrifuge tube. Then proteins from collected fractions were then analyzed using 15% SDS-PAGE gel to identify which fraction contained purified EGII. The protein concentration was estimated using a Bovine Serum Albumin (BSA) standard curve following Carter’s method (2013). The electropherogram of SDS-PAGE containing EGII proteins and a series of BSA with known concentration were subjected to ImageJ 1.53e software (Rasband, 1997-2018). By calculating the area under the curve (AUC) of the BSA protein band, the linear regression equation was made. The AUC of EGII was analyzed, and the value was entered into the equation to get the concentration of recombinant EGII.

2.6. Characterization of recombinant endoglucanase II

Characterization was done by following the NS method according to the reference by Bajaj et al. (2009). To determine pH’s effect on enzyme activity, various buffers including citrate, acetate, phosphate, tris, bicarbonate, and carbonate buffers with pH 3, 5, 6.8, 8, 9, and 10 respectively were used. To determine the optimum temperature of recombinant endoglucanase, the temperature used was varied (40 °C, 50 °C, 60 °C and 70 °C) under standard assay conditions. One percent of the CMC solution was used as a substrate to determine optimum pH and temperature. To determine the thermal stability of recombinant EGII, equal quantities of purified EGII were pre-incubated for 15, 30, 45, and 60 min at optimum temperature and pH, which already pre-determined. After pre-incubation, enzyme activities were measured using standard assay conditions. Enzyme activity can be calculated by using the formula as the following:

\[ \text{Enzyme activity (U/ml)} = \frac{\text{Enzyme concentration [mg/L]} \times \text{dilution factor} \times 1000}{\text{time (min)} \times \text{Molecular weight of substrate}} \]

Whereas enzyme-specific activity can be calculated by using the following formula:

\[ \text{Specific activity (U/mg)} = \frac{\text{Enzyme activity (U/ml)}}{\text{Protein concentration (mg/ml)}} \]

3. Results and Discussion

3.1. Selection of recombinant clones

A study conducted by (Sivashanmugam et al. 2009) suggested that long-time storage of glycerol stock utilized for pre-culture preparation often leads to low-yield protein expression, although the colony used was known to previously produce a high yield of the target protein. Therefore, colony selection should be re-conducted. In this study, we did colony selection from glycerol stock using colony PCR followed by enzyme activity and expression screening.

The recombinant clones from glycerol stock were spread on YPD plates containing zeocin (500 µg/mL) and ampicillin (100 µg/mL). Six random recombinant clones were selected for screening after incubation at 28 °C for 36 h. Colony PCR was performed using specific primers to detect the presence of the egII gene in transformant colonies. pLIPI-TrCel5A plasmid carrying egII gene was used as a template for positive control while P. pastoris SMD1168H nontransformed colony was used for negative control. The amplified products were visualized on an agarose gel as seen in Fig. S1, and were approximately 625 bp in length. The result ensures that all selected transformants were inserted with egII gene.

| Colony no. | Average of halo zones (cm) | Enzyme activity (U/ml) |
|------------|---------------------------|-----------------------|
| 1          | 2.98                      | 8.568                 |
| 2          | 2.65                      | 7.096                 |
| 3          | 2.62                      | 7.498                 |
| 4          | 2.65                      | 6.570                 |
| 5          | 0.92                      | 4.165                 |
| 6          | 0.77                      | 6.540                 |

TABLE 1 Average of halo zones and enzyme activity of clone 1-6.
Colony selection was then continued with a qualitative and quantitative enzymatic assay, namely plate diffusion and NS assay. In plate diffusion assay, clone selection was based on the ratio between well and halo zone diameter. Whereas in NS assay, clone selection was based on the activity of EGII to breakdown CMC and produce reducing sugars, by which the reducing sugars will react with the NS reagent. CMC was chosen as the substrate as it exhibits an amorph structure suitable for the hydrolysis mechanism by EGII (Biswas, 2014). This study showed that EGII was successfully expressed in all selected clones, as indicated by the halo zone formation. Clones 1-4 have a higher halo zone diameter, among others (Table 1 and Figure 1). Recombinant clones were then selected by determining their enzyme activities using the NS method (Table 1 and Figure 2). From those analyses, clones 1-3 showed higher endoglucanase activity compared to other colonies.

3.2. Heterologous expression of Endoglucanase II by recombinant clones in YPD media

The expression profile of recombinant protein produced by six recombinant clones in YPD media was analyzed through 15% SDS-PAGE gel after 96 h of incubation, including sampling every 24 h intervals (Supplementary Figure 2). Clone 2 was chosen for further expression analysis as it showed the thickest band of interest at a size of around 50 kDa with lesser non-target bands. Moreover, clone 2 also showed a high enzymatic activity, which refers to plate diffusion and NS assay (Figure 1 and Figure 2). Clone 2 was then used for recombinant EGII production through fed-batch and batch fermentation (Figure 3). The production of recombinant EGII through fed-batch and batch fermentation was done to compare both methods and know which method is less time-consuming and produce a higher level of the target protein. Feeding was included in fed-batch fermentation by adding a new 5×YPD medium right after sampling, whereas batch fermentation only included sampling. The maximum number of cells achieved through fed-batch fermentation was 3.76×10⁷ cells/mL after 72 h of incubation. Meanwhile, the maximum number of cells achieved through batch fermentation was only 2.71×10⁷ cells/mL after 48 h of incubation. The protein expression was done through fed-batch fermentation (Figure 4a) showed a thicker band of interest in each 24 h intervals compared to batch fermentation (data not shown). More protein bands were present in the protein expression profile from batch fermentation with less than 50 kDa in size. Macauley-Patrick et al. (2005) suggest that the secreted recombinant proteins can be proteolytically degraded in the culture medium. This might happen due to cell-bound proteases, extracellular proteases, and/or intracellular proteases from lysed cells. The problems due to proteolysis can be foreseen in the recombinant protein production: (a) reduction of product yield when product is degraded, and (b) reducing biological activity when the product is truncated. In this study, protease production may occur due to the insufficiency of nutrients that lead to cell lysis. In comparison, fed-batch fermentation supplied enough nutrients added during production periods, which promoted optimal growth of culture and higher protein expression (Hadiyanto et al. 2013). Figure 4a showed that P. pastoris expressed the highest concentration of recombinant EGII after 72 h of incubation. Thereby, fed-batch fermentation for 72 h of incubation was used for further production of recombinant EGII.
The expression profile of recombinant EGII by clone 2 every 24 h intervals was visualized through SDS-PAGE analysis (Figure 4a). The maximum expression of recombinant EGII was reached after 72 h with 3.49 mg/L protein secreted from 20 mL of fermentation culture. The growth of transformants reached stationary phase after 72 h; thereafter, it entered the death phase. The estimated molecular mass of recombinant EGII was 52 kDa. EGII is a glycoprotein with 397 amino acids and has a molecular weight of 40 kDa without glycosylation (Garvey et al. 2014). However, its molecular weight may increases up to 48 kDa with native glycosylation done by T. reesei (Akbarzadeh et al. 2014). Sun et al. (2018) reported that the expression system by P. pastoris might increase up to 4 kDa of molecular weight due to glycosylation. In this study, the SDS-PAGE analysis of recombinant EGII protein expressed in P. pastoris showed a thick band with a slightly larger molecular mass than that of native EGII from T. reesei. However, there was not enough evidence to show that the increase in molecular mass is due to glycosylation. Treatment of recombinant EGII protein with endoglycosidase H may be required as supporting data to signify if glycosylation takes place.

3.3. Purification and confirmation of recombinant endoglucanase II

The culture supernatant was purified manually using Ni-NTA sepharose resin. The column was eluted with an isocratic buffer containing 250 mM imidazole to release the recombinant protein bound to the resin. All fractions collected were analyzed using SDS-PAGE. Among 12 elution fractions collected, the elution fraction 3 was known to have the thickest band of interest, indicating the highest concentration of purified protein (data not shown). All samples collected from sampling for each 24 h and elution fraction 3 were analyzed using SDS-PAGE followed by Western blot to confirm the recombinant EGII protein (Figure 4b). The result exhibited a single band of 52 kDa for each lane, which corresponds to a theoretical molecular mass of EGII recombinant proteins expressed in P. pastoris (Bai et al. 2016). The concentration of purified recombinant EGII was then quantified using a BSA standard curve followed by ImageJ analysis, resulting in 0.21 mg/mL of purified EGII.

3.4. Characterization of recombinant endoglucanase II

The endoglucanase activity of recombinant EGII was assayed at different pH (3.0, 5.0, 6.8, 8.0, 9.0, 10.0). The results showed that the recombinant EGII exhibited optimum activity at pH 5.9 (Figure 5). Activity above pH 8.0 was negligible as it did not show enzymatic activity. The native endoglucanase by T. reesei also showed the same optimum pH (Li et al. 2013). The study conducted by Boonvitthya et al. (2013) revealed that the production of crude EGII in P. pastoris was around 10 U/ml under a controlled condition with the optimum pH at 5 to 6 and the temperature of 40-60 °C.

Endoglucanase activity measurement was using various temperatures ranging from 30 °C to 70 °C. The optimum temperature was found to be 40 °C to 50 °C, as shown in Figure 6. Native endoglucanase also showed a similar range of optimum temperature, which was at 45 °C to 55 °C (Kamal et al. 2017). The effect of temperature optimum on enzyme stability is shown in Figure 6. Pre-incubation was done at 40 °C and 50 °C with different time intervals up to 60 min. About 89% of endoglucanase activity was retained at 40 °C after 60 min, and more than 80% of endoglucanase activity was retained after 60 min at 50 °C (Figure 7). Based on the literature, native EGII secreted from T. reesei retained 60% of its enzymatic activity at 50
FIGURE 5 Optimum pH determination of endoglucanase II by clone 2. Values are averages of duplo assays; bars indicate SD.

FIGURE 6 Temperature optimum determination of endoglucanase II by clone 2. Values are averages of duplo assays; bars indicate SD.

FIGURE 7 Legend for Figure 7: Thermal stability determination of recombinant Endoglucanase II by clone 2. Temperature used were at 40 °C (red) and 50 °C (black). Values are averages of duplo assays; bars indicate SD.

°C after 40 min of incubation (Kamal et al. 2017). This evidence indicated the thermal stability of our recombinant EGII is slightly improved compared to native EGII derived from T. reesei as it can retain 80% endoglucanase activity after incubation at 50 °C for 60 min.

The result of the thermal stability assay obtained was slightly lower than the study conducted by Samanta et al. (2012), which showed that recombinant EGII could retain relative endoglucanase activity up to 90% at 50 °C. However, this study reported that purified EGII protein showed 24.3 U/mL of enzyme activity after pre-incubation at 50 °C in pH 5.0, which gave the specific activity of 115.7 U/mg. The specific activity of our recombinant EGII is higher than the result reported by using the E. coli expression system (Nakazawa et al. 2008).

N-glycosylation, a post-translational modification generally occurred in fungi, has important roles in enzyme stability. A study conducted by Han et al. (2020) demonstrated that the addition of N-glycosylation at particular amino acid sites has successfully increased the thermal stability of recombinant endoglucanase in P. pastoris. Optimized stabilization is associated with entropy which mostly dependent on glycosylation sites position (Shental-Bechor and Levy 2008). Generally, glycans attached to flexible regions within random coils would restrict the conformational space and promote entropic reduction to increase conformational stability at high temperatures (Dotsenko et al. 2016; Adney et al. 2009). This additional targeted N-glycosylation was deserved to be adopted in our future study to improve the thermal stability of our recombinant EGII.

Optimization of fermentation is also required to improve the expression of P. pastoris by exploring various carbon and nitrogen sources to enhance microbial growth and protein expression. Further study to maintain a specific growth rate is also highly suggested by implementing a short carbon-starving period through an exponential feeding strategy.

4. Conclusions

Recombinant EGII proteins were successfully expressed in the P. pastoris expression system with an optimum incubation time of 72 h using fed-batch fermentation. Characterization of recombinant EGII using SDS-PAGE and Western blot analyses showed that the EGII protein has a molecular weight of approximately 52 kDa. The enzymatic assay demonstrated that the crude enzyme has optimum activity at pH 5.0, which results in 16.3 and 14.6 U/mL activity at a temperature of 40 °C and 50 °C, respectively. The specific activity of recombinant EGII resulted in 115.7 U/mg in its optimum pH and temperature. Moreover, recombinant EGII can maintain 89% of its endoglucanase activity at 40 °C and more than 80% at 50 °C for 60 min, indicating the improvement in thermal stability compared to native EGII derived from T. reesei.

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Authors’ contributions

KAYT – practical works, analysis and data interpretation, article drafting, KSD – design of the study, practical works, analysis and data interpretation, article drafting, AMF and TA – conception and design of the study. All authors read and approved the final version of the manuscript.

Competing interests

The authors declare no competing interest.

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