



# Plagiarism Checker X Originality Report

**Similarity Found: 6%**

Date: Wednesday, February 24, 2021

Statistics: 203 words Plagiarized / 3568 Total words

Remarks: Low Plagiarism Detected - Your Document needs Optional Improvement.

---

/

INTRODUCTION Soap is a cleanser because it can remove dirt that sticks to parts of the body<sup>1</sup>. The use of liquid soap is more attractive to the public than solid soap because it is more practical, economical, not contaminated, easy to carry, and easy to store<sup>2</sup>. There have been many antibacterial soaps circulated in the market under various brand names.

Most of the antibacterial soap in the market still contains synthetic ingredients such as sodium lauryl sulfate (SLS) and triclosan, which have adverse effects on human skin. These side effects include sensitivity to the skin and turning off the protective layer on the skin to be more susceptible to exposure to harmful bacteria on the skin<sup>3,4</sup>. One of the efforts to overcome this problem is to utilize plants with antibacterial properties, one of which is ketapang (*Terminalia catappa* L.).

*Terminalia catappa* comes from the Combretaceae family, which is a large tree<sup>5</sup> that has horizontal branches with several levels, leaves of 15-25 cm long and 10-14 cm wide<sup>6</sup>. This plant is widely distributed in countries with tropical and sub-tropical climates, especially in coastal areas<sup>7</sup>. *Terminalia catappa* are often found on roadsides as decoration and shade trees<sup>8</sup>. *Terminalia catappa* shed their leaves every day, and most of them fall during the dry season<sup>9</sup>. *Terminalia catappa* is known as a plant with pharmacological effects and is used traditionally<sup>6</sup>. In Asian countries, T.

*catappa* leaves are usually used to treat dermatitis, hepatitis, diarrhea, and paresis. This plant is also included in the type of vegetable in the Caribbean, where T. *catappa* leaves are used in decoction to treat ulcers and urinary tract infections<sup>7</sup>. In India, T. *catappa* leaves are attached to the skin to treat scabies, leprosy wounds, and other skin diseases. Besides, in Malaysia, T. *catappa* leaves are used to treat diarrhea and fever<sup>10</sup>. *Terminalia catappa* has shown biological effects such as having antibacterial and antifungal activities<sup>7,11</sup>, antioxidants<sup>12</sup>, antipyretic, hemostatic, hepatitis<sup>5</sup>, anti-inflammatory, antidiabetic, antioxidant, hepatoprotective, and anticancer<sup>13,14</sup>, antiprotozoal, antiviral, anti-diarrhea, analgesic, antimalarial, and anticancer activities<sup>15</sup>.

*Terminalia catappa* leaves are known to contain chemical compounds such as tannins and flavonoids, which are thought to have antibacterial properties such as *Aeromonas hydrophila*, *Escherichia coli*, and *Staphylococcus aureus*<sup>11</sup>. *Terminalia catappa* also contains flavonoids, alkaloids, tannins, triterpenoids, steroids, resins, saponins, quinones, and phenolics<sup>16-18</sup>. According to several previous studies, giving T. *catappa* leaves extract has been shown to inhibit of several bacteria such as *Aeromonas salmonicida*, *A. hydrophila*, *E. coli*<sup>19</sup>, *S. aureus*, *Pseudomonas aeruginosa*<sup>5</sup>, and *Bacillus amyloliquefaciens*<sup>11</sup>.

*Terminalia catappa* leaves extract also has antifungal activity against *Candida sp*<sup>10</sup>. In

addition, *T. catappa* leaves extract can be used to increase the resistance of betta fish and tilapia to *A. hydrophila*<sup>20</sup>. In this study, the *T. catappa* leaves used were fallen leaves. Previous studies have shown that antibacterial and antifungal activity is higher in fallen *T. catappa* leaves than leaves still on trees<sup>21</sup>. Then the leaves are extracted with ethanol solvent and formulated into liquid soap. Subsequently, physical evaluation and antibacterial activity were carried out.

This study expects that the liquid soap products produced have good physical characteristics and antibacterial activity. **MATERIALS AND METHODS** **Materials** The materials used were *T. catappa* leaves, 96% ethanol, SLS, Comperland, CAB 30, NaCl, citric acid, glycerol, nipagin, Na<sub>4</sub> EDTA, distilled water, blank disc, strains of *S. aureus*, *S. epidermidis*, and *E. coli*, antibacterial body wash soap, and Mueller-Hinton agar (MHA) media. The tools used were rotary evaporator, analytical scale, pH meter, incubator, autoclave, oven, caliper, and laminar airflow.

**Methods** **Sample collection** *Terminalia catappa* leaves were collected in Pekanbaru city and determined at the Botanical Laboratory, Universitas Riau. Samples taken were *T. catappa* leaves that had fallen around the trees with brownish leaves characteristics. The leaves of *T. catappa* were shown in Figure 1. / Figure 1. *Terminalia catappa* leaves **Preparation of simplicia** The collected leaves were washed under running water to remove dirt on the leaves. The leaves were then chopped into small pieces to expand the surface and speed up the drying process. The samples were dried at room temperature until dry; then, they were sorting and ground into a powder.

The powders that had been produced were macerated with 96% ethanol for three days. Afterward, they were filtered and separated between the filtrate and the residue. Maceration was repeated three times with the same type and amount of solvent; then, the macerate was collected and evaporated using a rotary evaporator. **Phytochemical screening** **Alkaloids test:** A total of 0.5 mL of the sample was inserted into three test tubes. Each tube was then added by a few drops of Wagner's, Mayer's, and Dragendorff's reagents. **Flavonoids test:** A total of 0.5

mL of sample was heated for five minutes, then it was added with three drops of HCl concentrated and a little Mg powder. **Saponins test:** A total of 1 mL of sample was added with 2 mL of hot water, shaken, and let stand for five minutes. **Tannins test:** A total of 0.5 mL of sample was added with three drops of 1% FeCl<sub>3</sub>. **Terpenoids test:** A total of 0.5 ml of sample was added with three drops of Liebermann-Burchard's reagent through the test tube wall, and the results were observed<sup>22,23</sup>. **Liquid soap formulation** As much as 40 g of Comperland was mixed with 30 g of CAB 30 and shook until it was thick. Then, 180 g of SLS was added along with 100 mL of water; then, it was stirred until

blended.

A total of 1 g of Na<sub>4</sub> EDTA, 1 g of nipagin, and 400 mL of water were added and stirred until it was homogeneous. After it, 20 g of glycerin, 6 g of NaCl, 2 g of citric acid, and the remaining water were added and stirred until it was homogeneous. The mixture was stored in a tightly-closed container and waited until the shower gel's foam was gone. Furthermore, T. catappa leaves extract was added with a concentration of 1%, 2%, and 3%, stirred until it was homogeneous, and stored in a tightly closed container.

Physical evaluation of preparations Organoleptic test: An organoleptic test was carried out by observing the physical form of liquid soap preparations using the senses. Liquid soap preparations that have been formulated were observed in terms of color, odor, as well as dosage form<sup>24</sup>. pH test: Measurement of pH values was carried out using a pH meter in a 10% sample solution, which was made by dissolving 1 g of the sample in 9 mL of water.

Measurements were made by immersing the pH meter electrode, rinsed with distilled water into the solution. The pH value was determined after the numbers read on the pH meter have stabilized<sup>25</sup>. Foam stability test: The sample was weighed as much as 1 g, put into a test tube, then added with up to 10 mL of distilled water, shaken by turning the test tube back and forth, and immediately measured the level of foam produced. Then, the tube was left to stand for five minutes, then the height of the resulting foam was measured again after five minutes<sup>26</sup>.

???????? ?????????????????????= ?????????? ?????????? h????????h?? ?????????????????? ?????????? h????????h?? ?? 100% Skin irritation test: Testing was done using an open patch test. The open patch test was performed by applying the preparation to the inner forearm. A total of 1 cm of the preparation was applied and observed for 30-minute intervals for skin irritation, erythema, and redness. It was tested on the four formulations that have been made. Homogeneity test: A total of 1 g of liquid soap preparation was smeared on the surface of the mica plastic, then the coarse particles were observed by being touched, and the texture of the preparation was observed.

Antibacterial activity test All test bacteria (S. aureus, S. epidermidis, and E. coli) were respectively inoculated on MHA media. The 6 mm blank disc was dipped in liquid bath soap of T. catappa leaves extract and placed on the media's surface. The same thing was done with liquid bath soap sold in the market as a positive control and liquid soap base as a negative control. All samples were incubated at 35±2°C for 24 hours, and the formed inhibition zone was observed<sup>27</sup>. The interpretation was performed by looking at the clear area around the disc, indicating no bacterial growth. Then, the diameter of the

clear zone formed was measured by using a caliper.

**RESULTS AND DISCUSSION** Phytochemical screening **Phytochemical screening was carried out to determine secondary metabolite compounds contained in** a plant. Secondary metabolite compounds in one type of plant could vary, influenced by climate, soil, temperature, humidity, and others<sup>28</sup>. Therefore, this **phytochemical screening was carried out to determine the** secondary metabolite content of *T. catappa* leaves growing in the Pekanbaru area. Based on the test results, the compounds in the ethanol extract of *T. catappa* leaves were flavonoids, triterpenoids, saponins, and tannins, as shown in Table I. Table I. Phytochemical screening of *T.*

*catappa* leaves extract

No	Phytochemical test	Result	Conclusion
1	Alkaloid - Mayer's - Wagner's - Dragendorff's	white color - yellow color - brown color +	2
2	Flavonoid	orange color +	3
3	Triterpenoid/	purple color +	4
4	Steroid	blue color -	5
5	Tannin	green color +	5
5	Saponin	stable froth or foam is formed +	

Physical evaluation of preparations The liquid soap preparation formulation was made with four formulas, where F0 was a liquid soap base, F1 = base + 1% *T. catappa* leaves extract, F2 = base + 2% *T. catappa* leaves extract, and F3 = base + 3% *T. catappa* leaves extract. Physical evaluation of liquid soap preparations on the organoleptic test includes color, odor, and shape.

Based on the color of the preparations formed on the base, F0 was clear, while F1 was brown, and F2 and F3 were dark browns. This was because the *T. catappa* leaves extract was brown. While the preparation's odor had a distinctive aroma of extracts, F0, F1, and F2 were thick, while F3 was slightly liquid. The more the addition of the extract causes the soap to become more liquid. The resulting soap preparations could be seen in Figure 2. Furthermore, evaluation of the preparations was carried out, including organoleptic tests, pH, foam height, irritation, and homogeneity, as shown in Table II.

The pH formed in liquid soap preparations ranges from 4.6 to 5.2. The SNI 4085:201729 stipulates that the pH quality requirements for liquid soap range from 4 to 10 so that all formulas produced have a pH value that meets the requirements as liquid soap. Furthermore, in the homogeneity test, the preparation was mixed homogeneously, and **there were no coarse grains** on the preparation. Foam formation was not required and **had little effect on the** cleaning process, but it was more likely to patient acceptance of the product.

The criteria for good foam stability, which was within 5 minutes, the foam stability obtained ranges from 60 - 70%. In this case, F0, F1, F2 had met the criteria for good foam stability, which was in the range of 67-70%, except for F3. In the irritation test, the

preparations were given to ten panelists who did not have a history of allergies. The preparation was applied to the skin of the forearm and left for 30 minutes. The test results showed no signs of skin irritation in the ten participants, such as dry skin, pain, bleeding, and cracked skin. Thus, the preparation was declared not to irritate the skin. / Figure 2. Liquid soap from T. catappa leaves Table II.

Physical evaluation of liquid soap from T. catappa leaves extract Parameters \_F0 \_F1 \_F2 \_F3 \_\_ Color \_Clear \_Brown \_Dark brown \_Dark brown \_\_ Odor \_Odorless \_Distinctive of extracts \_Distinctive of extracts \_Distinctive of extracts \_\_ Shape \_Thick \_Thick \_Thick \_Slightly liquid \_\_ pH \_5.2 \_5 \_4.8 \_4.6 \_\_ Foam stability \_70% \_67% \_70% \_72% \_\_ Irritation \_Nothing happened \_Nothing happened \_Nothing happened \_Nothing happened \_\_ Antibacterial activity test The antibacterial activity test of T.

catappa leaves extracts liquid soap was carried out on three formulas, two controls with three times replication against the bacteria *S. aureus*, *S. epidermidis*, and *E. coli*. The test results showed that the preparation could inhibit bacterial growth. The higher the addition of T. catappa leaves extract concentration, the larger the inhibition zone's diameter, as presented in Table III. The antibacterial activity test of liquid soap preparations showed a difference in the inhibition zone diameter. The higher the concentration of T. catappa leaves extract was added, the higher the formed inhibition zone's diameter. Thus, the higher the ability of an extract to inhibit bacterial growth.

This finding was due to the presence of flavonoid compounds in the extract, which could be antibacterial<sup>30</sup>. This literature also supported by the results of phytochemical screening of T. catappa leaves extracts that were positive for flavonoid compounds. The inhibition zone formed at each concentration of 1%; 2%; 3%; positive; and negative control on *S. aureus* bacteria was 25.1; 28.13; 30.07; 40. 67; and 6.1 mm, respectively. The inhibition zone formed on *S. epidermidis* bacteria at each concentration was 12.17; 15.13; 19.17; 25.1; and 6.17 mm. Meanwhile, the inhibition zone against *E. coli* was 6.6; 7.17; 8; 15; and 6.2 mm.

The variation of inhibition zone diameter and SD values (>1 mm) yielded from three replications, which could be seen in Table III, was possibly generated by several factors such as the incubation temperature, diffusion ability of the extract, and volume of the medium used. The optimal temperature for bacterial growth is 35°C; hence, the lower temperature used could produce variation in inhibition zone diameter. If three or more discs were arranged in one pile in the experimental study, the middle disc would experience incubation temperature below 35°C<sup>31</sup>. Table III. Antibacterial activity of liquid soap from T.

catappa leaves extract Sample \_Inhibition zone diameter±SD (mm) \_ \_S. aureus \_S. epidermidis \_E. coli \_Control (+) \_40.67±1.15 \_25.1±1.31 \_15±0.85 \_F0 \_6.1±0.17 \_6.17±0.29 \_6.2±0.2 \_F1 \_25.1±0.96 \_12.17±0.87 \_6.6±0.53 \_F2 \_28.13±0.61 \_15.13±1.27 \_7.17±0.31 \_F3 \_30.07±1.01 \_19.17±0.76 \_8±1.25 \_ \_ Moreover, variation in inhibition zone diameter of bacterial growth also could be produced from the inconsistency of medium thickness used. The most effective medium thickness for bacterial inhibition study was approximately 4 mm thick, which thinner medium could quicken the extract solution diffusion while a thicker medium could slow it down<sup>32</sup>.

Unfortunately, in this research, the medium thickness used was not measured; hence, it was pretty hard to make sure the medium thickness. Based on the formed inhibition zone, it could be grouped into four groups; which were very strong (the inhibition zone >20 mm), strong (10-20 mm), moderate (5-10 mm), and weak (<5 mm)<sup>33</sup>. Therefore, liquid soap with a T. catappa leaves extract concentration of 1%, 2%, 3% in the preparation provides a very strong inhibitory power against S. aureus, a strong inhibitory power against S. epidermidis, and moderate inhibition against E. coli.

This result follows the literature and research objectives that the types of bacteria that could significantly infect the skin were S. aureus and S. epidermidis<sup>34</sup>. CONCLUSION The physical characteristics of T. catappa leaves extract liquid soap meets the requirement of SNI 4085:2017 with a pH value that was safe for the skin. The addition of T. catappa leaves extract variations did not affect the pH value, foam stability, and irritation. However, in the organoleptic test, the higher the concentration of T. catappa leaves extract was added, the liquid soap's color was getting more brownish, and the shape form was slightly liquid. The addition of T.

catappa leaves extract to liquid soap could increase the antibacterial activity. The highest antibacterial activity was shown by S. aureus with an inhibition zone diameter of more than 20 mm. ACKNOWLEDGMENT The authors thank Universitas Abdurrab Foundation for providing research grants. Furthermore, thanks to the Medicine and Health Sciences Faculty, Department of Pharmaceutical and Food Analysis, who have provided facilities for implementing this research.

Furthermore, thanks to the students who participated in helping carry out the research.

REFERENCES Mukhopadhyay P. Cleansers and Their Role in Various Dermatological Disorders. Indian J Dermatol. 2011;56(1):2-6. doi:10.4103/0019-5154.77542 Burton M, Cobb E, Donarchie P, Judah G, Curtis V, Schmidt WP. The Effect of Handwashing with Water or Soap on Bacterial Contamination of Hands. Int J Environ Res Public Health. 2011;8(1):97-104. doi:10.3390/ijerph8010097 Jing JIJ, Yi TP, Bose RJC, McCarthy JR, Tharmalingam N, Madheswaran T. Hand Sanitizers: A Review on Formulation Aspects,



Adverse Effects, and Regulations. *Int J Environ Res Public Health*. 2020;17(9):3326. doi:10.3390/ijerph17093326 Weatherly LM, Gosse JA. Triclosan Exposure, Transformation, and Human Health Effects. *J Toxicol Environ Health B Crit Rev*. 2017;20(8):447-69. doi:10.1080/10937404.2017.1399306 Allyn OQ, Kusumawati E, Nugroho RA.

Antimicrobial activity of Terminalia catappa brown leaf extracts against Staphylococcus aureus ATCC 25923 and Pseudomonas aeruginosa ATCC 27853. *F1000Res*. 2018;7:1406. doi:10.12688/f1000research.15998.1 Anand AV, Divya N, Kotti PP. An updated review of Terminalia catappa. *Pharmacogn Rev*. 2015;9(18):93-8. doi:10.4103/0973-7847.162103 Silva LP, de Angelis CD, Bonamin F, Kushima H, Mininel FJ, Dos Santos LC, et al. Terminalia catappa L.: a medicinal plant from the Caribbean pharmacopeia with anti-Helicobacter pylori and antiulcer action in experimental rodent models. *J Ethnopharmacol*. 2015;159:285-95. doi:10.1016/j.jep.2014.11.025 Nair R, Chanda S.

Antimicrobial Activity of Terminalia catappa, Manilkara zapota and Piper betel Leaf Extract. *Indian J Pharm Sci*. 2008;70(3):390-3. doi:10.4103/0250-474X.43012 Ng S, Lasekan O, Muhammad KS, Hussain N, Sulaiman R. Physicochemical properties of Malaysian-grown tropical almond nuts (Terminalia catappa). *J Food Sci Technol*. 2015;52(10):6623-30. doi:10.1007/s13197-015-1737-z Terças AG, Monteiro ADZ, Moffa EB, Dos Santos JRA, de Sousa EM, Pinto ARB, et al. Phytochemical Characterization of Terminalia catappa Linn. Extracts and Their antifungal Activities against Candida spp. *Front Microbiol*. 2017;8:595. doi:10.3389/fmicb.2017.00595 Tampemawa PV, Pelealu JJ, Kandou FEF.

Uji Efektivitas Ekstrak Daun Ketapang (Terminalia catappa L.) Terhadap Bakteri Bacillus amyloliquefaciens. *Pharmacon*. 2016;5(1):308-20. doi:10.35799/pha.5.2016.11324 Widyaningsih S, Chasani M, Diastuti H, Fredyono WN. Liquid Soap from Nyamplung Seed Oil (Calophyllum inophyllum L) with Ketapang (Terminalia catappa L) as Antioxidant and Cardamom (Amomum compactum) as Fragrance. *Molekul*. 2018;13(2):172-9. doi:10.20884/1.jm.2018.13.2.461 Divya N, Rengajaran RL, Radhakrishnan R, Allah EFA, Alqarawi AA, Hashem A, et al. Phytotherapeutic efficacy of the medicinal plant Terminalia catappa L. *Saudi J Biol Sci*. 2019;26(5):985-8. doi:10.1016/j.sjbs.2018.12.010 Chanda S, Rakholiya K, Nair R.

Antimicrobial Activity of Terminalia catappa L. Leaf Extracts against Some Clinically Important Pathogenic Microbial Strains. *Chin Med*. 2011;2(4):171-7. doi:10.4236/cm.2011.24027 Cock IE, Van Vuuren SF. A comparison of the antimicrobial activity and toxicity of six Combretum and two Terminalia species from southern Africa. *Pharmacogn Mag*. 2015;11(41):208-18. doi:10.4103/0973-1296.149740 Riskitavani DV,



Purwani KI. Studi Potensi Bioherbisida Ekstrak Daun Ketapang (*Terminalia catappa*) terhadap Gulma Rumpuk Teki (*Cyperus rotundus*). *Jurnal Sains dan Seni ITS*. 2013;2(2):E59-63. doi:10.12962/j23373520.v2i2.3593 Istarina D, Khotimah S, Turnip M.

Aktivitas Antibakteri Ekstrak Metanol Buah Ketapang (*Terminalia catappa* Linn.) Terhadap Pertumbuhan *Staphylococcus epidermidis* Dan *Salmonella typhi*. *Protobiont Jurnal Elektronik Biologi*. 2015;4(3):98-102. doi:10.26418/protobiont.v4i3.13321 Katiki LM, Gomes ACP, Barbieri AME, Pacheco PA, Rodrigues L, Verissimo CJ, et al. *Terminalia catappa*: Chemical composition, in vitro and in vivo effects on *Haemonchus contortus*. *Ver Parasitol*. 2017;246:118-23. doi:10.1016/j.vetpar.2017.09.006 Sumino, Supriyadi A, Wardiyanto. The Effectiveness of Ketapang (*Terminalia catappa* L.) Leave Extract for the Treatment of *Aeromonas salmonicida* Infection in Catfish (*Pangasioniodon hypophthalmus*). *Jurnal Sain Veteriner*. 2013;13(1):79-88. doi:10.22146/jsv.3503 Nugroho RA, Manurung H, Nur FM, Prahastika W. *Terminalia catappa* L. extract improves survival, hematological profile and resistance to *Aeromonas hydrophila* in *Betta* sp. *Arch Polish Fish*. 2017;25(2):103-15. doi:10.1515/aopf-2017-0010 Hardhiko RS, Suganda AG, Sukandar EY. Aktivitas antimikroba ekstrak etanol, ekstrak air daun yang dipetik dan daun gugur pohon ketapang (*Terminalia catappa* L.). *Acta Pharm Indones*. 2004;29:129-33.

Haro G, Iksen I, Rumanti RM, Marbun N, Sari RP, Gultom RPJ. Evaluation of Antioxidant Activity and Minerals Value from Watercress (*Nasturtium officinale* R.Br.). *Rasayan J Chem*. 2018;11(1):232-7. doi:10.7324/RJC.2018.1112011 Syahrina S, Asfianti V, Gurning K, Iksen I. Phytochemical Screening and Anti-Hyperuricemia Activity Test In Vivo of Ethanolic Extract of Shallot (*Allium cepa* L.) Skin. *Borneo J Pharm*. 2020;3(3):146-51. doi:10.33084/bjop.v3i3.1365 Muthmainnah R, Rubiyanto D, Julianto TS. Formulasi Sabun Cair Berbahan Aktif Minyak Kemangi Sebagai Antibakteri Dan Pengujian Terhadap *Staphylococcus Aureus*. *IJCR Indones J Chem Res*. 2014;1(2):44-50. doi:10.20885/ijcr.vol1.iss2.art6 Sasongko H, Mumpuni AS. Pengaruh penambahan sukrosa terhadap mutu sabun transparan dari ekstrak etanol herba pegagan (*Centella asiatica* L.). *Pharmaciana*. 2017;7(1):71-8. doi:10.12928/pharmaciana.v7i1.5795 Anggraeni Y, Nisa' F, Bertha OS.

Karakteristik Fisik dan Aktivitas Antibakteri Sabun Cair Minyak Nilam (*Pogostemon cablin* Benth.) yang Berbasis Surfaktan Sodium Lauril Eter Sulfat. *Jurnal Kefarmasian Indonesia*. 2020;10(1):1-10. doi:10.22435/jki.v10i1.499 Abu FA, Yusriadi, Tandah MR. Formulasi Sediaan Sabun Cair Antibakteri Minyak Atsiri Daun Kemangi (*Ocimum americanum* L.) dan Uji Terhadap Bakteri *Staphylococcus epidermidis* dan *Staphylococcus aureus*. *Jurnal Farmasi Galenika Galenika J Pharm*. 2015;1(1):1-8. doi:10.22487/j24428744.2015.v1.i1.4835 Sampaio BL, Edrada-Eel R, Da Costa FB. Effect of

the environment on the secondary metabolic profile of *Tithonia diversifolia*: a model for environmental metabolomics of plants. *Sci Rep*. 2016;6:29265. doi:10.1038/srep29265  
Badan Standarisasi Nasional.

Sediaan Sabun Cair. SNI 4085:2017. Jakarta, Indonesia: Badan Standarisasi Nasional; 2017. Saputra SA, Lailiyah M, Erivina A. Formulasi Dan Uji Aktivitas Anti Bakteri Masker Gel Peel-Off Ekstrak Daun Pacar Air (*Impatiens balsamina* linn.) Dengan Kombinasi Basis PVA dan HPMC. *Jurnal Riset Kefarmasian Indonesia*. 2019;1(2):114-22.

doi:10.33759/jrki.v1i2.20 Balouiri M, Sadiki M, Ibsouda SK. Methods for in vitro evaluating antimicrobial activity: A review. *J Pharm Anal*. 2016;6(2):71-9.

doi:10.1016/j.jpha.2015.11.005 Zeniusa P, Ramadhan MR, Nasution SH, Karima N. Uji Daya Hambat Ekstrak Etanol Teh Hijau Terhadap *Escherichia Coli* Secara In Vitro. *Med J Lampung Univ Majority*. 2019;8(2):136-43. Davis WW, Stout TR.

Disc Plate Method of Microbiological Antibiotic Assay: I. Factors Influencing Variability and Error. *Appl Microbiol*. 1971,22(4):659-65. Otto M. *Staphylococcus epidermidis* – the “accidental” pathogen. *Nat Rev Microbiol*. 2009;7(8):555-67. doi:10.1038/nrmicro2182

#### INTERNET SOURCES:

-----  
<1% -

<https://skinproductsreviews.com/best-of/23-toxic-cosmetic-ingredients-you-should-avoid/>

<1% - <https://canadianskin.ca/skin-conditions-and-diseases>

<1% - <https://iopscience.iop.org/issue/1755-1315/637/1>

<1% - [http://www.fao.org/tempref/KC/Reserved/Cabi/AgoraExport22\\_01\\_13.xml](http://www.fao.org/tempref/KC/Reserved/Cabi/AgoraExport22_01_13.xml)

<1% - <https://onlinelibrary.wiley.com/doi/full/10.1002/fsn3.770>

<1% -

[https://www.researchgate.net/publication/333619003\\_QUALITATIVE\\_AND\\_QUANTITATIVE\\_ANALYSIS\\_OF\\_THE\\_CONTENT\\_OF\\_CHEMICAL\\_COMPOUNDS\\_FROM\\_EXTRACTS\\_OF\\_HEXANE\\_ACETONE\\_ETHANOL\\_AND\\_WATER\\_FROM\\_AVOCADO\\_LEAVES\\_Persea\\_americana\\_Mill](https://www.researchgate.net/publication/333619003_QUALITATIVE_AND_QUANTITATIVE_ANALYSIS_OF_THE_CONTENT_OF_CHEMICAL_COMPOUNDS_FROM_EXTRACTS_OF_HEXANE_ACETONE_ETHANOL_AND_WATER_FROM_AVOCADO_LEAVES_Persea_americana_Mill)

<1% - <https://www.plantsjournal.com/archives/2018/vol6issue2/PartC/6-2-14-881.pdf>

<1% -

<http://images.pcmac.org/SiSFiles/Schools/AL/MadisonCity/BJHigh/Uploads/Forms/AP%20Chemistry%20Lab%20Manual%20and%20Syllabus%20for%20students%20Elegante.doc>

<1% -

[https://www.researchgate.net/publication/315962801\\_Preliminary\\_phytochemical\\_screening\\_total\\_phenolic\\_content\\_and\\_antibacterial\\_activity\\_of\\_thirteen\\_native\\_species\\_from\\_G](https://www.researchgate.net/publication/315962801_Preliminary_phytochemical_screening_total_phenolic_content_and_antibacterial_activity_of_thirteen_native_species_from_G)

uayas\_province\_Ecuador

<1% -

<https://jurnal.farmasi.umi.ac.id/index.php/fitofarmakaindo/article/download/727/421>

<1% - <https://iopscience.iop.org/article/10.1088/1742-6596/1402/5/055069/pdf>

<1% -

<https://www.dovepress.com/a-novel-ocular-delivery-of-brinzolamide-based-on-gellan-gum-in-vitro-a-peer-reviewed-fulltext-article-DDDT>

<1% -

[https://www.academia.edu/37645526/MICROBIOLOGY\\_PRACTICAL\\_GUIDE\\_A\\_2010\\_2\\_PRACTICAL\\_I](https://www.academia.edu/37645526/MICROBIOLOGY_PRACTICAL_GUIDE_A_2010_2_PRACTICAL_I)

<1% - <https://biologywise.com/what-does-zone-of-inhibition-mean>

<1% - <http://biodiversitas.mipa.uns.ac.id/D/D2110/D211020.pdf>

<1% -

<http://www.orientjchem.org/vol36no5/gel-formulation-of-jamblang-leaf-extract-syzygiumcumini-l-skeel-and-antioxidant-activity/>

<1% - <https://pubchem.ncbi.nlm.nih.gov/compound/p-toluenesulfonamide>

<1% - <https://iopscience.iop.org/issue/1755-1315/462/1>

<1% - <http://europepmc.org/articles/PMC4902980>

<1% - <http://www.fao.org/3/W3732E/w3732e06.htm>

<1% - <https://iopscience.iop.org/issue/1755-1315/347/1>

<1% -

[https://www.researchgate.net/publication/286201407\\_Antibacterial\\_Activity\\_of\\_an\\_Effective\\_Essential\\_Oil\\_Formulated\\_in\\_Liquid\\_Soap\\_Against\\_Skin\\_Bacteria](https://www.researchgate.net/publication/286201407_Antibacterial_Activity_of_an_Effective_Essential_Oil_Formulated_in_Liquid_Soap_Against_Skin_Bacteria)

1% - <https://www.frontiersin.org/articles/10.3389/fpubh.2020.574444/full>

1% -

[https://issuu.com/separ/docs/libro\\_20separ\\_20organizaci\\_c3\\_b3n\\_20en\\_20la\\_20covi](https://issuu.com/separ/docs/libro_20separ_20organizaci_c3_b3n_20en_20la_20covi)

1% - <https://ojs.jmolekul.com/ojs/index.php/jm/article/download/461/293>