Synthesis of L-Lactic Acid from Fermentation of Cassava Pulp by Using Tempeh Inoculum

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ABSTRACT – This study used cassava waste pulp as a fermentation substrate to produce lactic acid using a tempeh inoculum. Tempeh inoculum is a mixed culture of Rhizopus with Rhizopus oligosporus as the primary fungus. Lactic acid is an organic acid most widely used in the food, pharmaceutical, cosmetic and chemical industries. One of the important uses of lactic acid is as a raw material for producing Polyactic Acid (PLA) biopolymers, namely polymers that can decompose naturally in a relatively fast time. The analysis was performed using the Response Surface Methodology (RSM) method and the Box Behnken Design (BBD) experimental design with substrate concentration parameters, inoculum concentration, and incubation time on lactic acid. The fermentation process is carried out using a flask shaker at a temperature of 30°C, pH 6.0, and a rotational speed of 150 rpm. The optimum yield for lactic acid is 6.65 g/L. It was acquired at substrate 20 g/L, inoculum concentration 0.30 % (w/v) at an incubation time of 72 hours.

INTRODUCTION

Lactic Acid (LA) has a wide range of applications, especially as a raw material of bio-polymer production [1]. Synthesize of LA from starch-biomass or agricultural residues was begun since this material is cheap and abundant. One of the important LA derivatives (biopolymer Polyactic Acid. PLA) has profound properties; thus, it has been widely used in various applications [2]. PLA is a biodegradable and thermoplastic made by polymerization of LA. PLA is a promising material to overcome global pollution caused by conventional plastic [3].

In producing LA, the utilization of waste by biological processes such as fermentation has an important role [1]. Cassava waste pulp is a potential raw material in LA production. Cassava waste pulp is a solid waste of cassava starch production. The main components are carbohydrates (≥ 60-65%), and the remaining 40% contain moisture, minerals, and crude fiber. Meanwhile, cassava waste pulp is generally used only as fertilizer and animal feed, so it has not been optimally utilized [4, 5]. Several studies on cassava waste pulp utilization as substrate fermentation have started performed [6, 7]. This work chooses cassava waste pulp as a substrate due to its relatively cheap role in overcoming environmental pollution problems.

LA is usually produced by chemical synthesis from petrochemical sources or obtained by microbial fermentation. The LA produced by bacteria fermentation requires complex media, producing an LA racemic mixture [8]. However, fermentation is possibly carried out by using inoculum, such as tempeh-inoculum. The fungi usually cultivated in the inoculum such as R. oligosporus, R. oryzae, R. stolonifer, and R. arrhizus [9].

Some works reported that R.oryzae and R. oligosporus could utilize various agricultural residues such as L-LA, fumaric acid, and ethanol to produce LA [10, 11]. The use of Rhizopus has several advantages; it can suppress downstream processes because of the ease of separating mould filaments. Previous researchers have studied LA fermentation using strain Rhizopus sp, with different carbon sources, nitrogen sources, and operating conditions (pH, temperature, volume, and rotation speed) but mostly using acid hydrolysis to convert polysaccharides into monosaccharides [10-12]. Acid hydrolysis has drawbacks such as corrosion-resistant tools, carbohydrate degradation and product recombination [8].

Rhizopus sp has the shortest log phase, where the production is the highest mycelium mass occurs from 48 to 96 hours of incubation. After this time, the amount of mycelium remains even much lower than the log phase [13]. Some literature shows that the optimum result for fermentation using Rhizopus is at 72 hours, so that the observation of incubation time is chosen between 0-72 hours [14].

Several researchers have used this technique to optimize the yield of several parameters. This study used RSM-BBD with a factorial arrangement of three levels of fractions that can predict the best optimal value of the LA yield. This methodology could be employed to optimize media for LA fermentation [10, 15]. This method is to determine the effect of the independent variable on the response, obtain the model of the relationship between the independent variable and the response, and get the optimum condition process. In addition, the advantages of the RSM-BBD method does not require large number of experimental data and efficient time.

This study aims to ferment cassava-waste pulp into LA. The fermentation starter using tempeh inoculum, which has not been widely used in the fermentation process to produce LA.

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**EXPERIMENTAL METHOD**

The research was conducted in the bioprocess laboratory Research Unit for Clean Technology, National Research and Innovation Agency Republic of Indonesia (BRIN).

**Materials and Instruments**

Dried cassava waste pulp was obtained from Surya Pati Kencana Factories (Pati, Indonesia), sieved 80 mesh. The commercial tempeh inoculum of the Raprima brand, produced by PT Aneka Fermentasi Indonesia (AFI.) Bandung – Indonesia. Yeast extract, peptone, K$_2$HPO$_4$, KH$_2$PO$_4$, MgSO$_4$.7H$_2$O, (NH$_4$)$_2$SO$_4$, CaCO$_3$, Potato Dextrose Agar (PDA) were purchased from Merck (Darmstadt, Germany).

**Method and Procedure**

**Fermentation Process**

The fermentations at shake flask level (50 mL in a 250 mL Erlenmeyer flask) were carried out using the media described by Kumar et al. [14]. Media fermentation was sterilized using an autoclave (Hirayama H.V.E. 50) at a temperature of 121 °C for 15 minutes. Fermentation was carried out in a shaker incubator (Jeiotech SI-900R) at a temperature of 30 °C with a rotating speed of 150 rpm and pH 6.0.

**Statistical Optimization**

A three factorial experimental design using the RSM-BBD was applied to optimize the fermentation process using MINITAB 19.1 software. The Design of Experiment (DOE) was formed based on the RSM in doing optimization. The independent variable used is the substrate concentration (A), inoculum concentration (B), and incubation time (C). The response of independent variables will affect LA levels. In this study, the determination of the parameters used are shown in Table 1.

| Factor | Parameter | Min | Max |
|--------|-----------|-----|-----|
| A      | C (g/L)   | 20  | 60  |
| B      | Inoc (% w/v) | 0.10 | 0.30 |
| C      | t (h)     | 0   | 72  |

A : substrate concentration; B: inoculum concentration; and C: incubation time

A second-order polynomial model which includes all interaction terms was defined to fit the response, shown in Equation 1.

$$Y = \beta_0 + \sum \beta_iX_i + \sum \beta_{ij}X_iX_j$$  \hspace{1cm} (1)

Y is the predicted response (LA, g/L), $\beta_0$ is the interception coefficient, $\beta_i$ is the linear term, $\beta_{ij}$ is the quadratic term, and $\beta_{ij}$ is the interaction term. Calculation of the predicted responses, analysis of experimental design data, and the plotting of contour and surface plots were made using MINITAB 19.1 software.

Statistical analysis of the data: Data analysis where the response variable (Y) obtained was then analyzed by ANOVA. The coefficient of determination ($R^2$) was used to assess the fitting of second-order regression equations. By using three levels of BBD, it will produce 15 experiments.

**Characterization**

Cassava waste pulp was tested for proximate characteristics such as moisture, starch, ash, and crude fiber for preliminary testing. Dried cassava waste pulp containing moisture of 11.74%, starch content 59.40 %, ash content 1.95 %, and crude fiber content 16.42%. The tempeh inoculum was identified and tested for amylase enzyme activity using the Fuwa method [16]. Macroscopic observations were made of fungal colonies' color growth on the petri dish. Microscopic observations were analyzed using microscope Fluorosence Meiji Techno MT 6300 to know Rhizopus type in the tempeh inoculums [9]. Response testing LA production; The supernatant was measured with a Conductometer to determine its electrical conductivity, then plotted on the standard curve LA [17]. A regression equation is obtained from the standard curve of the conductivity measurement to calculate the yield of LA production (g/L). The equation leads to regression formula in Equation 2.

$$y = a \times x + b$$  \hspace{1cm} (2)

y is electrical conductivity (mS/cm), a is the slope, b is the y-intercept, and x is the yield of LA production (g/L).

FT-IR spectroscopy (Thermo Scientific Nicolet iS5, Madison, U.S.A.) identifies the functional groups resulting from LA fermentation[18].
RESULT AND DISCUSSION

Based on macroscopic and microscopic at 100x magnification, which was then compared with the standard, the type (a) resembled *R. oligosporus*, and (b) reached as *R. oryzae*. The results of the isolation of pure culture from the tempeh inoculum are shown in Figure 1. This study’s results indicated insulation on fungal isolates color tempeh inoculum greyish black and greyish-white in 48 hours of incubation. After more than 72 hours, the colony starts to turn grey-brown and greyish black.

![Figure 1](image-url)  
*Figure 1*. Macroscopic (left) and microscopic (right) morphology 100x magnification of (a) *R. Oligosporus* and (b) *R. oryzae* syn. *R. arrhizus* in Potato Dextrose Agar after seven days incubation using Microscope Fluorescence.

*R. oligosporus* has a shorter sporangiospores length than *R. oryzae*, which has short rhizoids. In seven days, the sporangium rupture causes the spores to leak out of the columella. In comparison, *R. oryzae* has a diameter of sporangium and columella larger than *R. oligosporus*. For *R. Oryzae*, sporangiospores were unequal, numerous, irregular, sub-globose or oval, angular with striations, 6–10 μm in length. Sporangiophores were usually straight, smooth-walled, simple or branched, non-septate, long, and long arose from stolons opposite rhizoids, usually in 3–5 μm or more. Sporangia were globose, white at first, and then turned black with many spores. Columella was globose to sub-globose in shape, pale brown in color. Rhizoids and stolons were dark brown (Figure 1) [19]. *R. oligosporus* has brownish-gray colonies with a height of 1 mm or more. A single colony or in a group have smooth or rather rough walls. Sporangia are brownish-black. Chlamydospores have single and short, colorless chains containing granules formed on hyphae, sporangiophores, and sporangia [9]. The form of Chlamydomyospora globose, ellipse, or cylindrical. For morphology, the details are shown in Table 2.

| Characteristics               | Identification of the commercial tempeh inoculum by microscope | Reference          |
|-------------------------------|-----------------------------------------------------------------|--------------------|
|                               | A                  | B                  | *R. oligosporus* [20] | *R. oryzae* [21] |
| Conidia color                 | Brownish grey      | Brownish grey to blackish-grey | Brownish-grey      | Brownish grey to blackish-grey |
| Mycelium color                | White              | White              | White               | White               |
| Sporangiospores Size          | 78–121 μm          | 171–420 μm         | 150–400 μm          | 150–1000 μm         |
| Sporangiospores shape         | Sub-globose to ellipsoidal | Sub-globose or oval | Sub-globose to ellipsoidal | Sub-globose or oval |
| Diameter sporangium           | 49–76 μm           | 79–120 μm          | 80–120 μm           | >150 μm             |
| Columella length              | 28 μm              | 36 μm              | 25–27 μm            | 80–110 μm           |
| Texture sporangiophore        | smooth             | smooth             | smooth              | smooth              |

Table 2. Comparison of the morphological characteristics of fungi isolated from Tempeh inoculum commercial with previous features of *R. oligosporus* and *R. Oryzae*.
Based on Table 2, comparing the macroscopic and microscopic analysis of the tempeh inoculum with reference states that the commercial tempeh inoculum was assumed to contain *R. oryzae* and *R. oligosporus* fungi type producing amylase and glucoamylase enzymes [22]. It was seen from the similarity of shapes, sizes, and colors based on reference, but there are also differences in morphology of *Rhizopus* compared to the standard due to the different strains of *Rhizopus* [19]. The tempeh inoculum's enzyme activity test result was amylase of 79.00 U/mL. Converting polysaccharides into monosaccharides (glucose) enzymatically involves a group of the amylase enzymes, where glucose will be used as a carbon source for the LA fermentation process. According to research by Freitas et al., the amylase enzymes produced from *R. oryzae* and *R. oligosporus* were converted 95% and 96% of the starch into glucose in 3.5 hours of reaction. This statement shows us that the commercial tempeh inoculums are the best choice due to converting 95% -96% of polysaccharides into monosaccharides in only 3.5 hours [23].

Figure 2 was LA produced by the fermentation process, which has brownish yellow color because it still contains sugar and nitrogen sources residues. Figure 3 shows the FT-IR spectrum of standard LA, L (+) - lactate, and the results of LA fermentation with calcium carbonate precipitates. FT-IR spectrum of L(+)-fermented calcium lactate has a higher carbonyl bond at 1500 ~ 1750 cm⁻¹ than others.
analyzed using statistical methods following the Design of Experimental. Multiple regression analysis of experimental data yields the following second-order polynomial equation, in Equation 3.

\[
LA \left( \frac{g}{L} \right) = -0.150 - 0.0261(A) + 9.96(B) + 0.1377(C) + 0.000960(A)^2 - 0.90(B)^2 - 0.000954(C)^2 - 0.2501(A)(B) - 0.000544(A)(C) + 0.0826(B)(C).
\] (3)

From Equation 3, the curve fitting for LA production can be calculated which the data can be compared with the experimental value (Table 3). If the value between the curve fitting and the experimental results is too far away, other factors may affect LA production. The experiment result was obtained from the measurement of electrical conductivity. Afterwards, a regression equation is obtained to calculate the yield of LA production (g/L) from the standard curve of the conductivity measurement. Equation 4 is the equation from the standard curve of conductivity measurement:

\[
y = 3.2133 . x + 5.5392
\] (4)

| Run orders | A (g/L) | B (% w/v) | C (h) | Conductivity mS/cm | LA production (g/L) |
|------------|---------|-----------|-------|--------------------|---------------------|
|            |         |           |       | Curve Fitting      | Experimental        |
| 1          | 60      | 0.20      | 0     | 7.12               | 0.69                | 0.49               |
| 2          | 40      | 0.10      | 72    | 19.14              | 5.11                | 4.23               |
| 3          | 40      | 0.10      | 0     | 7.19               | 0.33                | 0.51               |
| 4          | 40      | 0.30      | 36    | 6.63               | 0.25                | 0.34               |
| 5          | 40      | 0.20      | 36    | 17.43              | 3.83                | 3.70               |
| 6          | 60      | 0.30      | 36    | 17.40              | 3.58                | 3.69               |
| 7          | 40      | 0.30      | 72    | 22.40              | 4.65                | 5.24               |
| 8          | 40      | 0.20      | 36    | 18.17              | 3.83                | 3.93               |
| 9          | 20      | 0.20      | 0     | 7.44               | 0.67                | 0.59               |
| 10         | 60      | 0.20      | 72    | 20.25              | 5.28                | 4.58               |
| 11         | 20      | 0.20      | 72    | 25.60              | 5.26                | 6.24               |
| 12         | 20      | 0.30      | 36    | 22.64              | 4.56                | 5.32               |
| 13         | 40      | 0.20      | 36    | 17.93              | 3.83                | 3.85               |
| 14         | 60      | 0.10      | 36    | 18.67              | 4.85                | 4.09               |
| 15         | 20      | 0.10      | 36    | 17.48              | 3.83                | 3.72               |

A : concentration of cassava waste
B : concentration of tempeh inoculums
C : incubation time
LA : Yield of LA

| Source | Degree of Freedom | Adj SS. | Adj Mean square | F-Value | P-Value |
|--------|------------------|---------|-----------------|---------|---------|
| Model  | 9                | 52.3024 | 5.8114          | 125.89  | 0.000   |
| Error  | 5                | 0.2308  | 0.0462          |         |         |
| Pure Error | 2        | 0.0276  | 0.0138          |         |         |
| Total  | 14               | 52.5332 |                 |         |         |
| R²     |                  | 0.9956  |                 |         |         |
| Adj R² |                  | 0.9877  |                 |         |         |
| R² (prediction) |      | 0.9369  |                 |         |         |
| Lack of Fit | 3           | 0.2032  | 0.0677          | 4.91    | 0.174   |

Based on the prediction RSM to optimization of LA production (g/L) by tempeh inoculum was A= 20 g/L, B= 0.30 % (w/v), and C= 72 hours, will produce LA= 7.08 g/L. Three repetitions were carried out to confirm optimum conditions, and an average value of 6.65 g/L was obtained. This value is very close to the predicted optimum value, 7.08 g/L. The experiment validated the optimum conditions predicted by the BBD. The percentage of error between the experimental
results and the curve fitting is 0.38%. The R² value (0.9956) indicates that the sample variation for LA of 99.56% was attributed to the independent variables, and the model cannot explain only 0.44% of the total variation.

The adjusted determination coefficient (adj. R² = 0.9877) was also satisfactory for confirming the model's significance. The closer to number one, the higher the value of optimization accuracy. The analysis results also showed a significant interaction between cassava waste pulp substrate and the inoculum content. The relationships between independent variables can be better understood by examining the series's response surface and contour plots. Figures 4 to 6 represent surface plots and contour plots for optimizing LA fermentation conditions. The response surface's primary objective is to efficiently determine the variable's optimal value to maximize or minimize response. The smallest ellipse in the contour diagram is the maximum predicted value indicated by the boundary surface. Less substrate causes microorganisms to lack substrate in their reproduction; Conversely, more substrate concentration causes microorganisms to experience a long phase delay (adaptation). The interaction between the substrate concentration and the inoculum concentration can be seen in Figure 4.

![Figure 4](image1.png)

**Figure 4.** Response surface plot and contour plot showing the effect of substrate concentration, inoculum concentration, and their mutual impact on LA production (g/L).

![Figure 5](image2.png)

**Figure 5.** Response surface plot and contour plot showing the effect of substrate concentration, incubation time, and their mutual impact on LA production (g/L).

![Figure 6](image3.png)

**Figure 6.** Response surface plot and contour plot showing the effect of inoculum concentration, incubation time, and their mutual impact on LA production (g/L).
CONCLUSION

Commercial tempeh inoculums and cassava waste pulp as LA producers makes the effective fermentation process for bioplastic technology. The optimum condition at a cassava waste pulp concentration of 20 g/L and 0.30% (w/v) tempeh inoculum with 72 hours of incubation time yields a maximum LA production of 6.65 g/L. However, a well-established study on microorganisms producing LA with renewable raw material using the RSM technique still requires more research to be generalized and applied to a broader LA industry scope.

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