COMPARATIVE STUDY ON ACTIVITIES OF ANTI BACILLARY DYSENTERY SHIGELLA DYSENTERIAE OF SYZYGIUM POLYANTHUM AND DRACAENA ANGSTIFOLIA LEAVES ETHANOL EXTRACTS

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ABSTRACT

Objective: This study aims to determine the antibacterial activity of ethanol extract of bay (Syzygium polyanthum W.) and suji (Dracaena angustifolia Roxb.) leaves against Shigella dysenteriae and the amount of potassium to the discovery of anti-dysenteric drug candidates.

Methods: Testing activities and comparative value activities performed by the agar diffusion method, whereas the determination of minimum bactericidal concentration (MBC) was done with the subculture media incubation test followed with microdilution method on Mueller Hinton Agar medium sterile. Potassium levels of the extract were carried out quantitatively using atomic absorption spectrophotometry.

Results: The test results showed that the ethanol extract of both leaves had antibacterial activity against S. dysenteriae with MBC values were in the range of 10-20% w/v. Value comparative effectiveness suji leaf ethanol extract to the bay leaf was 1:0.4. Potassium levels ethanol extract of bay and suji leaves were 1.027% and 3.795%, respectively.

Conclusion: It can be concluded that the ethanol extract of bay and suji leaves has antibacterial activity against bacteria S. dysenteriae ATCC 13313. Rated comparative activity of the ethanol extract of the suji leaves with bay leaves ethanol extract was 1:0.4 which means to generating resistance to bacillary dysentery patients due to hypokalemia dysentery.

Keywords: Bay leaves, Suji leaves, Shigella dysenteriae, Potassium, Atomic absorption spectrophotometry.

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INTRODUCTION

Dysentery by cause divided into two, namely, amoebic dysentery and bacillary dysentery [1]. Both dysentery generally has the same symptoms, but bacillary dysentery is usually more common in children. Bacillary dysentery is an intestinal infection caused by Gram-negative bacilli of the Shigella dysenteriae [2]. About 1.1 million people are estimated to die each year, with 60% of the cases are from children under 5 years old [3]. Complications that often arise in bacillary dysentery are dehydrated [4]. Cases of dehydration cause the body to lose a lot of water with salts, especially potassium and sodium [2]. Hypokalemia reported cases are more prevalent. Hypokalemia occurs when the concentration of potassium in the blood of <3.5 mEq/L of blood and the risk of death [5]. This danger becomes great for babies and children because it can to lose more body fluid (15%) compared with adults (4%). Thus, in addition to a drug given anti-dysentery, required supply potassium to prevent hypokalemia due to dehydration.

Treatment of bacillary dysentery is generally performed by administering tetracycline, ampicillin, cotrimoxazole, ciprofloxacin, and chloramphenicol [6]. However, reported cases of S. dysenteriae resistance against the antibiotic is ampicillin (82%), cotrimoxazole (84%), and chloramphenicol (82%) [7]. Thus, the necessary searches anti bacillary dysentery potential candidates to overcome the resistance of existing cases. The use of natural materials is one of the solutions in the discovery of new drugs for anti-bacillary dysentery. Reported that the bay leaves (Syzygium polyanthum W.) and suji leaves (Dracaena angustifolia Roxb.) had a strong bacillary antidisenet activity. In addition, it was reported also that the test material has fairly high potassium content [8,9]. This open up a new interest to examine the anti-bacillary dysentery activity comparisons between two as well as their potassium content. Thus, the results of this study were obtained candidate anti bacillary dysentery capable of supplying potassium so as to reduce the death rate of bacillary dysentery patients due to hypokalemia.

METHODS

Plant material
Plant material samples used in this study was a bay (S. polyanthum W.), and suji leaves (D. angustifolia Roxb.) obtained from Plantation Manoko, Lembang, West Java and determination plants was performed in Taxonomy Laboratory, Department of Biology, Faculty of Mathematics and Natural Sciences, University of Padjadjaran.

Bacteria test
Bacteria test used is S. dysenteriae ATCC 13313.

Media growth of test bacteria was Shigella-Salmonella (SS) (Pronadisa), Mueller-Hinton agar (MHA) (Merck) and Mueller-Hinton broth (MHB) (Oxoid).

Extraction and extract parameters
The making extract was done by maceration method. The test material was extracted with ethanol 70% 3 × 24 hrs. Liquid extract was collected and then concentrated by rotary evaporator at a temperature of 60°C to...
obtain a thick extract. Viscous extract and then heated over a water bath at a temperature of 50°C to remove residual solvent remaining [10]. Inspection parameters included an examination of the organoleptic, yield, and moisture content of the extract.

**Phytochemical screening**

Phytochemical screening was conducted using modified phytochemical screening of plants [11].

**Thin layer chromatography (TLC)**

Determination profile TLC was performed using thin layer plate. The stationary phase used was silica gel 60 F254 and the mobile phase was a mixture of chloroform: methanol: 9:1.

**Confirmation of test bacteria**

~24 hrs at a temperature of 37°C. Colony morphology observed for its shape:

1. Observation morphology colonies
   Observations colony morphology was done by scraping the bacterial suspension in a sterilized MHB liquid medium on the SS surface. Media were then incubated for 18 and color.

2. Gram-staining
   Gram staining was performed by making a smear test bacteria suspension on the slide, and then the slide was passed on fire. Furthermore, a smear on the glass objects imbed with carbolic gentian violet for 1 minute. Excess dye was removed and rinsed with distilled water. Spreadable flooded with Lugol for 2 minutes. Lugol excess then removed and rinsed with distilled water. After this, the smear was washed with 95% alcohol drop by drop until the dye dissolved and then rinsed with distilled water. Last coloring stage was flooded smear methylene blue for 30 seconds and discharged excess dye and distilled water rinse. Further object glass filter paper and dried with heat immersion plus then observed under a microscope. Gram-positive bacteria in purple and Gram-negative bacteria were blue.

3. Biochemistry test
   Biochemistry test was conducted on the motility test, carbohydrate fermentation, Indol, TSA (Tripe Sugar Iron), Urace, methyl red, Voges Prokauer, and citrate.

**Antibacterial activities test**

1. Preparation equipment and materials
   Before testing the antibacterial activity, tools and materials used for the activity test sterilized using an autoclave at a temperature of 121°C for 15 minutes.

2. Preparation of bacterial growth media
   Solid medium used was that SS and MHA. 63 g that SS was dissolved in 1 L of distilled water. Media should not be sterilized by autoclaving. To reduce contamination, and the distilled water used must be sterile. Media in the mix with distilled water in a sterile Erlenmeyer flask aseptically and heated in a water bath until boiling. While the MHA media creation done by dissolving 30 g of media into 1 L of distilled water and then heated in a water bath until dissolved. Sterilization media performed by autoclaving at 121°C for 15 minutes. 21 g MHB dissolved in 1 L of distilled water and heated in a water bath until dissolved. Sterilization media is done with an autoclave at a temperature of 121°C for 15 minutes.

3. Preparation of bacteria test
   Bacteria embedded test in order SS 5 mL in a test tube (for oblique) by way of scraping a test bacteria in a zig-zag, then incubated for 18-24 hrs at a temperature of 37°C.

4. Preparation of standard solution McFarland
   McFarland standard solution consists of two components, namely, BaCl₂ solution of 1% and 1% H₂SO₄. A total of 0.05 mL of 1% BaCl₂ solution was mixed with 9.95 mL of 1% H₂SO₄ solution and shaken until homogeneous. The turbidity of the solution was measured at a wavelength of 620 nm using distilled water as blank. The absorbance value of the standard solution should be in the range of 0.08 up to 0.13. Standard solution equivalent to a 0.5 McFarland suspension of bacterial cells at a concentration of 1.5x10⁸ CFU/mL.

5. Preparation of bacteria suspension
   Dysentriae Shigella bacteria which have been cultured in the media so that the SS for 18-24 hrs at a temperature of 37°C, taken one Ose, then suspended in sterile MHB. Turbidity bacteria were measured using a standard 0.5 McFarland. Then, incubated at a temperature of 37°C for 18-24 hrs.

6. Testing antibacterial activity
   The method used to test the antibacterial activity of ethanol extract of bay and suji leaves was the agar diffusion method with perforation technique. Bay leaf ethanol extract dissolved in dimethyl sulfoxide (DMSO) to obtain the extract solution with a concentration of 80%, 40%, 20%, and 10% w/v. Sujil leaf ethanol extract dissolved in DMSO to obtain the extract solution with a concentration of 100%, 80%, 40%, 20% w/v. Sterile petri dish diameter of 10 cm was prepared and into it entered 20 mL test bacteria suspensions equivalent to McFarland 0.5, then poured agar medium MHA with a temperature of 20 mL at 400°C. Having shaken gently until a homogeneous, test medium was left to solidify. The medium test was divided into four zones and perforated using a perforator. A volume of 100 mL of the extract solution with different concentrations inserted into a hole on each cup. Grail incubated in an incubator at 37°C for 18-24 hrs. Tests conducted triplo for each cup. As a positive control, sterile petri dish diameter of 5 cm was prepared and poured into it order MHA 40-45°C temperature by 5 mL which already contains 5 mL bacterial suspension equivalent of 0.5 McFarland. While the negative control using only MHA media without bacterial suspension. Throughout the test and control media were incubated in an incubator at 37°C for 18-24 hrs.

**Comparison of antibacterial activity values**

The determination of the appeal done by way of one sterile petri dish diameter of 15 cm was prepared and put 45 mL of the bacterial suspension equivalent to a 0.5 McFarland on each cup, then added 45 mL order MHA temperature 40-45°C. The mixture was homogenized and then allowed to stand until solidified. In each cup, made 6 holes using the perforator. On hole number 1, 2, 3 into which is inserted a 100 mL solution of ethanol extract of leaves, whereas the number 4, 5, 6 holes included 100 mL solution of ethanol extract of the leaves suji. Extract solution which included an extract concentrations provide inhibitory diameter approaching the diameter of the resulting inhibition of the ethanol extract solution of other test substances at certain concentrations. Tests conducted comparative value triplo. Positive control, sterile petri dish diameter of 5 cm was prepared and poured into it MHA 40-45°C temperature by 5 mL which already contained 5 mL bacterial suspension equivalent of 0.5 McFarland. While the negative control using only MHA media without bacterial suspension. Throughout the test and control media were incubated in an incubator at 37°C for 18-24 hrs. Data diameter inhibition was observed and detected using a caliper. Then be made curve relationship between inhibition diameter (mm) on the X-axis against log concentration (ppm) on the X-axis. Furthermore, the linear regression line drawn searched the mathematical equations of the straight line.

**Determination of minimum inhibitory concentration (MIC) grow and minimum kill concentration (MBC)**

Determination of MIC grow conducted by preparing microplate and entirely filled with 100 mL of media MHB. Well first as a negative control containing only media MHB. In the second well containing MHB media and added 10 mL of the bacterial suspension equivalent to a 0.5 McFarland. Well, another containing MHB media and extract test. Extracts inserted into eppendorf tube and dissolved in DMSO to obtain the concentration of the extract solution stock. Then extract as much as 100 mL stock solution was taken and placed on a microplate well. Further dilution stratified to obtain the appropriate concentration range activity test results. Well into the whole test, inoculated 10 mL test bacteria. Microplate was incubated in an incubator at 37°C for 18-24 hrs. Growing measurement results (MIC be well with the smallest concentration which did not give a cloudiness in the test medium after an incubation period. Then from the well, was the determination of minimum kill concentration (MBC) by scraping the results of incubation
of the test medium on the surface of the solid MHA media. Positive and negative controls were also made as controls against sterility test medium and the media used. The test and control media were incubated in an incubator at 37°C for 18-24 hrs [12].

**Potassium levels in extract**
Quantitative measurement of potassium concentration was conducted to determine the content of potassium contained in the extract. Preparation of extracts made with dried destruction method and test extracts were analyzed using atomic absorption spectrophotometry (AAS).

**RESULTS AND DISCUSSION**
**Extraction and determination of parameter extract**
The extraction process is done by maceration method. 70% ethanol solvent selected for botanicals used are leaves and leaf suji is dried so that the required ethanol simplisia with large water content for the process of wetting. The water content was instrumental in the process of opening the dried simplisia pores so that the solvent would be able to log in and easily contact, dissolving and drawing secondary metabolites contained therein [13]. A viscous and thick extracts were obtained after rotary evaporated. This extract was then analyzed for its extract parameters including organoleptic, yield, and moisture content [14].

Organoleptic ethanol extract of bay leaves was blackish brown, typical aromatic extracts and thick texture. While the ethanol extract of the suji leaves the resulting green-brown, aromatic extracts and texture typical lumpy. The yield of bay and suji leaves were 13.23 and 8.04% w/w, respectively. The water content of bay and suji leaves were 8 and 5%, respectively, which met the specified conditions. The water content in the extract should not be more than 10% to avoid the rapid growth of the fungus in the extract [15].

**Phytochemical screening results**
Results of phytochemical screening botanicals and extracts from bay and suji leaves are shown in Table 1.

Our phytochemical screening results were consistent with the literature that says that the bay leaves contained secondary metabolites of tannins and flavonoids [16,17] while suji leaves contain secondary metabolites were flavonoids, and polyphenols [18]. However, both the samples and ethanol extracts bay leaf and ethanol extracts of suji leaves, not contained alkaloids, saponins, and quinones.

**TLC results**
The result of the determination of the profile TLC of ethanol extract of bay and suji leaves can be seen in Table 2.

From the results of the TLC profile, it might be concluded that there were at least five and six non-polar compounds in ethanol extract of bay and suji leaves, respectively.

### Table 1: Phytochemical screening of bay and suji leaves

| Secondary metabolites | Bay leaves | Suj leaves |
|-----------------------|------------|-----------|
| Simplisia             | Extract    | Simplisia | Extract |
| Alkaloids             | -          | -         | -       |
| Flavonoids            | +          | +         | +       |
| Tannins               | +          | +         | +       |
| Saponin               | -          | -         | -       |
| Triterpenoids         | -          | -         | +       |
| Steroids              | +          | +         |         |
| Polyphenols           | +          | +         | +       |
| Quinone               | -          | -         |         |
| Monoterpenoids & sesquiterpenoids | + | + | + |

+: Detected, -: Not detected

**Test results of confirm bacteria**
Early identification of *S. dysenteriae* ATCC 13313 was to grow them in media selectively-media Salmonella-Shigella agar. It was confirmed that *S. dysenteriae* could grow on selective media with colony morphology include color translucent colonies without black core and a spherical convex. This was consistent with *S. dysenteriae* colony morphology found in the literature [19]. Observations can be seen in Fig. 1.

Further identification by Gram-staining - When viewed under a microscope, Gram-positive bacteria in purple due to Gram-positive bacterial cell wall consists of peptidoglycan thick and will shrink by bleaching treatment/alcohol causes the pores shrink, so will retain the primary dye complex (CGV-Lugol). Gram-negative bacteria, on the other hand, have the structure of cell walls thinner and thicker lipid structures. Lipids in Gram-negative would be dissolved by the bleaching solution (alcohol), thus increasing the power seeped so under the microscope will be blue because the dye primer (CGV-Lugol) lost due to alcohol flushing and stained by the dye is methylene blue counter [20]. It was observed the test bacteria were Gram-negative because it showed the form of rods and blue corresponding to literature as shown in Fig. 2.

Furthermore, biochemical test conducted on the basis of metabolism caused by enzyme working power [19]. The results of biochemical tests can be seen in Table 3.
Table 2: TLC of bay and suji leaves ethanol extracts

| Spots | Bay leaves | Suji leaves | Bay leaves | Suji leaves |
|-------|------------|-------------|------------|-------------|
| 1     | 0.975      | 0.975       | Dark green | Dark green  |
| 2     | 0.958      |             | Yellow     | -           |
| 3     | 0.875      | 0.875       | Dark green | Dark green  |
| 4     | -          | 0.813       | -          | Light yellow|
| 5     | 0.575      | -           | Brown      | -           |
| 6     | -          | 0.463       | Light yellow| -           |
| 7     | -          | 0.35        | Light yellow| -           |
| 8     | 0.325      | -           | Brown      | -           |
| 9     | -          | 0.225       | Light yellow| -           |

- No spots, TLC: Thin layer chromatography, UV: Ultraviolet

Results of testing activities

Testing the antibacterial activity of ethanol extract of bay and suji leaves performed using agar diffusion method using a perforation technique, results of which shown in Fig. 3.

Based on this work, the ethanol extract of suji leaves began antibacterial activity at a concentration of at least 40% w/v. While the ethanol extract of bay leaves began to show antibacterial activity at a concentration of at least 10% w/v greater the concentration of the extract, the inhibition produced diameter increases. This was because, the ethanol extract of suji and bay leaves have diffusion into media so good, so the secondary metabolites contained therein were able to produce antibacterial activity against S. dysenteriae.

Results determination of comparative activities

The determination of the antibacterial comparative activity of ethanol extract of bay leaves of the ethanol extract of the suji leaves conducted to determine the best extracts activity at certain concentrations. The determination was carried out by the agar diffusion method. The results of the determination of the comparative activities can be seen in Table 4.

Based on the data in Table 4, then graphed the relationship between log concentration of the extract with a diameter of inhibition. It was found that the linear regression equation for the ethanol extract of bay leaves was y=10.217x−68.55 and for the ethanol extract of suji leaves was y=7.657x−49.77. As shown, the Table 5 indicated that at a concentration of 20% w/v, bay leaves ethanol extract provided inhibitory diameter 16.47 mm. Diameter value was then substituted at a concentration of 20% w/v, bay leaves ethanol extract provided inhibitory diameter 16.47 mm. Diameter value of which shown in Fig. 3.

Table 3: Biochemical test results of Shigella dysenteriae ATCC 13313

| Biochemical test | Shigella dysenteriae ATCC 13313 | Shigella dysenteriae (Kelly et al. 1995) |
|------------------|---------------------------------|---------------------------------------|
| Motility         | -                               | -                                     |
| Glucose          | +                               | +                                     |
| Lactose          | -                               | -                                     |
| Mannose          | -                               | -                                     |
| Malatose         | -                               | -                                     |
| Saccharose       | -                               | -                                     |
| Indol            | -                               | -                                     |
| TSIA             | -/H₂S                          | -/H₂S                                 |
| Urea             | -                               | -                                     |
| Metil red (MR)   | +                               | +                                     |
| Voges-Pnskauer (VP) | -                             | -                                     |
| Simon citrate    | -                               | -                                     |

+: React, -: Not react

Table 4: Results of determination of comparative value bay and suji ethanol extracts against Shigella dysenteriae ATCC 13313

| Concentration of bay leaves ethanol extract (%w/v) | 20 | 16.5 |
|---------------------------------------------------|----|------|
| Inhibition zone (mm)                              | 40 | 16.4 |
| Concentration of suji leaves ethanol extract (%w/v) | 16.1 |
| Inhibition zone (mm)                              | 16.2 |
| Concentration of suji leaves ethanol extract (%w/v) | 16.8 |
| Inhibition zone (mm)                              | 16.2 |
| Concentration of suji leaves ethanol extract (%w/v) | 16.47 |
| Inhibition zone (mm)                              | 16.27 |
| Concentration of suji leaves ethanol extract (%w/v) | 18.8 |
| Inhibition zone (mm)                              | 17.8 |
| Concentration of suji leaves ethanol extract (%w/v) | 18.7 |
| Inhibition zone (mm)                              | 17.4 |
| Concentration of suji leaves ethanol extract (%w/v) | 18.87 |
| Inhibition zone (mm)                              | 17.65 |
| Concentration of suji leaves ethanol extract (%w/v) | 80 |
| Inhibition zone (mm)                              | 22.8 |
| Concentration of suji leaves ethanol extract (%w/v) | 22.5 |
| Concentration of suji leaves ethanol extract (%w/v) | 22.5 |
| Inhibition zone (mm)                              | 22.6 |
| Concentration of suji leaves ethanol extract (%w/v) | 22.6 |
| Inhibition zone (mm)                              | 19.7 |

Results determination of comparative activities can be seen in Table 4.

Based on the data in Table 4, then graphed the relationship between log concentration of the extract with a diameter of inhibition. It was found that the linear regression equation for the ethanol extract of bay leaves was y=10.217x−68.55 and for the ethanol extract of suji leaves was y=7.657x−49.77. As shown, the Table 5 indicated that at a concentration of 20% w/v, bay leaves ethanol extract provided inhibitory diameter 16.47 mm. Diameter value was then substituted at a concentration of 20% w/v, bay leaves ethanol extract provided inhibitory diameter 16.47 mm. Diameter value of which shown in Fig. 3.

Determination of MIC growing and minimum kill concentration extract (MBC)

Testing grows MIC, and minimum kill concentration (MBC) were performed by microdilution method which was the method of turbidimetry with the view turbidity indicates growth of test bacteria (S. dysenteriae). The advantage of this method is the material needed microdilution less than using macrodilution. MIC was determined by observing the killing power of the smallest concentration of the extract was used to kill test bacteria. Killing ability was shown by the test medium that becomes clear in accordance with its negative. Clear controls stated that the bacteria had undergone lysis. However, turbidity difficult to observe because of the influence that disrupt observation. For this, it should be made a negative control containing media and extract to compare the turbidity of the extract or the growth of test bacteria. In addition, observations of the growth of test bacteria would clearly observe through the subculture of the results on the media to MHA sterile microplate with scratch plate method, which can simultaneously determine the value of the minimum kill concentration (MBC). MIC of both extracts could not be determined because of sediment and color of the extract that prevents bacterial growth.
**Table 5: MIC and MBC of ethanol extract of bay and suji leaves**

| Extract concentration (% b/v) | Colony growth | MIC | Extract concentration (%b/v) | Colony growth |
|------------------------------|---------------|-----|------------------------------|---------------|
|                              | Shigella dysenteriae |      |                              | Shigella dysenteriae |
| Bay leaves       | Suji leaves     |      | Bay leaves       | Suji leaves     |
| 0.625          | +             | 40.00| 5.00           | +             |
| 1.25           | +             | 20.00| 10.00          | +             |
| 2.5            | +             | 10.0 | 5.00           | +             |
| 5.00           | +             | 5.00 | +              | +             |
| 10.00          | +             | +   | +              | +             |
| 20.00          | -             | +   | +              | +             |
| 40.00          | -             | +   | +              | +             |

**MIC**: Minimum inhibitory concentration

**MBC**: Minimum kill concentration

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