Isolation of Kojic Acid Producing Mold using Complex Carbon Sources

Ines Dawiyah Suwarjo, Adnina Fithra Azzahra, Herman Suryadi*

ABSTRACT

Introduction: An independent effort in term of pharmaceutical raw materials procurement in Indonesia, especially skin brightening agent, is required due to the high demand of brightening skin care product. One of the skin brightening agent widely used in cosmetic skin care formulations is kojic acid. This study aimed to obtain the isolate of kojic acid producing mold from nature and its optimum fermentation condition by using various complex carbon substrates.

Methods: Aspergillus oryzae was used as reference. The isolates of fungi were screened with different substrates variation, namely sucrose, corn starch, cassava starch, and cellulose hydrolysate. Then, each of culture was dripped with FeCl₃ 1% and the most brownish-red color formed was selected as mold and media for further process. The preculture of selected isolate and A. oryzae, were inoculated into 100 ml of fermentation media respectively and incubated at room temperature, 180 RPM for 10 days. The concentration of substrate was varied to 5, 7.5, and 10%. The levels of kojic acid were determined by TLC densitometry with UV detector at 318 nm. Results: IHJ2K isolate in corn starch-yeast extract was selected as the best mold and media. However, the highest level of kojic acid was produced by Aspergillus oryzae with 10% of substrate, with kojic acid concentration of 5.22 g/L. The most efficient fermentation was obtained from A. oryzae with 7.5% of substrate, with the obtained yield of 0.53 g/g. Conclusion: Potential kojic acid producing mold namely IHJ2K was successfully isolated. The selected carbon source for optimum kojic acid fermentation condition with IHJ2K isolate was corn starch combined with yeast extract, KH₂PO₄, and MgSO₄.7H₂O.

Key words: Aspergillus oryzae, Complex carbon, Fermentation, Kojic Acid, Mold, TLC Densitometry.

INTRODUCTION

Skin Care Products are the first Best Cosmetic Sales Prospects in Indonesia, including brightening products.¹ This fact leads the high demand pharmaceutical raw materials such as brightening agents are required. Unfortunately, 90% of pharmaceutical raw material in Indonesia is still imported.² Therefore, an independent effort in term of the raw materials procurement in Indonesia’s pharmaceutical industry, especially skin brightening agent, is required.

Kojic acid is one of the skin brightening agent widely used in cosmetic skincare formulations. Kojic acid is a major secondary metabolite produced by mold such as Aspergillus and Penicillium species.³⁴ Glucose, sucrose, acetate, ethanol, arabinose, and xylose have been used as carbon sources for kojic acid production.⁵⁻⁷ However, acetate and ethanol may affect the formation of aflatoxin.⁶ Glucose, sucrose, arabinose, and xylose are also expensive to use as raw materials. In this study, an alternative fermentation of kojic acid from selected mold isolates with complex carbon sources was performed. Based on a study by Rosfarizan et al. in 1998, the kojic acid level produced with corn starch compared to glucose as the carbon source was relatively similar.⁹

The aim of this study was to explore further the use of complex carbon such as corn starch, cassava starch, and cellulose hydrolysate as a substrate on kojic acid fermentation. In addition, the isolation of kojic acid producing mold using starch, offers the advantage of reducing the cost of raw materials for the production of kojic acid.

MATERIALS AND METHODS

Chemicals

Chemicals used in this study where: potato dextrose agar (PDA) (Difco), yeast extract (Merck), urea (CO(NH₂))₂ (Merck), ammonium sulphate ((NH₄)₂SO₄) (Merck), magnesium sulphate (MgSO₄.7H₂O) (Merck), dikalium phosphate (KH₂PO₄) (Merck), ferric chloride(FeCl₃) (Merck), orthophosphoric acid(Merck), distilled water, and aquabidest.

Instruments

The instruments used in this study were autoclave (Hirayama), oven (WTB Binder), analytical balance

Cite this article: Suwarjo ID, Azzahra AF, Suryadi H. Isolation of Kojic Acid Producing Mold using Complex Carbon Sources. Pharmacogn J. 2018;10(6):1089-92.
(Acculab), vortex mixer (Barnsted), hotplate stirrer (Corning), pH meter (Hanna), centrifugator (Kubota 6800), incubator (Memmert), shaker (Orbit), ose, pipette volume and other glass wares commonly used in laboratories. The analytical instruments used were TLC-Densitometer (Camag TLC Scanner 3) and UV-Vis Spectrophotometer (Shimadzu).

**Screening Media**

The media consisted of 5% variation of carbon sources (sucrose, corn starch, cassava starch, and cellulose hydrolysate), 0.5% variation of nitrogen sources (yeast extract, urea, ammonium sulfate), 0.1% KH\(_2\)PO\(_4\) and 0.05% MgSO\(_4\).7H\(_2\)O. The media was prepared in twelve variations. The pH of media solution was adjusted to 5 with orthophosphoric acid.

**Isolation of Molds**

Molds were isolated from soil and rotten wood. The surface of the soil was cleaned from waste and dirt, then 100 mg of the moist layer beneath was put into the vial. A 100 mg of rotten wood were crushed in a sterile mortar. A hundred milligram of either the soil or the wood was diluted in 10 ml of aquabidest. The soil and wood suspension were diluted until 10\(^{9}\) of suspension was obtained.

Both of the suspension were scratched onto potato sucrose agar (PSA) media in a petri dish and incubated at 28°C for 5-7 days. Mold colonies grew were isolated based on their morphology. The colony was taken with inoculating loop and scratched onto the new PSA. The new PSA contained colonies was then incubated at 28°C for 5-7 days. The ideal colony obtained was transferred to the culture stock tube. The procedure was repeated if the colony obtained had not met the criteria. The purified colonies were transferred to a test tube containing the tilted potato dextrose agar (PDA) media and stored as a culture stock for further test preparation.

**Screening of Superior Mold and Media Combination**

The mold colonies seeded on PDA were cultured into 96-well plates containing a 200 μL of screening media, incubated for 6 days at 28°C. A total of 130 μL of supernatant was placed on a drop plate. The screening was done by dripping fresh FeCl\(_3\), 1%. The brownish red color formed was indicated a positive result and the most intense color was selected as the superior mold and media combination.

**Macroscopic and Microscopic Identification**

Macroscopic identification was done by observing the colony color, colony diameter, and colony texture. Microscopic identification was performed by examination and observation of mold preparations under a light microscope. Both macroscopic and microscopic observation were compared with *Aspergillus oryzae* (IPB Culture Collection).

**Kojic Acid Fermentation**

The pre-culture of the selected isolate was prepared with 50 mL of pre-culture media (superior media) in a 100 mL shake flask. Incubation was carried out by shaking at 180 rpm, 28°C for 48 h. Preparation of inoculum was also performed on *Aspergillus oryzae*. Fermentation was done with 100 mL media and 10% (v/v) inoculum in 250 mL shake flask. The level of carbon source was varied to 5, 7.5, and 10%. The flasks were incubated by rotary shaker at 180 rpm, 28°C for 10 days. At the same time, fermentation of *Aspergillus oryzae* was also performed.

**Analytical Methods**

The presence of kojic acid in the culture filtrate was determined using UV-Vis spectrophotometry and colorimetric method, and analyzed quantitatively using TLC-Densitometer (CAMAG III, Switzerland), using silica gel F\(_254\) as stationary phase, toluene-ethyl acetate-formic acid (3:6:1) as mobile phase, and UV detector at 318 nm. The calibration curve was made using kojic acid standard solution with a range of 20-80 ppm. Glucose concentration was determined using DNS method. Biomass was determined by the dry cell weight method. Culture samples were pipetted into a pre-weighed tubes and centrifuged at 7500 RPM for 15 min. Supernatant was used for kojic acid analysis, while biomass was washed with distilled water, centrifuged, and dried at 105°C to a constant weight.

**RESULTS AND DISCUSSION**

**Isolation of Molds**

Eight colonies were obtained based on the morphology. There were two black-spores colonies surrounded by white mycelia named IHIT (Figure 1.a) and IHJK (Figure 1.b), which secreted yellow pigment underneath. A green colony surrounded by a white mycelia was isolated, which was IHIJT (Figure 1.c). A different green colony that secreted yellow pigment underneath was also isolated and was named IHIJK (Figure 1.d). There was also a yellowish-green colony surrounded by white mycelia named IHJ2K (Figure 1.e), a yellow-green colony isolated from wood, named IHIJKK (Figure 1.f), a brownish-green that was isolated from wood called ICK (Figure 1.g), and a brown colony isolated from soil which was named ICT (Figure 1.h).

**Screening of Superior Mold and Media Combination**

The IHJ2K (Figure 2.e) and IHIJKK (Figure 2.f) were the types of mold which produced an intense brownish color. Re-screening was performed on IHJ2K and IHIJKK fermentation after two days. The results showed that IHJ2K in media containing 5% corn starch, 0.5% yeast extract, 0.1% KH\(_2\)PO\(_4\), and MgSO\(_4\).7H\(_2\)O 0.05% yielded the most intense brownish-red color (Figure 3.a).
The result showed that IHJ2K isolate contained amylase, glucoamylase, and other hydrolase enzymes that play a role in the metabolism of starch. Thus, IHJ2K could hydrolyze amylase and amylpectin into glucose, a precursor compound in the kojic acid biosynthetic pathway. This suggested that IHJ2K was able to produce kojic acid on complex carbon sources in a short time.

Corn starch was selected as a superior carbon source due to its low percentage of amylpectin compared to cassava starch. The degree of polymerization (DP) of amylpectin ranged from $10^2$ to $3 \times 10^6$ units of glucose. The more units required to be hydrolyzed on cassava starch the longer time required for kojic acid productivity. Meanwhile, β-cellulose had lower kojic acid productivity than corn starch due to the presence of cellulose which had not been fully hydrolyzed to glucose. Yeast extract was also selected considering its complex structure compared to urea and ammonium sulfate to provide a richer nutritional intake.

**Macroscopic and Microscopic Identification**

The macroscopic identification results showed that the growth of IHJ2K on PSA media with temperature of 28°C grew to 8 cm in diameter in 7 days. The colony had a yellowish green color surrounded by white mycelium and velvety texture (Table 1). IHJ2K had the same spores as *Aspergillus oryzae*. *Aspergillus* had a type of asexual spore called conidia.

**Kojic Acid Fermentation**

The UV spectrophotometric absorption spectra of the kojic acid standard and culture sample (IHJ2K in 10% corn starch) indicated the same maximum wavelength, which was 268.8 nm.

The spectrocolorymetry absorption spectra of kojic acid standard and culture sample (IHJ2K in 10% corn starch) also showed the same maximum wavelength at 503.6 nm. This result showed that the IHJ2K culture contained kojic acid.

The biomass produced by IHJ2K was higher compared to *A. oryzae*. The higher substrate concentration, the higher biomass produced by IHJ2K which could be seen in a pattern (Table 2). Meanwhile, the escalation of *A. oryzae*’s biomass was not as patterned as IHJ2K’s. Too much sugar led to culture leakage to the *A. oryzae* fermentation in 10% of corn starch was less efficient. In addition, the more level of corn starch substrate used by IHJ2K, the more the level of kojic acid was produced.

Overall, the yields obtained by IHJ2K fermentation were less than by *Aspergillus oryzae*. Although the IHJ2K isolate showed an increase in kojic acid concentration and its yield by increasing the substrate concentration, but it had not been able to outperform the efficiency of kojic acid productivity resulted by *Aspergillus oryzae*.

**CONCLUSION**

In conclusion, kojic acid producing mold called IHJ2K was successfully isolated. The carbon source for optimum kojic acid fermentation condition with IHJ2K isolate was corn starch combined with yeast extract, KH$_2$PO$_4$, and MgSO$_4$, $7\text{H}_2\text{O}$.

**ACKNOWLEDGEMENT**

The authors are grateful to DRPM Universitas Indonesia for grant “Hibah PITTA 2017” which supporting this study.

**CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest.

**ABBREVIATIONS**

- TLC: Thin Layer Chromatography
- PSA: Potato Sucrose Agar
- PDA: Potato Dextrose Agar
- UV: Ultraviolet
- UV-Vis: Ultraviolet-Visible
- DNS: 3.5-dinitro salicylic acid

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**Table 1: Macroscopic and Microscopic Comparison of IHJ2K with *Aspergillus oryzae***

<table>
<thead>
<tr>
<th>Mold</th>
<th>Morphology</th>
<th>Microscopy</th>
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</thead>
<tbody>
<tr>
<td>IHJ2K</td>
<td><img src="image1.png" alt="Image" /></td>
<td><img src="image2.png" alt="Image" /></td>
</tr>
<tr>
<td><em>Aspergillus oryzae</em> (IPB Culture Collection)</td>
<td><img src="image3.png" alt="Image" /></td>
<td><img src="image4.png" alt="Image" /></td>
</tr>
</tbody>
</table>

**Table 2: Growth of IHJ2K and *A. oryzae* and their kojic acid production.**

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Corn starch (g/L)</th>
<th>Biomass (g/L)</th>
<th>Kojic acid (g/L)</th>
<th>Glucose remained (g/L)</th>
<th>Glucose consumed (g/L)</th>
<th>Yield (g/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IHJ2K</td>
<td>50</td>
<td>17.84</td>
<td>-</td>
<td>0.19</td>
<td>49.81</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>75</td>
<td>25.78</td>
<td>0.204</td>
<td>0.23</td>
<td>74.77</td>
<td>2.73 x 10$^3$</td>
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<tr>
<td></td>
<td>100</td>
<td>30.58</td>
<td>3.781</td>
<td>74.74</td>
<td>25.26</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>15.37</td>
<td>-</td>
<td>0.22</td>
<td>49.78</td>
<td>-</td>
</tr>
<tr>
<td><em>Aspergillus oryzae</em></td>
<td>75</td>
<td>22.13</td>
<td>3.158</td>
<td>69</td>
<td>6</td>
<td>0.53</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td>22.23</td>
<td>5.22</td>
<td>72.74</td>
<td>27.26</td>
<td>0.19</td>
</tr>
</tbody>
</table>

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**Figure 3:** Screening results of IHJ2K and IHJKK culture filtrate.
REFERENCES


SUMMARY

• Isolation of molds from soil and rotten wood. Eight colonies were obtained
• Screening of superior mold and media combination in producing kojic acid. A colony called IHJ2K which fermentated in corn starch combined with yeast extract, KH₂PO₄, and MgSO₄·7H₂O was selected.
• Identification of the superior mold macroscopically and microscopically
• Qualitative and quantitative analysis of kojic acid

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Cite this article: Suwarjo ID, Azzahra AF, Suryadi H. Isolation of Kojic Acid Producing Mold using Complex Carbon Sources. Pharmacog J. 2018;10(6):1089-92.