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# Therapeutic Effect of Topical Ointment Ethanol Extract From Patiwala Leaves (*Lantana camara*) on Histological Profile of Incision Wounds in Diabetic Rat Models

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## Abstract

Diabetic wounds are one of the neurovascular complications of diabetes. Hyperglycemia complicates the healing of diabetic wounds, which are susceptible to infection and chronic inflammation. Topical treatment using herbal extracts aims to reduce the side effects of surrounding tissue damage. A topical ointment made from ethanol extract of Patiwala (*Lantana camara*) leaves, which has been tested for product quality, has antibacterial and anti-inflammatory effects that have been proven to help accelerate the healing of diabetic wounds in rats through parameters such as reduced wound diameter, changes in the number of fibroblast cells, and collagen fiber formation seen in the histological profile during 21 days of therapy with a concentration of 15% ( $p > 0.05$ ) compared to the positive control of 10% betadine.

**Keywords:** Topical ointment; *Lantana camara*; Histological profile; Diabetic wounds; Fibroblast cells

## INTRODUCTION

Diabetes is a major cause of damage to various organs such as the heart, blood vessels, nerves, eyes, and kidneys (Harreiter & Roden, 2023; Cloete, 2022). One complication of diabetes related to blood vessel and nerve damage is diabetic wounds or diabetic neuropathy (Urso et al., 2021; McDermott et al., 2023). Sensory and motor neuropathy can cause various changes in the skin and muscles, which then lead to changes in pressure distribution in the peripheral area that triggers ulcers (Kurz, 2020; Akkus & Sert, 2022). The healing of diabetic ulcers or diabetic wounds is a serious concern for clinicians today. This is because the speed of diabetic wound healing is not the same as the healing of wounds in general. Diabetic wounds contain high blood glucose levels (hyperglycemia) which is known to be an energy source for pathogenic bacteria, making diabetic wounds very susceptible to infection and inflammation (Raja et al., 2023; Wang et al., 2025).

Topical treatment of diabetic wounds using chemical antibiotic ointments often causes side effects, including damage to the surrounding skin tissue. Therefore, various studies on herbal ointments are ongoing, with the hope that they can become an alternative treatment for diabetic wounds with minimal side effects. Herbal plant extracts intended to help accelerate diabetic wound healing must be able to inhibit pathogenic bacteria that trigger infection and inflammation in diabetic wounds in vitro (Ramachandran et al., 2023; Norman et al., 2021).

One of the local plants often used by the people of Southeast Sulawesi to treat wounds is the Patiwala leaf (*Lantana camara*). Several previous studies have reported the potential of *Lantana camara* in various preparations for incision wounds (Tamuntuan et al., 2021; Arifin et al., 2023; Saranani et al., 2023) but there has been no specific research on diabetic wounds. The ability of *Lantana camara* in treating wounds is influenced by its phytochemical compounds which functions as an antibacterial and anti-inflammatory (Sari et al., 2023; Hasnaeni et al., 2024). The results of phytochemical screening of *Lantana camara* vary quite a bit, depending on the geographical conditions of the place of growth as well as the analytical method used (Kapitan et al., 2024; Orji et al., 2024). The phytochemical compounds of ethanol extract of Patiwala leaves from Southeast Sulawesi has been reported in a study by Rosanty et al. (2025) with active compounds of alkaloids, flavonoids, tannins, saponins and terpenoids (Rosanty et al., 2025). Antibacterial activity test of the extract against several pathogenic bacteria causing diabetic ulcers in vitro including *Staphylococcus aureus*, *Staphylococcus epidermis*, *Proteus mirabilis*, *Proteus vulgaris*, *Pseudomonas aeruginosa*, *Klebsiella pneumoniae* and *E.Coli* with ciprofloxacin as a positive control showed quite significant results. This study is a follow-up study of Rosanty (2025) which was conducted in vivo using a white rat model of the Wistar type with diabetic wounds to observe the histological profile after topical ointment therapy with ethanol extract of Patiwala leaves (*Lantana camara*).

54

## MATERIALS AND METHODS

### 55 Research Ethics and Design

56 This research was an experimental study involving 30 male Wistar rats. The research was conducted from June to  
 57 December 2024 at the Biomedical Research Laboratory, Faculty of Medicine, Halu Oleo University. **This study**  
 58 **received ethical approval from the Health Research Ethics Committee,** Tanjung Karang Ministry of Health  
 59 Polytechnic, under No. 451/KEPK-TJK/VIII/2023.

60

### 61 Materials and equipment

62 The main material used in this study was a topical ointment of ethanol extract of Patiwala (*Lantana camara*) leaves  
 63 formulated based on the results of in vitro extract effectiveness tests in the study by Rosanty (2025) with the following  
 64 formula:

| Materials  | % ingredients in every 10 grams of formula |        |        | Function        |
|--|--|--------|--------|-----------------|
|  | A  | B      | C      |                 |
| Ethanol extract of Patiwala leaves ( <i>Lantana camara</i> ) | 5  | 10     | 15     | Active compound |
| Propil paraben   | 0.01                                       | 0.01   | 0.01   | Preservative    |
| Cera alba  | 2  | 2      | 2      | Base            |
| Vaselin album  | Ad 100                                     | Ad 100 | Ad 100 | Base            |

65

66 Other materials and equipment include measuring flasks, Erlenmeyer flasks, test tubes, stirring rods and dropper  
 67 pipettes for making ointments, weighing rats using a MACS 1.5/W digital scale, scalpels for making incision wounds,  
 68 calipers for measuring wound diameter. Making a diabetic rat models using an induction agent streptozotocin/STZ  
 69 (Sigma-Aldrich) dissolved in 0.9% NaCl, **measuring blood glucose levels** in using a **point of care testing (POCT)**  
 70 FORA 6 Plus (Switzerland). Histological observation of incision wounds using hematoxylin-eosin staining technique  
 71 with natural buffered formalin 10% (Indopath-Paraform) for tissue fixation, ethanol 70%, 80% and 95% (Merck) for  
 72 dehydration-rehydration, xylol (Bio-Analitika) for clearing, paraffin (Paraf flakes) and basemold for embedding,  
 73 microtome tool (Accu-Cut), object glass and deck glass for microtomy and preparation of preparations, hematoxylin  
 74 (Biognost) and eosin (Indoreagen).

75

### 76 Treatment of experimental animals

77 **The experimental animals used in this study were 30 male white Wistar rats with body weights ranging from 90-200**  
 78 **grams.** The rats **were divided into 3 groups with 10 rats each group consisting of** K-7 for the intervention group for 7  
 79 days, K-14 for the intervention group for 14 days, and K-21 for the intervention group for 21 days. Then all rats were  
 80 acclimatized in standard cages, fed standard food with a light/dark cycle for 2 weeks. Body weight (BW) and random  
 81 blood glucose level measurements (RBG) were carried out on all rats before induction. Each group was then divided  
 82 into 5 subgroups consisting of the 5% Patiwala ointment intervention group (K1), the 10% Patiwala ointment  
 83 intervention group (K2), the 15% Patiwala ointment intervention group (K3), the 10% Betadine ointment positive  
 84 control (KP) and the untreated negative control (KN), each subgroup consisting of 2 rats. STZ induction was  
 85 performed on all rats with a single intraperitoneal dose of 50 mg/kgBW. After 5 days, BW and RBG levels were  
 86 remeasured to confirm diabetes (RBG >200 mg/dL). A 5 cm incision wound was made in the low back of the diabetic  
 87 rats. Treatment was administered twice daily to both **the intervention and positive control groups.** **On the final day of**  
 88 **the intervention,** the rats **were** euthanasia using carbon monoxide. Skin fragments from the incision wounds were  
 89 removed and fixed in 10% NBF for histological analysis. Intracardial blood was collected using gel separator tubes,  
 90 other organs and tissues were also collected and stored in 10% NBF for further research.

91

### 92 Measurement of RBG levels

93 RBG levels were measured in each group two times: before STZ induction and 5 days after STZ induction. RBG  
 94 levels were measured using the Fora 6 Plus device using the POCT method, which is based on the principle of  
 95 enzymatic reactions. RBG levels are showed in milligrams per deciliter (mg/dL).

96

### 97 Measurement of wound diameters

98 Wound diameters were measured in each group two times: before STZ induction and 5 days after STZ induction.  
 99 Wound diameter measurements using calipers, the results are showed in millimeters (mm).

100

### 101 Histological analysis

102 The skin tissue that had been fixed in 10% NBF solution for 1x24 hours was then dehydrated using ethanol in stages  
 103 starting from 70%, 80%, and 95% for 2 hours each. Then, the dehydrated tissue was cleared using xylol solution for 2  
 104 cycles, each for 1 hour. Next, embedding was carried out in a basemold using liquid paraffin at a temperature of 40°C.  
 105 After the tissue block was formed, tissue ribbons were cut on a microtome with a size of 5 µm. The tissue ribbons

107 were placed on a glass object, fixed on a hotplate at a temperature of 40°C to remove residual paraffin, and then  
 108 continued with hematoxylin and eosin staining. Histological observations were carried out under a microscope with a  
 109 magnification of 40x (Orno, 2023). The number of fibroblast cells was counted in 15 fields of view on each slide and  
 the results were presented as an average value.

110 **Data analysis**

111 The effects of STZ induction on rats BW and RBG levels were analyzed using the Wilcoxon test. Wound diameters  
 112 and number of fibroblast cells after therapy in each subgroup within a group were analyzed using the One-Way  
 113 ANOVA test and for comparison of wound diameter between groups using the Kruskal Wallis test. The histological  
 114 results are presented in the form of micrographs followed by notes.  
 115

116 **RESULTS AND DISCUSSION**

117 **Characteristics of topical ointment products**

118 The topical ointment made from ethanol extract of patiwala leaves used in this study has been tested in vitro  
 119 to have antibacterial effects against several bacteria that cause diabetic wounds. Below is a description of the topical  
 120 ointment product and the results of the quality test for the ointment preparation.  
 121

122 **Table 1.** Product quality report of Topical ointment of Patiwala leave ethanol extract

| Product  | Product Quality Report   |             |      |                |                    |                  |
|--|--|-------------|------|----------------|--------------------|------------------|
|  | Organoleptic   | Homogeneity | pH   | Viscosity (cP) | Spreadability (mm) | Stickiness (sec) |
| Topical ointment of Patiwala leave ethanol extract | Lightly greenish brown color, distinctive odor, soft hydrocarbon ointment form, smooth without coarse particles, and easy to apply | Homogen     | 6.42 | 8.745          | 41                 | 4                |



123 Ointment physical properties testing is conducted as part of efforts to ensure the quality and stability of the  
 124 preparation and to ensure that the final product meets predetermined quality criteria. This evaluation is important  
 125 because the physical characteristics of the ointment, such as visual appearance, texture, pH, and ability to spread and  
 126 adhere to the skin surface, significantly influence the comfort of use and therapeutic effectiveness of topical  
 127 preparations (Savitri et al., 2025; Wahyuni et al., 2025). The test results showed that the ointment could be applied to  
 128 experimental rats and can be used as a reference for future formulation needs.  
 129

130 **Effect of STZ induction on Body weight and Random blood glucose**

131 This study used STZ as an induction agent in an animal model of diabetes mellitus. Table 2 explains the  
 132 effect of STZ induction on increasing RBG levels with a mean level of 310.23 ± 59.12 (p<0.001) which is in line with  
 133 the increase in BW of rats (p=0.013).  
 134

135 **Table 2.** Effect of STZ induction on Body weight and Random blood glucose

| Variables                    |          | Mean ± SD      | p value |
|------------------------------|----------|----------------|---------|
| Body weight (g)              | Pre-STZ  | 135.75 ± 49.39 | 0.013   |
|                              | Post-STZ | 144.22 ± 41.12 |         |
| Random blood glucose (mg/dL) | Pre-STZ  | 86.13 ± 17.74  | <0.001* |
|                              | Post-STZ | 310.23 ± 59.12 |         |

136 \*Wilcoxon test, the level of significance at p< 0.05

137 Several studies have reported STZ is cytotoxic to pancreatic β-cells, and its effects are visible 72 hours after  
 138 administration and are dose-dependent. The toxic effects of STZ begin with the uptake of STZ into cells via the low-  
 139 affinity glucose transporter-2 (GLUT2) found in the plasma membrane of β-cells, hepatocytes, and renal tubular cells.  
 140 This has been demonstrated by studies showing that insulin-producing cells that do not express GLUT2 are resistant to  
 141 STZ induction (Al-Awar et al., 2016; Pandey & Dvorakova, 2020).  
 142  
 143

## 144 The therapeutic effect of ointment on the diameter of rat wounds

145 Hyperglycemic conditions in diabetes reduce the healing rate of incision wounds in rats this is caused by  
 146 several conditions such as impaired angiogenesis, neuropathy, chronic inflammatory responses and bacterial  
 147 infections (Burgess et al., 2021). Changes in the diameter of the incision wound are a visual indicator of the healing  
 148 effect of the ointment used, as presented in table 3.

149  
 150 **Table 3.** Comparison of wound diameter between groups and subgroups

| Groups/<br>subgroups | K-7 (mm)          | K-14 (mm)         | K-21 (mm)     | p value |
|----------------------|-------------------|-------------------|---------------|---------|
|                      | Mean ± SD         | Mean ± SD         | Mean ± SD     |         |
| KP (betadine 10%)    | 13.23 ± 0.01      | 1.23 ± 0.07       | 0.000 ± 0.00  | 0.095** |
| K1 (patiwala 5%)     | 18.90 ± 0.05      | 13.43 ± 0.07      | 4.85 ± 0.86   | 0.102** |
| K2 (patiwala 10%)    | 16.13 ± 0.15      | 9.54 ± 0.42       | 1.57 ± 0.07   | 0.102** |
| K3 (patiwala 15%)    | 15.91 ± 0.09      | 5.63 ± 0.07       | 0.00 ± 0.00   | 0.095** |
| <b>p value</b>       | <b>&lt;0.001*</b> | <b>&lt;0.001*</b> | <b>0.001*</b> |         |

\*One-way Anova test, \*\*Kruskal Wallis test, the level of significance at  $p < 0.05$

152 The effectiveness of the ointment was estimated against 10% betadine as a positive control. The difference in  
 153 wound diameter between subgroups in each group using the One-way ANOVA test showed a significant difference  
 154 ( $p < 0.001$ ). In the 7-day therapy group (K7), the average KP wound diameter was  $13.23 \pm 0.01$  mm, significantly  
 155 different from the K1, K2, and K3 groups. The 14-day therapy group (K14) showed almost perfect wound closure in  
 156 the KP ( $1.23 \pm 0.07$  mm), while the K3 therapy group showed quite good results with an average wound diameter of  
 157  $5.63 \pm 0.07$  mm. A linear pattern was seen in the 21-day therapy group (K21) where the KP wound diameter was  
 158  $0.000 \pm 0.00$  mm in line with K3, followed by K2 and K1. The results of the Kruskal-Wallis test between groups  
 159 showed significant results ( $p > 0.05$ ) which proved that in all groups there was a significant change in wound diameter.  
 160 Changes in wound diameter indicate the activity of the active compounds contained in the ointment. Several previous  
 161 studies have examined the role of active ingredients in herbal extracts in in vivo diabetic wound healing (Nurwahita et  
 162 al., 2024; Palupi et al., 2022; Wulandari et al., 2023) explains the role of active compounds such as alkaloids,  
 163 flavonoids, tannins, saponins and terpenoids in accelerating the wound tissue regeneration process which is visible in  
 164 changes in wound diameter. Physiological wound tissue regeneration involves muscle fiber repair, extracellular matrix  
 165 remodeling, and collagen deposition. Theoretically, physiological wound closure occurs on days 10-14, but under  
 166 hyperglycemic conditions, the remodeling process can take longer to compensate for the inflammation (Giha et al.,  
 167 2022; Sun et al., 2024). Patiwala topical ointment, with its antibacterial and anti-inflammatory properties, helps  
 168 prevent infection and chronic inflammation during the remodeling process.

## 169 The effect of ointment therapy on the number of fibroblast cells in rats

170 Unlike changes in wound diameter, which can be observed and measured visually, the number of fibroblast  
 171 cells can only be observed and counted microscopically. The number of fibroblast cells is one indicator of successful  
 172 wound healing, calculated in 15 large fields of view (HPF), as presented in table 4.

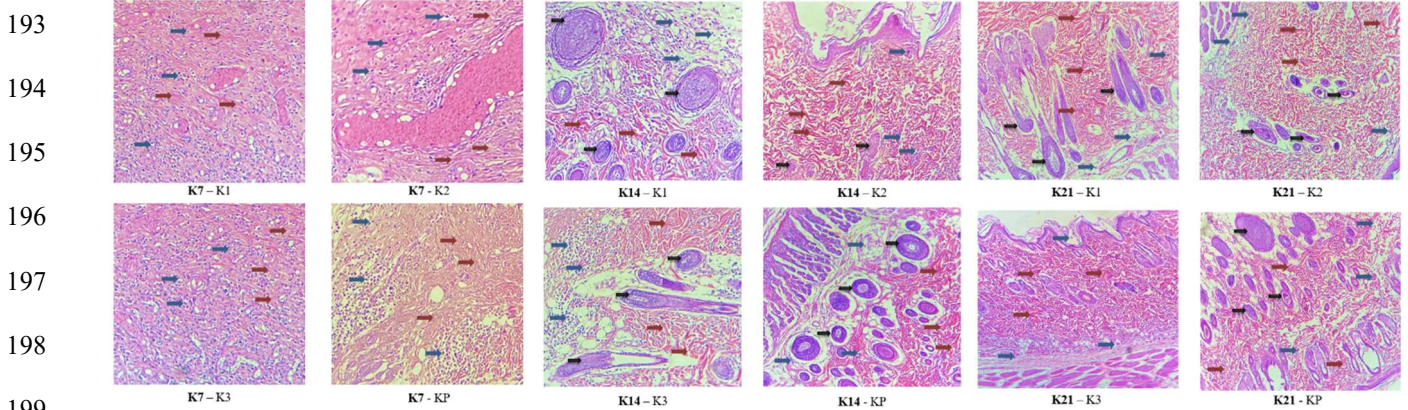
173  
 174  
 175 **Table 4.** Comparison of fibroblast cells between groups and subgroups

| Groups/<br>subgroups | K-7 (cells/HPF) | K-14 (cells/HPF) | K-21 (cells/HPF) | p value |
|----------------------|-----------------|------------------|------------------|---------|
|                      | Mean ± SD       | Mean ± SD        | Mean ± SD        |         |
| KP (betadine 10%)    | 178.50 ± 3.53   | 95.00 ± 4.24     | 108.00 ± 1.41    | 0.102** |
| K1 (patiwala 5%)     | 158.00 ± 4.24   | 110.00 ± 11.31   | 110.00 ± 9.89    | 0.180** |
| K2 (patiwala 10%)    | 167.50 ± 14.84  | 88.50 ± 4.94     | 75.00 ± 12.72    | 0.102** |
| K3 (patiwala 15%)    | 175.50 ± 12.02  | 156.50 ± 62.93   | 90.50 ± 16.42    | 0.180** |
| <b>p value</b>       | <b>0.303*</b>   | <b>0.283*</b>    | <b>0.036*</b>    |         |

\*One-way Anova test, \*\*Kruskal Wallis test, the level of significance at  $p < 0.05$

176 Table 4 shows that there was no significant difference in the number of fibroblast cells in K7 and K14  
 177 ( $p = 0.303$  &  $p = 0.283$ ). On the 7th day of therapy, all groups experienced a significant increase in the number of  
 178 fibroblast cells, K3 was the group with the highest number of fibroblast cells ( $175.50 \pm 12.02$ /HPF) approaching the  
 179 number of KP fibroblast cells ( $178.50 \pm 3.53$ /HPF). The pattern of changes in the number of fibroblast cells began to  
 180 appear non-linear in K14 where KP showed a decrease in the number of fibroblast cells ( $95.00 \pm 4.24$ /HPF) which  
 181 was not in line with the decrease in the number of fibroblast cells in K3 and K1 which on average experienced a  
 182 decrease but anomalous with K2. Although statistically there was no difference in the K14 group, K2 appeared to  
 183 experience a very sharp decrease in the number of fibroblast cells ( $88.50 \pm 4.94$ /HPF). Variations in fibroblast cell  
 184 count patterns during wound healing are theoretically possible (Liu et al., 2021). In the initial phase of wound healing,  
 185 fibroblasts are produced in large numbers to compensate for the inflammation (Guillamat-Prats, 2021; Liu et al.,  
 186 2022). In the subsequent phase (10-14 days later), the number of fibroblasts decreases, and they are replaced by  
 187 collagen fiber formation. In the 21-day therapy group (K21), there was a significant difference between treatment  
 188 subgroups ( $p = 0.036$ ), this may be due to wound re-epithelialization and granulation tissue formation during the

191 experimental period differing between animals. The distribution pattern of fibroblast cells and collagen fiber  
 192 formation can be observed more clearly in the histological profile below.



199  
 200  
 201 **Figure 1.** Changes in the number of fibroblast cells in mice after therapy with ethanol extract ointment of patiwalá leaves at. K7 (7-day therapy  
 202 group), K14 (14-day therapy group), K21 (21-day therapy group). K1 (5% patiwalá ointment subgroup), K2 (10% patiwalá ointment subgroup), K3 (15%  
 203 patiwalá ointment subgroup), KP (10% betadine positive control). Fibroblast cells ( → ) collagen fibers ( → ) dan hair follicles ( → )  
 204

205 Proliferating fibroblasts accompany these vessels and begin to deposit collagen. During the proliferation  
 206 phase, a special type of tissue that characterizes healing, called granulation tissue, appears. The term granulation tissue  
 207 originates from its histological appearance, characterized by the proliferation of fibroblasts, smooth, thin-walled  
 208 capillaries within a loose extracellular matrix. Granulation tissue then progressively accumulates a connective tissue  
 209 matrix, ultimately producing dense fibrosis, which can undergo further remodeling over time (Kunkemoeller &  
 210 Kyriakides, 2017). After injury, exposure of fibrillar collagen to the blood causes platelet aggregation and activation  
 211 and releases chemotactic factors that initiate the wound healing process. Collagen fragments release leukocytic  
 212 collagenase to attract fibroblasts to the wound area. Collagen then forms the foundation for a new extracellular matrix,  
 213 thus accelerating the formation of granulation tissue. The saponin content in the ethanol extract ointment of patiwalá  
 214 leaves prevents wound infection (Chanu et al., 2023; Kunkemoeller & Kyriakides, 2017). According to Chanu (2023),  
 215 the more connective tissue there is in a wound, the greater the contractile force of the wound so that the sides of the  
 216 wound will be pulled and cause the wound to shrink. Fibroblast proliferation in the wound healing process is naturally  
 217 stimulated by interleukin-1b (IL-1b), platelet-derived growth factor (PDGF), and fibroblast growth factor (FGF).  
 218 Fibroblast migration in the injured area is stimulated by transforming growth factor (TGF), a growth factor produced  
 219 by granulation tissue formed during the inflammatory process (Gökşen et al., 2017). The wound healing process is  
 220 greatly influenced by the role of fibroblast migration and proliferation in the injured area (Gökşen et al., 2017; Parmar  
 221 et al., 2018). The content of the ethanol extract ointment of patiwalá leaves applied to the wounds of test animals  
 222 stimulates the synthesis of growth factors including FGF thereby increasing the activity of fibroblast cells to produce  
 223 collagen and form connective tissue so that the wound heals quickly.

224 Ethanol extract ointment from patiwalá leaves contains many phytochemical compounds such as flavonoids,  
 225 tannins, phenols, terpenoids, saponins, and other nutrients that significantly affect health, including healing cuts in test  
 226 animals. Research conducted by Saranani (2023) and Arifin (2023) found that flavonoids and tannins are among the  
 227 components of *Lantana camara* that influence wound healing, particularly wound moisture. Flavonoids can stop  
 228 bleeding in wounds and act as anti-inflammatories, influencing the production of inflammatory cells during the wound  
 229 healing phase. The presence of flavonoids in the cream can help change the condition of wet wounds to become  
 230 moister more quickly. The flavonoid content of *Lantana camara* is believed to play a significant role in the wound  
 231 healing process. In addition to flavonoids, tannins act as astringents, reducing mucosal permeability and strengthening  
 232 inter-mucosal bonds, thus preventing irritation. Therefore, tannins indirectly influence changes in moisture levels. In  
 233 addition to affecting mucosal permeability, tannins can also affect the permeability of bacterial walls or membranes,  
 234 causing bacteria (Zubair & Ahmad, 2019). These antibacterial properties can prevent wound infections. The saponins  
 235 contained in *Lantana camara* can influence collagen production in the early stages of tissue repair and stimulate  
 236 epithelial cell regeneration in the skin, thereby accelerating the wound healing process in test animals. Phenolic  
 237 compounds play a role in preventing cell damage caused by free radicals, thus preventing inflammation (Kuan et al.,  
 238 2025). In addition to saponins and phenolic compounds, *Lantana camara* also contains terpenoids, which are useful  
 239 for reducing inflammatory activity (Zubair & Ahmad, 2019). The anti-inflammatory properties of *Lantana camara*  
 240 can inhibit the inflammatory process in cuts in white Wistar rats, allowing the wound healing process to occur more  
 241 quickly.

242

243

## CONCLUSIONS

244 Based on the results of the research conducted, it was concluded that topical ointment of ethanol extract of patiwala  
245 leaves (*Lantana camara*) at a dose of 15% was able to accelerate the healing of incision wounds in diabetic rats as  
246 evidenced by changes in wound diameter, number of fibroblast cells and collagen fiber formation on the 21st day after  
247 therapy.

248

## ACKNOWLEDGEMENTS

249 The author would like to thanks to Poltekkes Kemenkes Kendari for funding this research through DIPA number  
250 HK.02.03/F.XXXVI/1666/2024.

251

## AUTHORS' CONTRIBUTIONS

252 Tuty Yuniarty & Zulfikar Ali Hasan carried out the laboratory work. Theosobia Grace Orno wrote the manuscript. Ahmat  
253 Rediansya Putra & Ratih Feraritra Danu Atmaja analyze the data. Theosobia Grace Orno & Anita Rosanty designed the  
254 study. Theosobia Grace Orno supervised the manuscript. **All authors read and approved the final version of the manuscript.**

255

## COMPETING INTERESTS

256 **The authors declare that there are no competing interests.**

257

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