

Standardization, Quantification of Total Phenolics, Flavonoids Content and Antibacterial Activity of Yellow Root (*Arcangelisia flava* (L.) Merr.)

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Abstract

Yellow root (*Arcangelisia flava* (L.) Merr.) has been used empirical in West Sumatera, Indonesia for stomach disease, anti-malaria, and tumors. The present study is to characterized non-specific, specific parameters, total phenolics content (TPC), total flavonoids content (TFC), and antibacterial activity from yellow root. Standardization of yellow root extract tests followed to Indonesian Herbal Pharmacopoeia. TPC was determined by Folin-Ciocalteu colorimetric method. TFC was calculated by aluminum chloride colometric assay, antibacterial activity was conducted by the agar diffusion method from extract and fractions against *Propionibacterium acnes* ATTC 1223. The results for non-specific parameters showed the water content of the extract (10.1%), shrinkage drying extract (12%). Specific parameters showed that extract have a specific odor, bitter in taste, brownish-yellow in color, and have a thick physical appearance. Microscopic parameters showed fragments such as fibrous sklerotic cells, parenchymal cells, exocarpic stone cells, stone cells, essential oil cells, and tracheal cells. The value of water-soluble compounds (5.25%), and the value of the ethanol-soluble (19.75%). Thin layer chromatography from extract showed the presence spots of alkaloids, phenolics, flavonoids, and saponins. *From this research we can obtained that Yellow-yellow* root extract has a total phenolic content of 87.37 mg GAE/g and flavonoid content of 73.80 mg QE/g. Extract has better antibacterial activity against *Propionibacterium acnes* than its fraction in 800 ppm with diameter inhibition 20.23 ± 0.58 mm. Standardization of yellow root extract revealed that all parameters matched the criteria requirements. *Extract has high quantities of phenolics and flavonoids. The antibacterial activity from extract was higher from fraction*

Keywords: *Arcangelisia flava* (L.) Merr.; ~~*Arcangelisia flava*~~; flavonoids; phenolics; antibacterial; standardization

Introduction

Indonesia has one of the world's most extensive tropical forest areas. The spread of Indonesian flora and fauna is very broad, and there are some varieties that are endemic, which means that these species can only grow or be discovered in one place. Indonesia's rich ethnic diversity has given rise to a wide range of cultural practices, traditions, and indigenous knowledge, including the use of local biodiversity for the treatment of various ailments (1)(1)(1)(1)(1)(1). Yellow root (*Arcangelisia flava* (L.) Merr.) belonging to Menispermaceae family, is one of the therapeutic plants being studied in this study. Yellow root is used as a jaundice treatment, anti-inflammation, anti-malaria treatment, and a cancer treatment (1-3)(1-3)(1-3)(1-3)(1-3)(1-3).

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To maintain their qualities, medicinal plants require procedures to prove identity, purity, and quality.

Herbal medicine ~~Drug~~ standardization entails confirming its identity as well as determining its quality and purity. Because of advances in crude drug chemical understanding, many methods such as botanical, chemical, spectroscopic, and biological methods are currently employed to estimate active ingredients present in crude medications in addition to physical constants ~~(4)(4)(4)(4)(4)(4)~~.

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Numerous studies have been conducted on yellow root, particularly pharmacological activity tests, (5) (Yuliana et al), but standardization of yellow root has not been conducted. Therefore, this study was conducted to determine the specific and nonspecific parameters, chromatographic patterns, and antibacterial activity of yellow root extract against the acne causing bacteria, *Propionibacterium acnes*.

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The purpose of this research was to evaluate both specific and non-specific parameters of herbal therapeutic substances in accordance with the standards of the Indonesian Herbal Pharmacopoeia. Non-specific properties like shrinkage, drying time, and water content were measured. Specific properties are extract identity and organoleptic evaluation, microscopic identification, detection of compounds dissolved in certain solvents, phytochemical screening, and chromatogram profile. This study also analyze the total phenolics content (TPC), total flavonoids content (TFC) and antibacterial activity *Propionibacterium acnes* ATCC 1223 of yellow root.

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Methods

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Materials

Arcangelisia flava radix (yellow root) were collected from Perum Perhutani Pontianak, Southeast Pontianak, Kubu Raya District, West Kalimantan in April 2021. The plant was identified by Plant Taxonomy Laboratory, Faculty of Mathematics and Natural Sciences, Universitas Padjadjaran, Indonesia, with No. No.21/HB/03/2021 which stated that the plant used was *Arcangelisia flava* (L.) Merr. ~~All chemicals were analytical grade and purchased from Merck (Germany), i.e. Mmethanol, ethanol, hydrochloride acid, acetic acid, sodium hydroxide, sodium ferric chloride, magnesium powder, and aluminium chloride and Silica gel F254 purchased from Merck (Germany).~~ Gallic acid, quercetin, caffeine was purchased from Sigma Aldrich (USA) ~~and Silica gel F254 were purchased from Merck KGaA (Germany).~~ *Propionibacterium* ATCC 1223 were obtained from the Microbiology Laboratory, Padjadjaran University. Analytical-grade sulfuric acid, barium chloride, and dimethyl

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sulfoxide were purchased from Sigma-Aldrich (Germany) and bacteriology-grade Mueller–Hinton agar (MHA) from Oxoid (UK).

Determination

The plant was identified by Plant Taxonomy Laboratory, Faculty of Mathematics and Natural Sciences, Universitas Padjadjaran, Indonesia, with No. No.21/HB/03/2021 which stated that the plant used was *Arcangelisia flava* (L.) Merr

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Extraction

Yellow root was extracted through maceration using 70% ethanol over a period of 4 x 24 hours, with the solvent being replaced every 24 hours. The extracts were concentrated using a rotary evaporator at 40–50 °C until reaching a stable weight, after which the yield was calculated ~~(5)(5)(5)(5)(5)(5)~~

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Fractionation

The viscous ethanolic extract (10 g) was mixed with distilled water to obtain a final solution volume of 100 mL. This solution underwent triple liquid–liquid extraction using both n-hexane and ethyl acetate. All extracts were then collected and evaporated ~~(7)(6)(6)(6)(6)(6)~~

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Shrinkage drying measurement

To determine drying shrinkage, 1 g of extract was introduced into a closed weighing bottle that had previously been conditioned by heating and weighed. The sample was then oven-dried at 105 °C with the bottle's lid removed. Following heating, the bottle was placed in a desiccator to allow it to return to ambient temperature. This cycle was repeated until a constant mass was recorded, with a weight variation of less than 0.0005 g between successive measurements ~~(8)(7)(7)(7)(7)(7)~~

Water content

The extract (3 g) is wrapped in aluminium foil and put in a dry round bottom flask. Fifty mL of toluene is added to the flask through a cooler (vertical condenser) and heated carefully for 60 minutes. Toluene is used to rinse the inside of the condenser, and the setup is left undisturbed until distinct toluene and water layers form. The amount of the water layer is then recorded ~~(8)(7)(7)(7)(7)(7)~~

Extract identity and organoleptic evaluation of extracts

Extract identity includes nomenclature description, plant's other names, including the Indonesian name, as well as the plants parts used. The organoleptic properties evaluated include the extract's color, smell, and taste ~~(8)(7)(7)(7)(7)(7)~~

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Microscopic identification

The powdered yellow root was positioned on a microscope slide and treated with chloral hydrate. After covering it with a cover slip, the preparation was gently heated over a Bunsen burner using tube clamps, avoiding boiling. Once dried, the slide was observed microscopically at 10x magnification ~~(9)(8)(8)(8)(8)(8)~~

Identification of compounds that are soluble in specific solvents

Two samples of yellow root extract (1.5 g each) were weighed and designated as Extract 1 and Extract 2. Extract 1 underwent 24-hour maceration in a stoppered flask with water–chloroform mixture (50 mL), whereas Extract 2 was macerated using 50 mL of 96% ethanol. Following filtration, 10.0 mL of each extract was placed in a pre-weighed container and evaporated. The residue was then dried in an oven at 105 °C to achieve a stable weight ~~(8)(7)(7)(7)(7)(7)~~

Phytochemical screening

The identification of chemical components in yellow root ethanol extract involved phytochemical testing for the presence of flavonoids, alkaloids, tannins, saponins, steroids, triterpenoids, and glycosides ~~(10)(9)(9)(9)(9)(9)~~

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Alkaloid test

On a porcelain cup, 100 mg simplicia and viscous extract placed. After that, samples are dissolved in 5 mL HCl 2 M. The obtained solution was divided into two test tubes. As a blank, three drops of HCl 2 M are added to the first tube. Three drops of Dragendorff reagent are added to the second tube. The Dragendorff reagent will form orange deposits.

Phenolic test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL hot water. Filter up the samples, add with a few drops of 10% iron (III) chloride solution. The presence of tannins is indicated by a dark blue or greenish-black color.

Flavonoids test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL of hot water. Filter up the samples, add 0.05 mg Mg powder and 1 mL concentrated HCl, and vigorously shake. Positive tests produce the colors red, yellow, or orange.

Tannin test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL of hot water. Filter up the samples, add with 1 mL of 1 % gelatine solution. The presence of tannins is indicated by white precipitate.

Saponin test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL of hot water. Filter up the samples, followed by 5 ml of water. The test tube is then vigorously shaken for 10 seconds, and one drop of HCl 2 N is added. A positive test forms a stable froth that reaches a height of 1-10 cm and remains stable for at least 10 minutes.

Steroid test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL of ether. Filter up the samples, added along with two drops of glacial CH₃COOH. The solution is gently shaken and set aside for a few minutes. Blue or green indicates steroids.

Glycosides test

On test tube, 100 mg simplicia and viscous extract placed. Add 5 mL of ethanol following evaporating. The residue added 1 mL of glacial CH₃COOH and 10 drops of concentrated H₂SO₄. Blue or green indicates steroids.

Chromatogram profile

In order to separate different phytochemicals from the ethanol extract of yellow root, the sample was applied manually with a capillary tube onto silica gel GF254 plates (15x5 cm, 3 mm thick). The plates were then placed in a mobile phase system based on Table 1 (9)(8)(8)(8)(8)(8) mobile phase system based on Karthika et al (2014) Karthika et al (2014) Karthika et al (2014) Karthika et al (2014) Karthika et al (2014) Karthika et al (2014) Karthika et al (2014) guidelines (Table 1). After Thin Layer Chromatography (TLC) development, specific visualization reagents—Dragendorff's and 5% ferric chloride—were sprayed onto the plates to detect alkaloids and phenolics, respectively (11)(10)(10)(10)(10)(10). The developed spot colors were noted, and the Rf values were calculated using formula 1 (11)(10)(10)(10)(10)(10):

$$\text{Retention factor (Rf)} = \frac{\text{distance travelled by the solute}}{\text{distance travelled by the solvent}} \quad (1)$$

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the corresponding Rf values were:

$$\text{Retention factor (Rf)} = \frac{\text{distance travelled by the solute}}{\text{distance travelled by the solvent}}$$

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Table 1: Determination of phytochemicals yellow root extract with suitable mobile phases through TLC

Compounds	Mobile Phases	Spraying Reagents	Colour of the spot
Alkaloids	Chloroform: Methanol (3:2)	Dragendorff	Orange/ Brown
Polyphenols	Chloroform: Methanol (7: 3)	FeCl ₃	Blackish blue
Flavonoids	Ethylacetate: Methanol: Water: Glacial acetic acid (1,7: 0,5: 0,5: 0,5)	FeCl ₃	Grey
Saponins	Kloroform: Metanol (2,2: 0,5)	Vanillin H ₂ SO ₄ reagent	Violet Blue

Total phenolics content (TPC)

To evaluate the total phenolics, the Folin-Ciocalteu technique according to Hanifah et al (2020)Chun et al(2003)- with modifications (12)(11)(11)(11)(11)(11) The test sample (2500 ppm in 70% ethanol) was mixed with 5 mL of the Folin-Ciocalteu reagent (diluted 1:10 with distilled water) and 4 mL of 1M sodium carbonate. After 15 minutes of incubation, the absorbance was recorded at 776.5 nm#s maximum wavelength. The absorbance of samples was calculated on standard curve through linear regression. The total phenolic content was expressed in mg GAE/g using the following formula 2.The phenolic concentration was calculated based on a standard curve obtained through linear regression.

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$$TPC (mg\ GAE/g) = \frac{c \times v \times fp}{g} \tag{2}$$

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Description:

- TPC : Total Phenolic Content (mg GAE/g)
- c : concentration of gallic acid compound (mg/L)
- v : volume of the sample used (L)
- fp : dilution factor
- g : weight of the sample (g)

Total flavonoids content (TFC)

The determination of flavonoids content was done using Chang et al (2020) method with modifications (13)(12)(12)(12)(12)(12) The sample was produced at a concentration of 5000 ppm using 70 percent ethanol. An aliquot of 0.5 mL of sample was added to a test tube containing 1.5 mL of 70% ethanol. Subsequently, 0.1 mL of 10% aluminum chloride, 0.1 mL of 1 M sodium acetate, and

2.8 mL of distilled water were added. The mixture was incubated at room temperature for 30 minutes.

Absorbance was then measured at 435.5 nm the maximum absorbance wavelength using a UV-Vis spectrophotometer, and the total flavonoid concentration was determined using a quercetin standard curve via linear regression analysis.

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The absorbance of samples was calculated on standard curve through linear regression. The total flavonoid content was expressed in mg QE/g using the following formula 3.]

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$$TFC (mg QE/g) = \frac{c \times v \times fp}{g} \quad (3)$$

Description:

TPC : Total Flavonoid Content (mg QE/g)

c : concentration of quercetin compound (mg/L)

v : volume of the sample used (L)

fp : dilution factor

g : weight of the sample (g)

Preparation of P. acnes Culture

P. acnes ATCC 1223 pure culture was inoculated onto slant MHA medium and incubated for 24 hours at 37 °C. The bacterial colonies were then suspended in 5 mL of sterile 0.9% NaCl solution. The turbidity of the bacterial suspension was standardized to the 0.5 McFarland scale, corresponding to 1.5×10^8 CFU/mL. The 0.5 McFarland standard was created by mixing 1.175% barium chloride with 1% sulfuric acid in a volumetric ratio of 0.05 to 9.95 (13)(13)(13)(13)(13)

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Antibacterial Activity Assay

A petri dish containing Mueller Hinton Agar (MHA) was inoculated with 20 µL of the bacterial suspension and left to solidify. A perforator (6 mm diameter) was used to make wells in the agar. For each well, 50 µL of solution was added: 25% clindamycin phosphate served as the positive control, 1% DMSO as the negative control, and the experimental groups included plant extract, and water, ethyl acetate, and n-hexane fractions at 800 ppm and 600 ppm concentrations. The culture plates were incubated for 18–24 hours at 37 °C, and the size of the inhibition zones was determined with a calliper (13)(13)(13)(13)(13)

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Results and discussion

From 1.5 kg of dried yellow root, a total of 151.86 g of concentrated extract was obtained, resulting in a yield of 10.12%. The appearance of yellow root simplicia and its powdered form is shown in Figure 1.

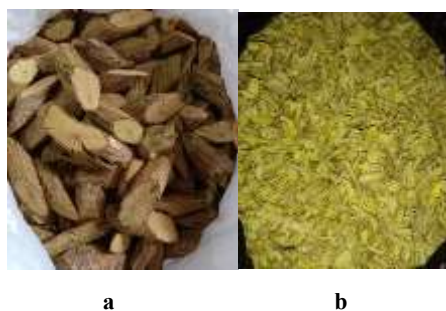


Figure 1: Yellow Root (a); Dried Yellow Root (b)

Maceration is an extraction technique where the sample is soaked in a solvent (~~menstruum~~) until it is fully immersed. After extraction, the solution (micelle) is separated from the solid residue (marc) through filtration. Evaporation is then used to remove the micelle from the menstruum ~~(15)(14)(14)(14)(14)(14)~~. This technique was chosen to protect thermolabile chemicals while also adhering to Indonesian herbal pharmacopoeia ~~(9)(8, 11+11111111)~~. Ethanol was chosen because it is polar, miscible with water, nontoxic at low concentrations, and requires only a modest quantity of heat to concentrate the extract ~~(1)(1)(1)(1)(1)(1)~~. Extract yield from this research was 10.12%. In this study, separation funnel method was used to separate the component in the extract. The solvents used for fractionation were *n*-hexane, ethyl acetate, and water. Those fractions were used to analyze antibacterial activity. Yield of each fraction showed in table 2.

Table 2: Yield Fraction from Yellow Root Extract

Fraction	Yield (%)
<i>n</i> -hexane	2.075
Ethyl acetate	5.075
Water	81.025

The fraction yield value does not reach 100%, this is likely because the natural raw material consists of various structural components and metabolites. Only certain active compounds are dissolved, while the dregs (such as fiber and complex carbohydrates) are left behind or discarded. (16). Agila Alviola Bani, Asni Amin, Abdul Mun'im, Maksun Radji.

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This study looked at non-specific factors such as shrinkage drying and moisture measurement from simplicia and extract. The drying shrinkage measurement provides insight into the amount of compounds that may be lost throughout the drying procedure. Meanwhile, the water content of simplicia and extract is related to their quality and storage capacity. The water content and shrinkage drying of simplicia do not exceed 10%, but the water content ranges from 5-30% in the thick extract category. So, according to Table 3, both simplicia and extract met the Indonesian Herbal Pharmacopoeia criteria-

Table 3: The Characterization of Non-Specific Parameters from Yellow Root

Parameters	Results (%)	
	Simplicia	Extract
Shrinkage drying	3.0	12.0
Water content	3.8	10.1

To objectively confirm the identity of a material, a specific parameter such as extract identification and organoleptic evaluation must be conducted. The parameter must be determined in order to provide objective identity from the compound's name and specific identity. The organoleptic properties of yellow root ethanolic extract were evaluated. The yellow root plant has not been included in the Indonesian herbal pharmacopoeia, so the determination of specific and non-specific parameters is only carried out based on observations by researchers. The results of the form, colour, smell, and taste tests are shown in Table 4.

~~The results of the form, colour, smell, and taste tests are shown in Table 4.~~

Table 4: The Characterization of Specific Parameters from Yellow Root Extract

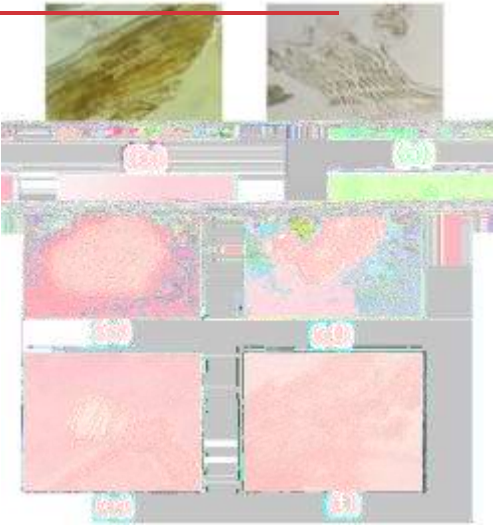
Parameters	Characteristics
Identity	
Scientific name for plants	<i>Arcangelisia flava</i> L. (Merr)
Used parts	Radix
Indonesian name plants	Akar kuning (Yellow root)
Organoleptic	
Form	Thick
Color	Brownish-yellow
Smell	Specific
Taste	Bitter

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The purpose of assessing dissolved compound levels is to estimate the amount of compounds that can be extracted using particular solvents. Water primarily dissolves polar molecules, while ethanol is effective for solubilizing both semi-polar and non-polar compounds ~~(17)(15)(15)(15)(15)(15)~~. Yellow root is more soluble in water than ethanol, as seen in Table 5. According to these observations, the compounds detected in yellow root extract are polar. This result was accordanced with the fractionation results as seen in table 2 previously.

Table 5: Parameters Levels of Dissolved Compounds in Specific Solvents

Parameter	Characteristics
Soluble ethanol compounds	5.25%
Soluble water compounds	19.75%



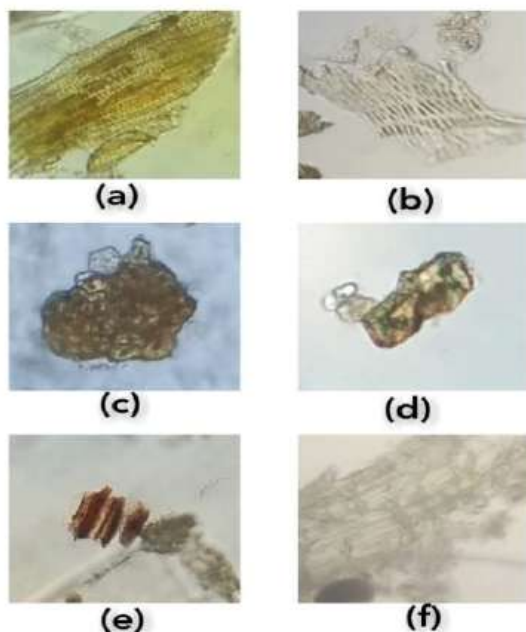


Figure 2: Microscopic from yellow root powder fragments (a) fibrous sklerotic cells; (b) parenchymal cells; (c) exocarpic stone cells; (d) stone cells; (e) essential oil cells; and (f) tracheal cells.

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The purpose of using a microscope to examine yellow root powder is to discover the properties of herb identification fragments. The chloral hydrate solution is meant to remove cell components such as protein and starch, allowing cell identification fragments on the herbs to be clearly visible under a microscope (18)(16)(16)(16)(16)(16). Figure 2 showed the presence of fibrous sclerotic cells, parenchymal cells, exocarpic stone cells, stone cells, essential oil cells, and tracheal cells in yellow root.

Table 6 showed the results of the phytochemical screening. Phytochemical screening not only reveals the components of plant extracts and which ones predominate, but it also aids in the search for chemical compounds that can be used in the development of powerful pharmaceuticals. During phytochemical screening, the colour or changes caused by a reaction with a certain response are

observed qualitatively. This screening aimed to detect and characterize secondary metabolite constituents within the simplicia and extracts.

Secondary metabolites are available in medicinal plants, and alkaloids, flavonoids, glycosides, saponins, and terpenoids are among several bioactive substances present. The chromatographic method is widely used for isolating components from plants. Thin-layer chromatography (TLC) was utilized to distinguish secondary metabolites such as alkaloids, phenolics, flavonoids, and saponins in yellow root ethanolic extract. Various mobile phase formulations were tested (see Table 1), and the samples were spotted onto TLC plates for analysis. Following derivatization with specific spraying reagents, color development was observed, indicating the presence of alkaloids, phenolics, flavonoids, and saponins. These compounds exhibited varying R_f values across different solvent systems in the TLC analysis of ethanolic yellow root extract (Figure 3).

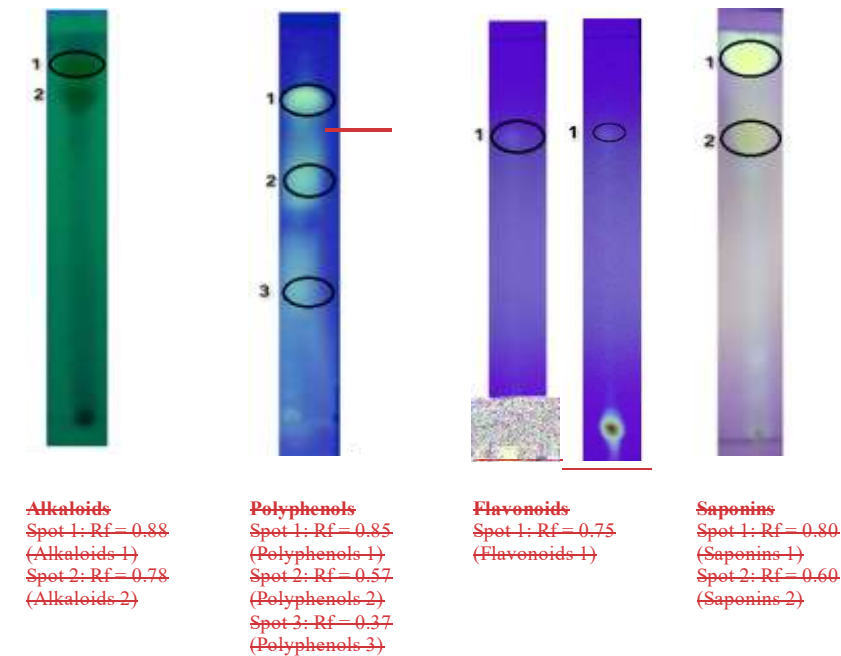
Table 6: Results of Identification Bioactive Compounds from Yellow Root

Phytochemical Test	Simplicia	Extract
Alkaloids	+	+
Phenolics	+	+
Flavonoids	+	+
Tannins	-	-
Saponins	+	+
Steroids	-	-
Glycosides	-	-

*(+) = positive test (exist); (-) = negative test (not exist)

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Figure 3: Rf Values of Yellow Root Extract on Various Secondary Metabolites

Table 7: Rf Value of Yellow Root Extract on Various Secondary Metabolites

Metabolites	Number of Spot	Rf Value
Alkaloids	1	0.88
	2	0.78
Polyphenols	1	0.85
	2	0.57
	3	0.37
Flavonoids	1	0.75
Saponins	1	0.80
	2	0.60

Total phenolic content (TPC) is frequently determined using the Folin-Ciocalteu assay, a well-established analytical technique. Since high phenolic levels are often linked to strong antioxidant properties, this assay plays a key role in evaluating overall antioxidant potential. The Folin-Ciocalteu reaction involves electron transfer from phenolic compounds and other reductants to phosphomolybdic-phosphotungstic acid complexes, forming a blue complex with peak absorbance (λ_{max}) between 700 - 800 nm. As a result, the absorption spectra of yellow root extract were compared to that of gallic acid, a reference component (20)(18)(18)(18)(18)(18). The calibration curve for acid gallic was obtained from the study data, as shown in Figure 4. Using the gallic acid

calibration curve with the linear regression model $y = 0.0857x + 0.2997$ ($R^2 = 0.9945$), the total phenolic content (TPC) of yellow root extract was determined to be 87.372.280 mg gallic acid equivalents (GAE) per gram of dried extract (Table 7).

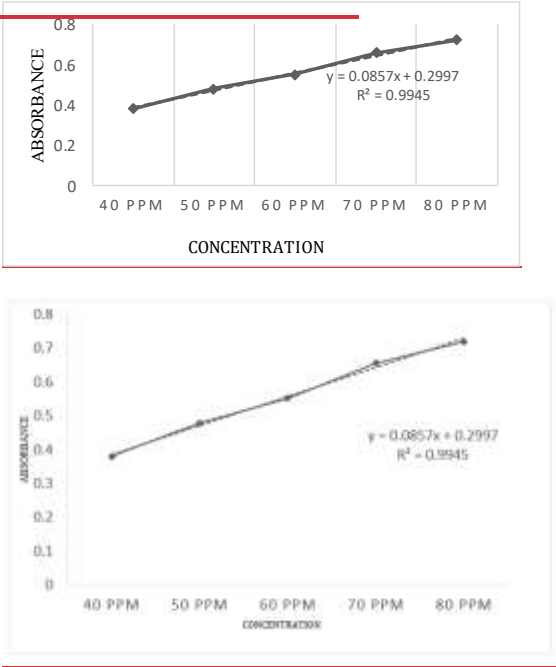


Figure 4: Calibration Curve of Gallic Acid (n=3)

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Table 7: Results of TPC on Yellow Root Extract

Sample	Replication	Concentration (ppm)	Absorbance	Total Phenolics Content (mg GAE/g)	Average of Total Phenolics Content (mg GAE/g)±SD
Extract	1	2500	0.317269	2.52386.92	2.28087.37±0.971011
	2		0.321	3.10787.72	
	3		0.308	1.21187.52	

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Pls give more explanation of this, as you mentioned in the paragraphs using this linier regresion for determining TPC

Total flavonoid content (TFC) was determined using a colorimetric technique with quercetin as a reference flavonoid component ~~(21)(19)(19)(19)(19)~~ TFC in the extracts were calculated from the regression equation of the calibration curve ($y=0.0753x+0.0082$; $R^2=0.997$) as shown in Fig. 5 and expressed as mg quercetin equivalents (QE) each gram of sample in dry weight (mg/g). The

absorption of quercetin was measured at a maximum wavelength of 435 nm. The flavonoids concentration of the leaf extract was determined to be 73.871.071 mg QE/g, as indicated in Table 8.

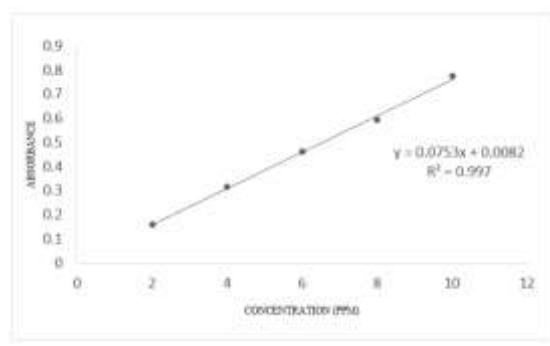
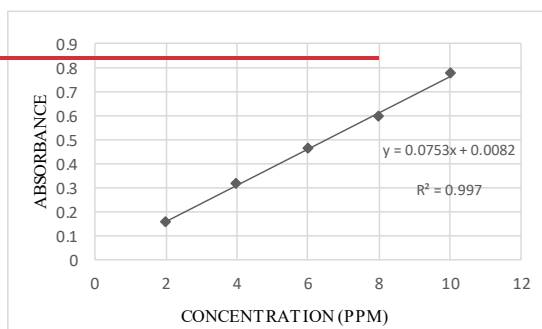


Figure 5: Calibration Curve of Quercetin (n=3)

Table 8: Results of TFC on Yellow root Extract

Sample	Replication	Concentration (ppm)	Absorbance	Flavonoids content (mg QE/g)	Average Flavonoids content (mg QE/g)±SD
Extract	1	2500	0.438	<u>74.0871.348</u>	<u>73.8071.071±0.002253</u>
	2		0.436	<u>73.7471.016</u>	
	3		0.435	<u>73.5670.850</u>	

Clindamycin phosphate, employed in this study as a positive control, is approved by the U.S. Food and Drug Administration for treating conditions such as sepsis, intra-abdominal and lower respiratory tract infections, gynecological infections, and infections of the skin, bones, and joints. The antibiotic may exhibit bacteriostatic or bactericidal properties, contingent upon the organism, infection location, and concentration. The predominant bacteria in the pilosebaceous unit of normal human skin is *P.*

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acnes, a Gram-positive anaerobic bacterium. *P. acnes* is linked to several disorders, including skin diseases, and is the most frequent source of opportunistic infections, such as acne vulgaris (13)(13)(13)(13)(13)(13).

The inhibition zones for *P. acnes* were significantly affected by the concentration of both extract and fraction (p-value = 1.02×10^{-4}). A positive correlation was noted between concentration and inhibition diameter, as presented in Table 9. The differences in antibacterial activity may be due to variations in the secondary metabolites of yellow root contained within the extract and fraction, influencing their effectiveness against *P. acnes* (22)(20)(20)(20)(20)(20).

Table 9: Antibacterial Activity of Extract and Fractions against *P. acnes* (n = 3)

Sample	Inhibition Zone (mm)±SD	
	Concentration 800 (ppm)	Concentration 600 (ppm)
Ethanol extract	20.53±0.58	18.90±0.62
n-hexane fraction	14.83±0.50	13.26±0.05
Ethyl Acetate fraction	15.93±0.35	14.46±0.66
Water fraction	18.60±0.20	17.16±0.66
Clindamycin phosphate (10%)	21.90±0.40	21.36±0.80

Conclusions

From this research we can obtained that yellow root extract has a total phenolic content of 87.37 mg GAE/g and flavonoid content of 73.80 mg QE/g. Extract has better antibacterial activity against *Propionibacterium acnes* than its fraction in 800 ppm with diameter inhibition 20.23 ± 0.58 mm. Standardization of yellow root extract revealed that all parameters matched the criteria requirements. Extract has high quantities of phenolics and flavonoids. The antibacterial activity from extract was higher from fraction.

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Conflict of interest

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Authors declare no conflict of interests

References

1. [Diliarosta S, Sudarmin S S, Efendi A, Dillasamola D, Oktomalioputri B, Ramadhani R. Reconstruction and Scientific Explanation of Akar Kuning \(Arcangelisia flava Merr.\) From West Sumatra as Ethnomedicine and Source of Science Learning. Pharmacognosy Journal. 2021 Jan 8;13\(1\):206–11. doi:10.5530/pj.2021.13.29](#)

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3. [Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning \(Arcangelisia flava L\) Stems from West Kalimantan. Jurnal Biologi Tropis. 2022;22\(4\):1334–9.](#)

4. [Sakshi Y. Pathrikar, Vaishnavi S. Mukhmale, Nandakishor B. Deshmukh, Swati P. Deshmukh. Standardization extraction and phytochemical screening for herbal drugs. GSC Advanced Research and Reviews. 2025 Jan 30;22\(1\):084–91. doi:10.30574/gscarr.2025.22.1.0518](#)

5. [Yabansabra Y, Bowaire A, Simaremare ES, Yaam DY, Tiris A, Nadeak ESM. Phytochemical and Pharmacological Review of Arcangelisia flava \(L.\) Merr: Insights into Its Bioactive Compounds and Therapeutic Potential. JURNAL BIOLOGI PAPUA. 2025;17\(1\):82–92.](#)

6. [Saptarini NM, Mustarichie R, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of Pereskia bleo \(Kunth\) DC. International Journal of Applied Pharmaceutics. 2022 Nov 26;106–10. doi:10.22159/ijap.2022.v14s4.PP19](#)

7. [Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy \(Hemigraphis colorata Hall. F.\) leaf using 1, 1-diphenyl-2-picrylhydrazyl. Drug Invention Today. 2018;10\(5\):3791–3.](#)

8. [Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.](#)

9. [Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.](#)

10. [Nur RM, Dewi R, Kaliu S. Antibacterial activity of methanol extract Rhizophora mucronata leaves toward Salmonella typhi: leading the typhoid fever. Pharmacia. 2022 Nov 14;12\(3\):364. doi:10.12928/pharmacia.v12i3.22475](#)

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11. [Karthika, S J, S P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of Solena amplexicaulis. Journal of Pharmacognosy and Phytochemistry JPP. 2014;3\(31\):198–206.](#)
12. [Hanifah HN, Hadisoebroto G, Dewi L. Comparison of phenolic, flavonoid, and tannin contents from ethanol extract of Kratom stem \(Mitragyna speciosa Korth.\) and senggani flower \(Melastoma malabathrium L.\). J Phys Conf Ser. 2021 Apr 1;1869\(1\):012002. doi:10.1088/1742-6596/1869/1/012002](#)
13. [Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colometric methods. J Food Drug Anal. 2020 Jul 14;10\(3\). doi:10.38212/2224-6614.2748](#)
14. [Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig \(Ficus carica L.\) From Ciwidey Distric, West Java, Indonesia. RASAYAN Journal of Chemistry. 2022;\(Special Issue\):172–9. doi:10.31788/RJC.2022.1558205](#)
15. [Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation Procedures for Experimental Purposes. J Pharm Bioallied Sci. 2020 Jan;12\(1\):1–10. doi:10.4103/jpbs.JPBS 175 19](#)
16. [Bani AA, Amin A, Mun'im A, Radji M. Rasio Nilai Rendamen dan Lama Ekstraksi Maserat Etanol Daging Buah Burahol \(Stelecocharpus burahol\) Berdasarkan Cara Preparasi Simplisia. Makassar Natural Product Journal. 2023;1\(3\):176–84.](#)
17. [Muhtadi M, Ningrum U. Standardization of durian fruit peels \(Durio zibethinus Murr.\) extract and antioxidant activity using DPPH method. Pharmacia. 2019 Nov 30;9\(2\):271. doi:10.12928/pharmacia.v9i2.12652](#)
18. [Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction \(Moringa oleifera L.\) . Indonesian Journal of Pharmaceutical Science and Technology. 2021;1\(1\):66–74.](#)
19. [Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumpit mutiara \(Oldelandia corymbosa L.\). Pharmacia. 2025 Jul 31;15\(2\):332–41. doi:10.12928/pharmacia.v15i2.27939](#)
20. [Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of Commiphora mollis \(Oliv.\) Engl. resin. BMC Chem. 2022 Dec 25;16\(1\):48. doi:10.1186/s13065-022-00841-x](#)
21. [Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SI, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethyl acetate extracts from accessions of Amomum compactum fruits. Annals of Agricultural Sciences. 2021 Jun;66\(1\):58–62. doi:10.1016/j.aos.2021.04.001](#)

22. [Khotimah TK, Krisridwany A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of Hylocereus Polyrhizus Peel Against E. Coli and S. aureus. Journal of Fundamental and Applied Pharmaceutical Science. 2021 Apr 23;1\(2\). doi:10.18196/jfaps.v1i2.11001](#)
1. [Diliaresta S, Sudarmin S S, Efendi A, Dillasamola D, Oktomalioputri B, Ramadhani R. Reconstruction and Scientific Explanation of Akar Kuning \(Arcangelisia flava Merr.\) From West Sumatra as Ethnomedicine and Source of Science Learning. Pharmacognosy Journal. 2021 Jan 8;13\(1\):206–11. doi:10.5530/pj.2021.13.29](#)
2. [Saepudin S, Hidayat TS, Al-Azzahra Y, Cahyati A. Uji Aktivitas Antiinflamasi Ekstrak Batang Akar Kuning \(Arcangelisia flava \(L.\) Merr.\) Secara In Vitro. Jurnal Buana Farma. 2024 Mar 26;4\(1\):1–10. doi:10.36805/jbf.v4i1.934](#)
3. [Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning \(Arcangelisia flava L\) Stems from West Kalimantan. Jurnal Biologi Tropis. 2022;22\(4\):1334–9.](#)
4. [Sakshi Y, Pathrikar, Vaishnavi S, Mukhale, Nandakishor B, Deshmukh, Swati P, Deshmukh. Standardization extraction and phytochemical screening for herbal drugs. GSC Advanced Research and Reviews. 2025 Jan 30;22\(1\):084–91. doi:10.30574/gscarr.2025.22.1.0518](#)
5. [Saptarini NM, Mustarichie R, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of Pereskia bleo \(Kunth\) DC. International Journal of Applied Pharmaceutics. 2022 Nov 26;106–10. doi:10.22159/ijap.2022.v14s4.PP19](#)
6. [Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy \(Hemigraphis colorata Hall. F.\) leaf using 1, 1-diphenyl-2-picrylhydrazyl. Drug Invention Today. 2018;10\(5\):3791–3.](#)
7. [Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.](#)
8. [Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.](#)
9. [Nur RM, Dewi R, Kalis S. Antibacterial activity of methanol extract Rhizophora mucronata leaves toward Salmonella typhi: leading the typhoid fever. Pharmacia. 2022 Nov 14;12\(3\):364. doi:10.12928/pharmacia.v12i3.22475](#)
10. [Karthika, S J, S P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of Solena amplexicaulis. Journal of Pharmacognosy and Phytochemistry JPP. 2014;3\(31\):198–206.](#)
11. [Hanifah HN, Hadisoebroto G, Dewi L. Comparison of phenolic, flavonoid, and tannin contents from ethanol extract of Kratom stem \(Mitragyna speciosa Korth.\) and senggani flower \(Melastoma malabathrium L.\). J Phys Conf Ser. 2021 Apr 1;1869\(1\):012002. doi:10.1088/1742-6596/1869/1/012002](#)

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12. [Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colometric methods. J Food Drug Anal. 2020 Jul 14;10\(3\). doi:10.38212/2224-6614.2748.](#)
13. [Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig \(Ficus carica L.\) From Ciwidey Distric, West Java, Indonesia. RASAYAN Journal of Chemistry. 2022;\(Special Issue\):172–9. doi:10.31788/RJC.2022.1558205.](#)
14. [Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation Procedures for Experimental Purposes. J Pharm Bioallied Sci. 2020 Jan;12\(1\):1–10. doi:10.4103/jpbs.JPBS-175-19.](#)
15. [Muhtadi M, Ningrum U. Standardization of durian fruit peels \(Durio zibethinus Murr.\) extract and antioxidant activity using DPPH method. Pharmacia. 2019 Nov 30;9\(2\):271. doi:10.12928/pharmacia.v9i2.12652.](#)
16. [Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction \(Moringa oleifera L.\). Indonesian Journal of Pharmaceutical Science and Technology. 2021;1\(1\):66–74.](#)
17. [Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumput mutiara \(Oldelandia corymbosa L.\). Pharmacia. 2025 Jul 21;15\(2\):332–41. doi:10.12928/pharmacia.v15i2.27939.](#)
18. [Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of Commiphora mollis \(Oliv.\) Engl. resin. BMC Chem. 2022 Dec 25;16\(1\):48. doi:10.1186/s13065-022-00841-x.](#)
19. [Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SI, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethyl acetate extracts from accessions of Amomum compactum fruits. Annals of Agricultural Sciences. 2021 Jun;66\(1\):58–62. doi:10.1016/j.aos.2021.04.001.](#)
20. [Khotimah TK, Krisridwany A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of Hylocereus Polyrhizus Peel Against E. Coli and S. aureus. Journal of Fundamental and Applied Pharmaceutical Science. 2021 Apr 23;1\(2\). doi:10.18196/jfaps.v1i2.11001.](#)
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2. [Saepudin S, Hidayat TS, Al-Azzahra Y, Cahyati A, Uji Aktivitas Antiinflamasi Ekstrak Batang Akar Kuning \(Arcangelisia flava \(L.\) Merr.\) Secara In Vitro. Jurnal Buana Farma. 2024 Mar 26;4\(1\):1–10. doi:10.36805/jbf.v4i1.934.](#)
3. [Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning \(Arcangelisia flava L.\) Stems from West Kalimantan. Jurnal Biologi Tropis. 2022;22\(4\):1334–9.](#)
4. [Sakshi Y, Pathrikar, Vaishnavi S, Mukhale, Nandakishor B, Deshmukh, Swati P, Deshmukh. Standardization extraction and phytochemical screening for herbal drugs.](#)

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- GSC Advanced Research and Reviews. 2025 Jan 30;22(1):084–91. doi:10.30574/gscarr.2025.22.1.0518
5. Saptarini NM, Mustarichie P, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of Pereskia bleo (Kunth) DC. International Journal of Applied Pharmaceutics. 2022 Nov 26;106–10. doi:10.22159/ijap.2022.v14s4.PP19
 6. Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy (Hemigraphis colorata Hall. F.) leaf using 1, 1-diphenyl 2-picrylhydrazyl. Drug Invention Today. 2018;10(5):3791–3.
 7. Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.
 8. Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.
 9. Nur RM, Dewi R, Kaliu S. Antibacterial activity of methanol extract Rhizophora mucronata leaves toward Salmonella typhi: leading the typhoid fever. Pharmacia. 2022 Nov 14;12(3):364. doi:10.12928/pharmacia.v12i3.22475
 10. Karthika, S J, S P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of Solena alexicaulis. Journal of Pharmacognosy and Phytochemistry JPP. 2014;3(31):198–206.
 11. Hanifah HN, Hadisoebroto G, Dewi L. Comparison of phenolic, flavonoid, and tannin contents from ethanol extract of Kratom stem (Mitragyna speciosa Korth.) and senggani flower (Melastoma malabathrium L.). J Phys Conf Ser. 2021 Apr 1;1869(1):012002. doi:10.1088/1742-6596/1869/1/012002
 12. Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colometric methods. J Food Drug Anal. 2020 Jul 14;10(3). doi:10.38212/2224-6614.2748
 13. Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig (Ficus carica L.) From Ciwidey Distric, West Java, Indonesia. RASAYAN Journal of Chemistry. 2022;(Special Issue):172–9. doi:10.31788/RJC.2022.1558205
 14. Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation Procedures for Experimental Purposes. J Pharm Bioallied Sci. 2020 Jan;12(1):1–10. doi:10.4103/jpbs.JPBS_175_19
 15. Muhtadi M, Ningrum U. Standardization of durian fruit peels (Durio zibethinus Murr.) extract and antioxidant activity using DPPH method. Pharmacia. 2019 Nov 30;9(2):271. doi:10.12928/pharmacia.v9i2.12652
 16. Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction (Moringa oleifera L.). Indonesian Journal of Pharmaceutical Science and Technology. 2021;1(1):66–74.
 17. Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumput mutiara (Oldelandia corymbosa L.). Pharmacia. 2025 Jul 31;15(2):332–41. doi:10.12928/pharmacia.v15i2.27939

18. [Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of Commiphora mollis \(Oliv.\) Engl. resin. BMC Chem. 2022 Dec 25;16\(1\):48. doi:10.1186/s13065-022-00841-x](#)
19. [Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SI, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethyl acetate extracts from accessions of Amomum compactum fruits. Annals of Agricultural Sciences. 2021 Jun;66\(1\):58–62. doi:10.1016/j.aas.2021.04.001](#)
20. [Khotimah TK, Krisidwani A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of Hylocereus Polyrhizus Peel Against E. Coli and S. aureus. Journal of Fundamental and Applied Pharmaceutical Science. 2021 Apr 22;1\(2\). doi:10.18196/jfaps.v1i2.11001](#)
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3. [Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning \(Arcangelisia flava L\) Stems from West Kalimantan. Jurnal Biologi Tropis. 2022;22\(4\):1334–9.](#)
4. [Sakshi Y, Pathrikar, Vaishnavi S, Mukhmal, Nandakishor B, Deshmukh, Swati P, Deshmukh. Standardization extraction and phytochemical screening for herbal drugs. GSC Advanced Research and Reviews. 2025 Jan 30;22\(1\):084–91. doi:10.30574/gscarr.2025.22.1.0518](#)
5. [Saptarini NM, Mustarichie R, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of Pereskia bleo \(Kunth\) DC. International Journal of Applied Pharmaceutics. 2022 Nov 26;106–10. doi:10.22159/ijap.2022.v14s4.PP19](#)
6. [Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy \(Hemigraphis colorata Hall. F.\) leaf using 1, 1 diphenyl 2 picrylhydrazyl. Drug Invention Today. 2018;10\(5\):3791–3.](#)
7. [Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.](#)
8. [Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.](#)
9. [Nur RM, Dewi R, Kalii S. Antibacterial activity of methanol extract Rhizophora mucronata leaves toward Salmonella typhi: leading the typhoid fever. Pharmacia. 2022 Nov 14;12\(3\):364. doi:10.12928/pharmacia.v12i3.22475.](#)
10. [Karthika, S J, S P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of Solena amplexicaulis. Journal of Pharmacognosy and Phytochemistry JPP. 2014;3\(31\):198–206.](#)
11. [Chun OK, Kim DO, Lee CY. Superoxide radical scavenging activity of the major polyphenols in fresh plums. J Agric Food Chem. 2003;51\(27\):8067–72.](#)

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12. [Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colorimetric methods. J Food Drug Anal. 2020 Jul 14;10\(3\). doi:10.38212/2224-6614.2748](#)
13. [Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig \(Ficus carica L.\) From Ciwidey Distric, West Java, Indonesia. RASAYAN Journal of Chemistry. 2022;\(Special Issue\):172–9. doi:10.31788/RJC.2022.1558205](#)
14. [Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation Procedures for Experimental Purposes. J Pharm Bioallied Sci. 2020 Jan;12\(1\):1–10. doi:10.4103/jpbs.JPBS_175_19](#)
15. [Muhtadi M, Ningrum U. Standardization of durian fruit peels \(Durio zibethinus Murr.\) extract and antioxidant activity using DPPH method. Pharmaciana. 2019 Nov 30;9\(2\):271. doi:10.12928/pharmaciana.v9i2.12652](#)
16. [Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction \(Moringa oleifera L.\) . Indonesian Journal of Pharmaceutical Science and Technology. 2021;1\(1\):66–74.](#)
17. [Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumpit mutiara \(Oldelandia corymbosa L.\). Pharmaciana. 2025 Jul 31;15\(2\):332–41. doi:10.12928/pharmaciana.v15i2.27939](#)
18. [Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of Commiphora mollis \(Oliv.\) Engl. resin. BMC Chem. 2022 Dec 25;16\(1\):48. doi:10.1186/s13065-022-00841-x](#)
19. [Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SJ, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethyl acetate extracts from accessions of Amomum compactum fruits. Annals of Agricultural Sciences. 2021 Jun;66\(1\):58–62. doi:10.1016/j.aas.2021.04.001](#)
20. [Khotimah TK, Krisnidwary A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of Hylocereus Polyrhizus Peel Against E. Coli and S. aureus. Journal of Fundamental and Applied Pharmaceutical Science. 2021 Apr 23;1\(2\). doi:10.18196/jfaps.v1i2.11001](#)
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2. [Saepudin S, Hidayat TS, Al-Azzahra Y, Cahyati A. Uji Aktivitas Antiinflamasi Ekstrak Batang Akar Kuning \(Arcangelisia flava \(L.\) Merr.\) Secara In Vitro. Jurnal Buana Farma. 2024 Mar 26;4\(1\):1–10. doi:10.36805/jbf.v4i1.934](#)
3. [Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning \(Arcangelisia flava L\) Stems from West Kalimantan. Jurnal Biologi Tropis. 2022;22\(4\):1334–9.](#)
4. [Pandey A, Tripathi S. Concept of standardization, extraction and pre phytochemical screening strategies for herbal drug. J Pharmacogn Phytochem. 2014;2\(5\):115–9.](#)

Formatted: Fontcolor: Black

5. [Saptarini NM, Mustarichie R, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of Pereskia bleo \(Kunth\) DC. International Journal of Applied Pharmaceutics. 2022 Nov 26;106–10. doi:10.22159/ijap.2022.v14s4.PP19](#)
6. [Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy \(Hemigraphis colorata Hall. F.\) leaf using 1, 1-diphenyl 2-picrylhydrazyl. Drug Invention Today. 2018;10\(5\):3791–3.](#)
7. [Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.](#)
8. [Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.](#)
9. [Nur RM, Dewi R, Kaliu S. Antibacterial activity of methanol extract Rhizophora mucronata leaves toward Salmonella typhi: leading the typhoid fever. Pharmacia. 2022 Nov 14;12\(3\):364. doi:10.12928/pharmacia.v12i3.22475](#)
10. [Karthika, S.J, S.P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of Solena amplexicaulis. Journal of Pharmacognosy and Phytochemistry JPP. 2014;3\(31\):198–206.](#)
11. [Chun OK, Kim DO, Lee CY. Superoxide radical scavenging activity of the major polyphenols in fresh plums. J Agric Food Chem. 2003;51\(27\):8067–72.](#)
12. [Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colorimetric methods. J Food Drug Anal. 2020 Jul 14;10\(3\). doi:10.38212/2224-6614.2748](#)
13. [Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig \(Ficus carica L.\) From Ciwidey Distric, West Java, Indonesia. RASAYAN Journal of Chemistry. 2022;\(Special Issue\):172–9. doi:10.31788/RJC.2022.1558205](#)
14. [Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation. Procedures for Experimental Purposes. J Pharm Bioallied Sci. 2020 Jan;12\(1\):1–10. doi:10.4103/ipbs.JPBS. 175–19.](#)
15. [Muhtadi M, Ningrum U. Standardization of durian fruit peels \(Durio zibethinus Murr.\) extract and antioxidant activity using DPPH method. Pharmacia. 2019 Nov 30;9\(2\):271. doi:10.12928/pharmacia.v9i2.12652](#)
16. [Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction \(Moringa oleifera L.\). Indonesian Journal of Pharmaceutical Science and Technology. 2021;1\(1\):66–74.](#)
17. [Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumput mutiara \(Oldelandia corymbosa L.\). Pharmacia. 2025 Jul 31;15\(2\):332–41. doi:10.12928/pharmacia.v15i2.27939](#)
18. [Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of Commiphora mollis \(Oliv.\) Engl. resin. BMC Chem. 2022 Dec 25;16\(1\):48. doi:10.1186/s13065-022-00841-x](#)
19. [Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SI, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethyl acetate extracts from accessions of Amomum compactum fruits. Annals of Agricultural Sciences. 2021 Jun;66\(1\):58–62. doi:10.1016/j.aas.2021.04.001](#)
20. [Khotimah TK, Krisdriwany A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of Hylocereus Polyrhizus Peel Against E. Coli and S. aureus. Journal of Fundamental and Applied Pharmaceutical Science. 2021 Apr 23;1\(2\). doi:10.18196/jfaps.v1i2.11001](#)

1. Diliarosta S, Sudarmin SS, Efendi A, Dillasamola D, Oktomalioputri B, Ramadhani R. Reconstruction and Scientific Explanation of Akar Kuning (*Arcangelisia flava* Merr.) From West Sumatra as Ethnomedicine and Source of Science Learning. *Pharmacognosy Journal*. 2021 Jan 8;13(1):206–11.
2. Saepudin S, Hidayat TS, Al Azzahra Y, Cahyati A. Uji Aktivitas Antiinflamasi Ekstrak Batang Akar Kuning (*Arcangelisia flava* (L.) Merr.) Secara In Vitro. *Jurnal Buana Farma*. 2024 Mar 26;4(1):1–10.
3. Tavita GE, Lestari D, Linda R, Apindiati RK, Rafdinal R. Phytochemical Testing and In Vitro Anti-inflammatory Activity on Ethanol Extract of Akar Kuning (*Arcangelisia flava* L) Stems from West Kalimantan. *Jurnal Biologi Tropis*. 2022;22(4):1334–9.
4. Pandey A, Tripathi S. Concept of standardization, extraction and pre phytochemical screening strategies for herbal drug. *J Pharmacogn Phytochem*. 2014;2(5):115–9.
5. Saptarini NM, Mustarichie R, Herawati IE, Hadisoebroto G. Isolation, Identification, and Quantification of Major Flavonoid In Leaves of *Pereskia bleo* (Kunth) DC. *International Journal of Applied Pharmaceutics*. 2022 Nov 26;106–10.
6. Herawati IE, Hanifah HN. Antioxidant activity from ethanol extract and fractions of red flame ivy (*Hemigraphis colorata* Hall. f.) leaf using 1, 1-diphenyl-2-picrylhydrazyl. *Drug Invention Today*. 2018;10(5):3791–3.
7. Departemen Kesehatan. Parameter Standar Umum Ekstrak Tumbuhan Obat. 2000. 77 p.
8. Kemenkes RI. Farmakope Herbal Indonesia Edisi Kedua. 2nd ed. Jakarta: Departemen Kesehatan Republik Indonesia; 2017. 209–213 p.
9. Novaryatiin S. Phytochemical Screening and Antibacterial Activity of Bawang Dayak (*Eleutherine* sp.) and Hati Tanah (*Angiopteris* sp.) and Their Combination Against *Propionibacterium acnes*. *International Journal of Applied Pharmaceutics*. 2019 Jul 10;11–3.
10. Karthika, S J, S P. TLC and HPTLC Fingerprint Profiles of Different Bioactive Components from the Tuber of *Solena amplexicaulis*. *Journal of Pharmacognosy and Phytochemistry JPP*. 2014;3(31):198–206.
11. Chun OK, Kim DO, Lee CY. Superoxide radical scavenging activity of the major polyphenols in fresh plums. *J Agric Food Chem*. 2003;51(27):8067–72.
12. Chang CC, Yang MH, Wen HM, Chern JC. Estimation of total flavonoid content in propolis by two complementary colorimetric methods. *J Food Drug Anal*. 2020 Jul 14;10(3).
13. Saptarini NM, R. Mustarichie, Aulifa DL, R. Hendriani, Herawati IE. Analysis of Antioxidant and Antibacterial Activity of Leaves Of Fig (*Ficus carica* L.) From Ciwidey Distric, West Java, Indonesia. *RASAYAN Journal of Chemistry*. 2022;(Special Issue):172–9.
14. Abubakar AR, Haque M. Preparation of Medicinal Plants: Basic Extraction and Fractionation Procedures for Experimental Purposes. *J Pharm Bioallied Sci*. 2020 Jan;12(1):1–10.
15. Muhtadi M, Ningrum U. Standardization of durian fruit peels (*Durio zibethinus* Murr.) extract and antioxidant activity using DPPH method. *Pharmaciana*. 2019 Nov 30;9(2):271.
16. Fatmawati A, Sucianingsih D, Kurniawati R, Abdurrahman M. Microscopic Identification and Determination of Total Flavonoid Content of Moringa Leaves Extract and Ethyl Acetate Fraction (*Moringa oleifera* L.). *Indonesian Journal of Pharmaceutical Science and Technology*. 2021;1(1):66–74.
17. Herawati IE, Saepudin S. Analysis of antioxidant and standardization of ethanol extract of rumpun mutiara (*Oldelandia corymbosa* L.). *Pharmaciana*. 2025 Jul 31;15(2):332–41.
18. Molole GJ, Gure A, Abdissa N. Determination of total phenolic content and antioxidant activity of *Commiphora mollis* (Oliv.) Engl. resin. *BMC Chem*. 2022 Dec 25;16(1):48.

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19. Nurcholis W, Sya'bani Putri DN, Husnawati H, Aisyah SI, Priosoeryanto BP. Total flavonoid content and antioxidant activity of ethanol and ethylacetate extracts from accessions of *Amomum compactum* fruits. *Annals of Agricultural Sciences*. 2021 Jun;66(1):58–62.
20. Khotimah TK, Krisnidwary A, Orbayinah S, Harimurti S. Antibacterial Activity of Fractions from Extract Ethanolic of *Hylocereus Polyrhizus* Peel Against *E. Coli* and *S. aureus*. *Journal of Fundamental and Applied Pharmaceutical Science*. 2021 Apr 23;1(2).