

2 **Chemical composition, Antibacterial and Antioxidant activities of**
3 **extracts from dry leaves and ash-dry leaves of *Luffa cylindrica* (L.)**
4 **Roem cultivated in Vietnam**

5
6 **Abstract**

7 *Luffa cylindrica* (L.) Roem was traditionally used to treat stomachaches, as antihyperlipidemic
8 and antioxidant, particularly for atherosclerosis therapy, as a suppository to cure constipation and
9 spleenopathy, as an anthelmintic, carminative, emmenagogue, galactagogue, and as an antiseptic.
10 Therefore, the aim of this study was to study the chemical composition, antibacterial and
11 antioxidant properties of an extract from dry leaves (LuL) compared to ash-dry leaves (LuA) of
12 *Luffa cylindrica* (L.) Roem.

13 Our results showed the physical-chemical and phytochemical properties, antioxidant activity,
14 antibacterial activity and the metal ion content of both extracts. The comparison between the
15 extract from dry leaves (LuL) and ash-dry leaves (LuA) of *Luffa cylindrica* (L.) Roem, showed
16 difference in quantitative phytochemical determination of cardiac glycosides, alkaloids,
17 phenolics, flavonoids, and triterpenoids. This findings may be related to the LuA sample being
18 burned out incompletely into ash. Furthermore, this study showed that the activity of extracts
19 from dry leaves (LuL) and ash-dry leaves (LuA) contained both bacteriostatic and bactericidal
20 effects. The antioxidant properties observed may be related to , the flavonoids content. The
21 presence of metal ions in both extracts, which may contribute to the known wound healing
22 effects, deserves further study.

23 **Keywords:** *Luffa cylindrica* (L), inorganic herbal metal ions , antibacterial, antioxidant.

24 **1. Introduction**

25 *Luffa cylindrica* (L.) Roem belongs to the family *Cucurbitaceae* [1]. The origin of *Luffa*
26 *cylindrica* is believed to be in South America [2]. *Luffa cylindrica* is commonly grown in
27 Guinea, Ivory Coast, the Philippines, India, and China [3]. The flowers, buds, and young leaves
28 can be used as food [4]. When the fruit is old and dry, cleared of its epidermis and seeds, it gives
29 an excellent sponge called "vegetable sponge," which can be used as a body scrub, pot, or
30 appliance. It is also used as a heavy metal absorber for dehydration [5], [6]. The seed oil is
31 edible. In America, oil is used as an ingredient in soapmaking [7], [8]. The traditional use has
32 been reported in Africa, China, Vietnam, Cambodia, Thailand, Laos, and the Philippines [9]–
33 [11]. The fruit is used as a galactagogue, the roots as a hydragogue and purgative [12], and the
34 root and the whole plant as a suppository to cure constipation [13]. Seed acts as an anthelmintic
35 drug, an inducing vomiting drug, and a laxative [9], [14], [15].

36 The leaves are prescribed for skin diseases, treat wounds, reduce swelling, treat
37 stomachaches, antihyperlipidemic and antioxidant, particularly for atherosclerosis therapy [16].
38 Freshly crushed leaves act as emmenagogues, blood detoxifiers, and are used to treat papules and
39 swelling skin [17]. A decoction of leaves is used as a diuretic [18]. Past research has found that
40 leaf extract contains saponin, flavonoids, alkaloids, and cardiac glycosides, and the extract can
41 inhibit *Bacillus subtilis*, *Escherichia coli*, *Staphylococcus aureus*, and *Salmonella typhi* [19].
42 Aqueous extracts also have an oxytocic activity [13], [20].

43 The present knowledge of the wound healing process comprises coagulation, inflammation,
44 proliferation, formation and accumulation of fibrous tissues, collagen deposition,
45 epithelialization, contraction of the wound with the formation of granulation tissues, remodeling,
46 and maturation [17], [21]. The constituents of the plant extracts modulate one or more of the
47 above stages.

48 It was the endeavor to identify the active constituents responsible for antimicrobial activity,
49 free radical scavenging properties, stimulators of enhanced collagen production, and/or
50 angiogenesis promoters through the identification of lead scaffold chemical structures [20], [21].

51 Some studies have shown that *Luffa Cylindrica* is able to affect wound healing, which is a
52 wisdom of folk medicine in many countries [22], but in Vietnam it is used in a different way by
53 using only the leaves [23] to make the wound heal faster. Minerals in organics are known to have
54 an effect on wound healing, such as zinc and chromium shots, which speed up wound healing
55 [24]. However some report already showed on the trend of using *Luffa* leaves for wound healing
56 suggest that this may be a product that helps with diabetes [22], [25]. In our previous research,
57 the trace of traditional use of *Luffa* leaves was done in Hai Duong province, which is located in
58 the center of the Red River Delta with a total area of 1,668.28 km² and a population of more than
59 1.9 million people. The province has good conditions for agriculture, transportation, and
60 industrial production and plays an important role in the social and economic development of the
61 country. A total of six traditional medicine practitioners were interviewed for this survey.
62 Informed consent was obtained from all, and the survey was explained to them in detail,
63 including the information that the survey results may be published internationally. Findings
64 showed that *Luffa* leaf was used long ago by both traditional medicine doctors and the old
65 generation themselves to treat open wounds that were affected long-term by bacteria or fungi.
66 The conservative burned ash from *luffa leaf* was pound into dried powder and then applied to the
67 acne, boils, pressure ulcers, and fungal infection areas. The treatment was very effective in many
68 cases of pressure ulcers and fungal infections in the area between the toes during flood season.
69 The wounds were quickly healed and recovered. The aim of this study was to study the chemical
70 composition, antibacterial and antioxidant effects of extracts from dry leaves (LuL) compared to
71 ash-dry leaves (LuA) of *Luffa cylindrica* (L.) Roem. as a preliminary study of sample extracts
72 obtained from Vietnam with possible properties that may contribute to wound healing.

73 **2. Materials and Methods**

74 **Plant material, extraction, and chemicals**

75 *Luffa* leaves were collected from a Vietnamese farm in Hui Doung Province, Vietnam. Voucher
76 specimen No. 0023302 was identified and kept at the Herbarium of the Faculty of Pharmacy,
77 Chiang Mai University, Thailand. The chemical ingredients and solvent used for extraction of
78 the leaves and ash were of pharmaceutical grade and were purchased from Union Sciences Co.
79 Ltd., Thailand. The leaf was dried in a hot air oven at 60 °C and ground to powder (LuL). *Luffa*
80 ash (LuA) was prepared by burning the *Luffa* dried leaves at a normal temperature in open air
81 until the blackish-grey ash was obtained in an uncompleted burning condition. This process was
82 done by the local people.

83 **The Pharmacogenetic Evaluation of the Raw Material of Crude Dried Leaves**

84 A microscopic examination of powdered LuL was studied. The Thin Layer Chromatography,
85 moisture content, and extractive value were done according to Thai Herbal Pharmacopoeia V.I.
86 in order to prove the scientific database for further use. Two systems of developing solvents for
87 TLC plates were used: hexane: ethyl acetate (6:4) and dichloromethane: ethyl acetate (9:1). TLC
88 patterns were determined under UV light at 254 and 366 nm detectors. The plate was sprayed
89 with a freshly prepared anisaldehyde-sulfuric acid reagent (AS).

90 **Sample extraction**

91 Samples of LuL and LuA were ground to 60–80 mesh size with an electric grinder. Each sample
92 was extracted with 95% ethanol in a ratio of 1:10. Sonication was done under an ultrasonic
93 device for 1 hour, separated the clear parts, repeated three times, and then evaporated under
94 pressure.

95 **Quantitative Phytochemical Determination**

96 **Determination of Total Phenolic Content (TPC)**

97 The total phenolic content of the sample was examined by the Folin-Ciocalteu colorimetric
98 method modification [26]. The sample solutions (1 mL) were mixed with 5 mL of the Folin-
99 Ciocalteu reagent (diluted with distilled water in a ratio of 1:10). After 8 min, a sodium
100 carbonate solution (4 mL, 7.5% w/v) was added and incubated in the dark at room temperature
101 for 2 hrs. Finally, the absorbance of the test samples was measured at 765 nm by a Milton Roy
102 Spectronic 21D spectrophotometer. The gallic acid equivalent values (GAE mg/100g) were
103 calculated and compared with the standard curve of gallic acid. All tests were done in triplicate.

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105 2) Determination of Total Flavonoids Content: The extract solution (1 mg/mL in ethanol:
106 water 1:1, 1 ml) was mixed with a 2% AlCl₃ solution (1 ml) and kept in the dark at ambient
107 temperature for 25 minutes. The absorbance was determined at 415 nm compared with rutin.

108 3) Determination of Total Alkaloids Content: The extract solution (0.1 µg /ml in purified
109 water, 1 ml) was mixed with phosphate buffer solution (pH 4.7, 2 ml). The bromocresol green
110 solution (2 ml) was added to the mixture and then extracted with 1, 2, and 2 ml of chloroform.
111 The absorbance was determined at 415 nm by using berberine chloride as a standard.

112 4) Determination of Total Triterpenoids Content: The extract solution (1 mg/ml in glacial
113 acetic acid, 200 µl) was mixed with a 5% vanillin-acetic acid solution (1 ml) and sulfuric acid
114 (1.8 ml). The sample solutions were allowed to stand at 70°C for 30 minutes and then cooled
115 down to room temperature before adding glacial acetic acid (2 ml). The absorbance of sample
116 solutions was measured at 573 nm by using ursolic acid (Tokyo Chemical, Japan) as a standard.

117 5) Determination of Total Cardiac Glycoside Content: The extract (1 mg/ml in 50%
118 aqueous ethanol, 1 ml) was mixed with 1 ml of freshly prepared Baljet's reagent (95 mL of 1%
119 picric acid and 5 ml of 10% sodium hydroxide solution). The reaction mixture was incubated for
120 1 h, then diluted with 2 ml of purified water. The absorbance was quantitatively determined at
121 495 nm by using digoxin as a standard.

122 **Biological activities of LuL and LuA extract**

123 **Determination of Antioxidant Activity**

124 The *Diphenylpicryl-hydrazyl* (DPPH) radical scavenging assay was used for determination using
125 the method described by Wu et al., 2005 [27]. The solution of DPPH radicals was prepared in
126 methanol (81.2 mM in methanol). The sample solution (1 mL) was mixed with 5 mL of DPPH
127 solution. The mixtures were vigorously shaken and left for 30 minutes in the dark. The
128 absorbance was measured at 517 nm using methanol as a blank. 5 mL of DPPH solution in 5 mL
129 of methanol were used as controls.

130 Percent inhibition = [(A control-A sample)/A control] x 100. Where A control is the absorbance
131 of only DPPH radical solution, A sample is the absorbance of a sample mixed with DPPH radical

132 solution. The sample concentration providing 50% inhibition (IC₅₀) was calculated from the
133 graph of inhibition percentage against sample concentration.

134 Antibacterial activity

135 The antibacterial activity of LuA extract and LuL extract were determined at minimal
136 inhibitory concentration (MIC) and minimal bactericidal concentration (MBC) using the broth
137 dilution method (CLSI, 2018). A single colony of tested bacteria were inoculated in Mueller-
138 Hinton broth (MHB) and incubated at 37°C for 18 to 24 hours. The extracts were dissolved with
139 dimethyl sulfoxide (DMSO) at a concentration of 500 mg/mL as a stock solution. The MHB (0.1
140 mL) was dispensed in a sterile 96-well plate. The stock solution of the extract (0.1 mL) was
141 added to the first test well and the extract was serial two-fold diluted in each well in the 96-well
142 plate. The bacterial culture (McFarland standards No. 0.5) was added to all wells. The mixture of
143 extract and bacterial culture were incubated at 37°C for 18 to 24 hours. The MIC was determined
144 from the lowest concentration of the extract not showing any bacterial growth. Therefore, MBC
145 value was determined from each well showing no visible bacteria growth of as compared with
146 the bacterial growth control. The culture broth not displaying bacterial growth was streaked on
147 Mueller-Hinton agar (MHA), and bacteria colonies were determined after incubating. The MBC
148 value was the lowest concentration inhibiting bacterial growth by 99.9%. [36]

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150 Identification of the metal ions in the samples

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152 The LuL and LuA dried samples were sent for checking for metal ions at the Central
153 Laboratories (Thailand) Co., Ltd. The analysis was performed based on the EPA 3052 method (to
154 prepare the samples) and the analysis performed using Inductively Coupled Plasma Optical Emission
155 spectroscopy (ICP-OES).

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158 3. Result

159 3.1 Microscopic Identification.

160 The Microscopic characteristics of *Luffa leaves* are shown in Figs. 1-3.

161 The diagnostic characters are:

- 162 1. In surface view, the fragments of the lamina in the upper polygonal epidermis and lower
163 epidermis were wavy in outline. Anomocytic stomata were also present on both surfaces.
- 164 2. Palisade mesophyll was usually found in surface view; it is composed of cells with thin
165 walls, circular in outline, containing abundant chloroplasts.
- 166 3. The fragments of spongy mesophyll show thin-walled parenchyma containing moderately
167 large chloroplasts with large intercellular spaces and air chambers.
- 168 4. The vascular strand was found in various sizes and views, some of which are associated
169 with spongy mesophyll.
- 170 5. The fragments of spiral and reticulated vessels in longitudinal view were not very
171 frequent.
- 172 6. The occasional fibers could be found in groups or solitary.
- 173 7. The occasional glandular trichome appeared as whole trichomes with stalk and head, or
174 fragments of them.
- 175 8. The very occasional tracheid fragments in longitudinal view.

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9. Starch grains were seldom found and accumulate in parenchymatous tissue.

UNDER PEER REVIEW

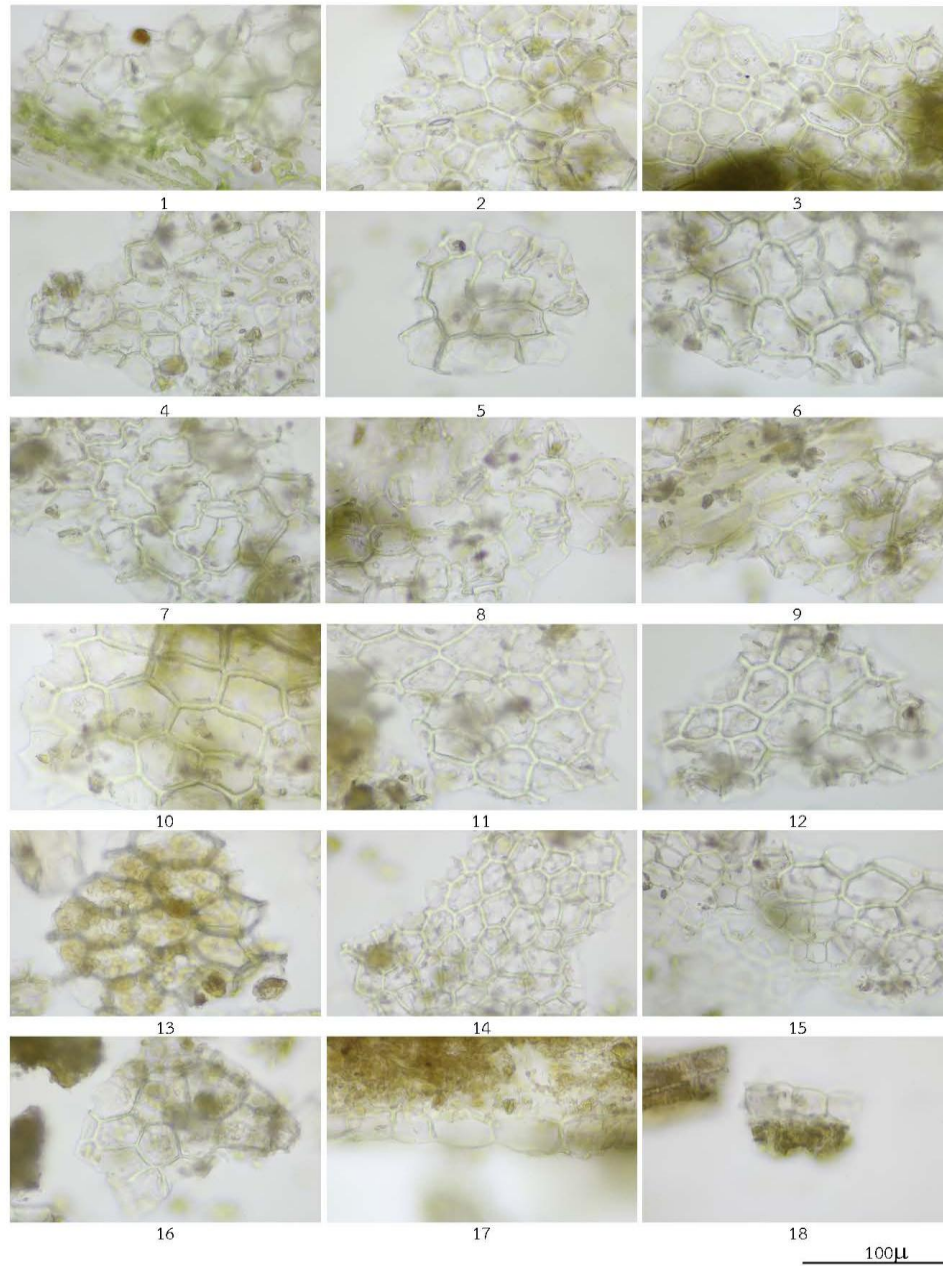


Fig.1 Powder drug of *Luffa cylindrica* leaf; 1-6 upper epidermis showing stoma, 7-8 lower epidermis showing stoma and wavy epidermis, 10 upper epidermis over vein with stoma, 11-16 upper epidermis with palisade underneath, 17-18 epidermis in sectional view

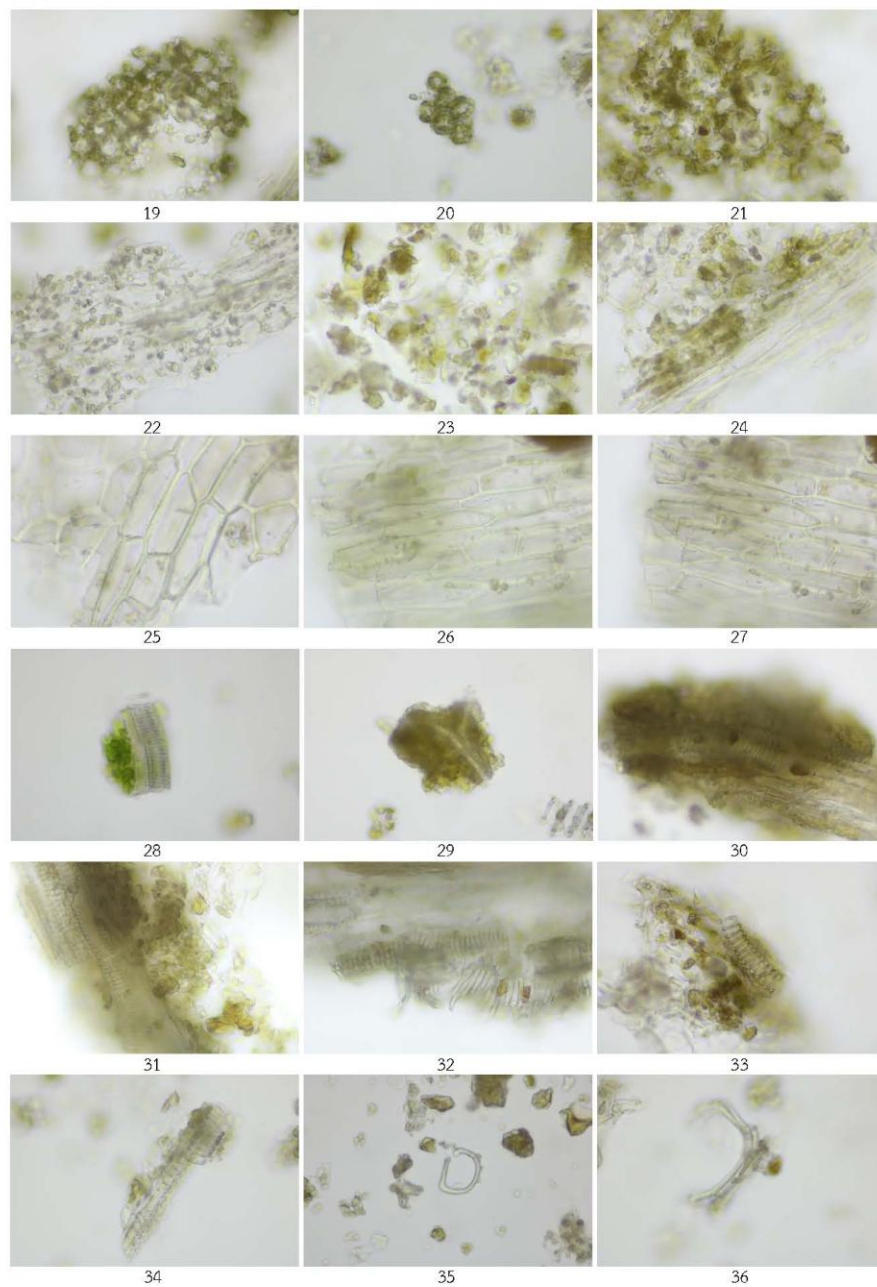


Fig.2 Powder drug of *Luffa* sp. Leaf (continuee) ; 19-21 palisade in surface view, 22-24 spongy mesophyll, 25-27 polygonal epidermis, 28-29 vascular strand of mesophyll, 30-32 vascular bundle, 33-34 spiral vessel associated with chlorenchyma, 35-36 fragments of vessel

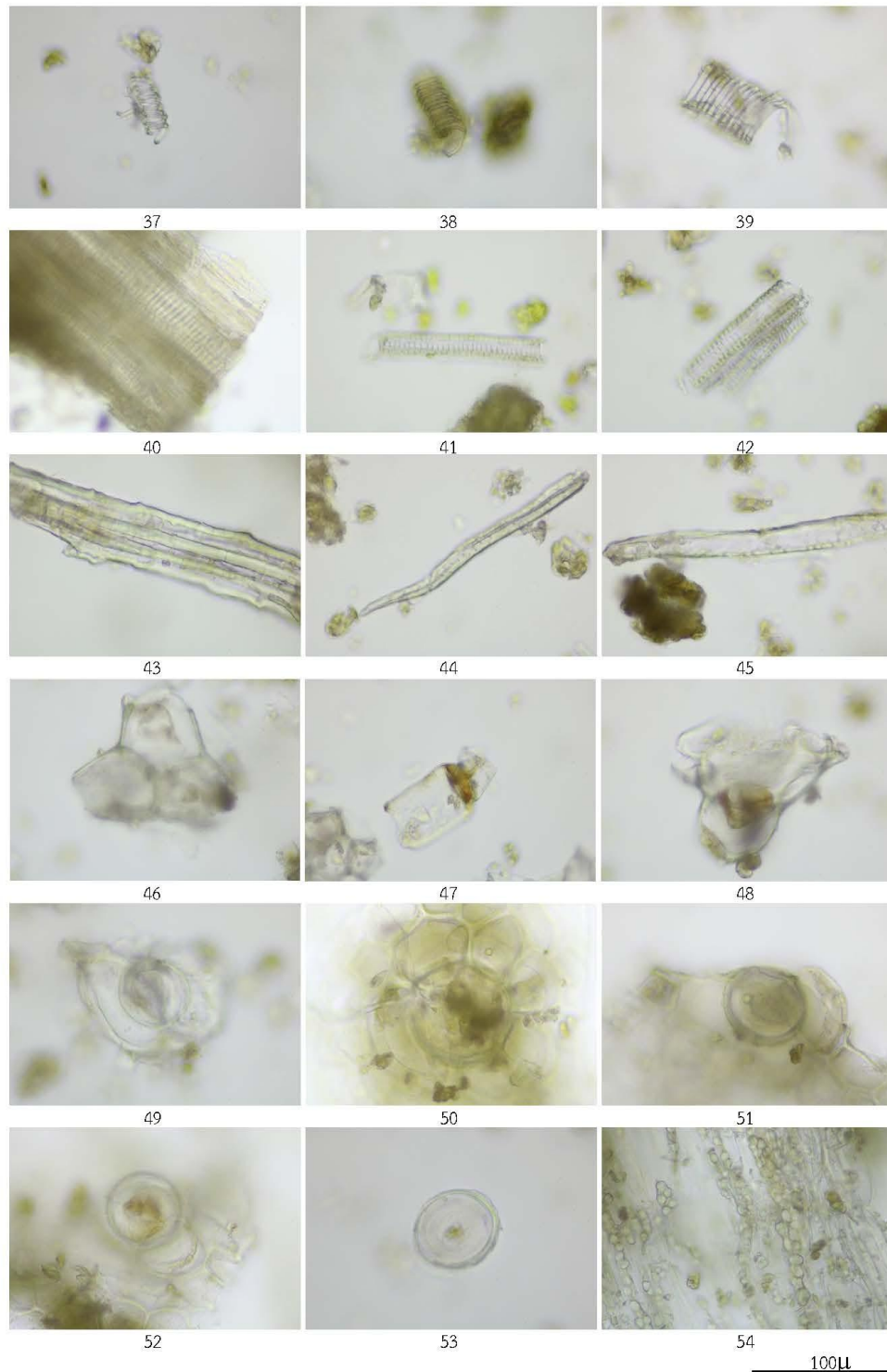


Fig.3 Powder drug of *Luffa cylindrica* Leaf (continuee) ; 37-38 spiral vessel, 39 reticulated vessel, 40 xylem element showing reticulated vessel and xylem fiber, 41-42 tracheid, 43 group of fiber, 44-45 solitary fiber, 46-48 trichome stalk, 49-52 capitate stalked trichome in surface view, 53 capitate stalked trichome head, 54 starch grains

194 **3.2 Physico-chemical Identification**

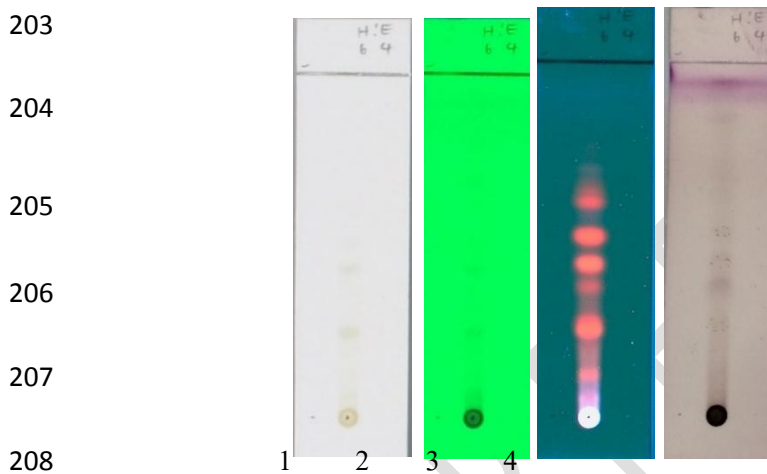
195 The samples were tested as in the Thai Herbal Pharmacopeia. The physico-chemical examinations were
196 as follows: Loss on drying, total ash, ethanol-soluble extractive value, and chloroform water extractive
197 value The mean values are presented in Table 1. TLC is shown as in Fig. 4, Fig. 5, Table 2, and Table 3.
198 From the results, the optimum system used in quality control of raw materials, LuL, should be: Solvent
199 system: Hexane: ethyl acetate (6:4), with a UV 366 detector.

200 **Table 1** Pharmacogenetic characteristic of LuL

Specification	Content (%)
Loss on drying	13.01±0.20 (W/W)
Total ash	21.9144±0.59 (W/W)
Ethanol-soluble extractive value	11.7900±0.17 (W/W)
Chloroform water extractive value	18.3633±0.22 (W/W)

201 The data represent mean values of three replicates ± SD.

202 Thai Herbal Pharmacopoeia 1995 Volume 1 pp.123, 126



209 Fig.4 Sample before (1) and after (2) detected with UV 254 detector, with UV
210 366 detector (3) and after spraying with Anisaldehyde – sulfuric acid (AS)
211 (4). Solvent system: Hexane: Ethyl acetate (6:4)

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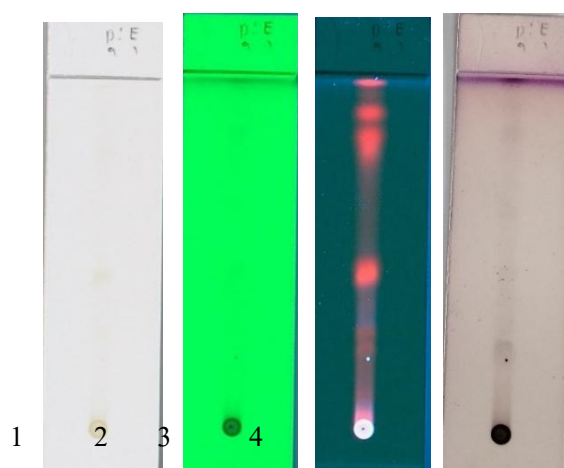
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Fig.5 Sample before (1) and after (2) detected with UV 254 detector, with UV 366 detector (3) and after spraying with Anisaldehyde – sulfuric acid (AS) (4). Solvent system: Dichloromethane: Ethyl acetate (9:1)

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Table 2 TLC of LuL extract in Hexane: Ethyl acetate (6:4)

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Rf value	Visual inspection	Spot color UV254	Spot color UV366	Spraying with Anisaldehyde – sulfuric acid
0.14	-	-	orange	-
0.26	opaque	opaque	orange	-
0.36	-	-	orange	purple-gray
0.48	opaque	opaque	orange	-
0.50	opaque	opaque	orange	-
0.60	-	-	orange	-

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Table 3 TLC of LuL-extract in Dichloromethane: Ethyl acetate (9:1)

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Rf value	Visual inspection	Spot color UV254	Spot color UV254	Spraying with Anisaldehyde – sulfuric acid
0.26	-	-	orange	purple-gray
0.30	-	-	orange	-
0.44	opaque	opaque	orange	-
0.84	-	-	orange	-
0.90	-	-	orange	-
0.98	-	-	orange	-

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3.3 Quantitative Phytochemical Determination

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Phytochemical screening of LuL and LuA demonstrated the presence of cardiac glycosides, alkaloids flavonoids, phenolics and triterpenoids as shown in Table 4.

261 Table 4 Contents of bioactive compounds referenced to their standards

Extract No.	Name of the extract	Cardiac glycosides		Alkaloids		Phenolics		Flavonoids		Triterpenoids	
		mg digoxin /g extract	SD	mg berberine /g extract	SD	mg gallic acid /g extract	SD	mg Rutin /g extract	SD	mg Ursolic acid/g extract	SD
1	LuA	22.3	0.17	10.7	0.59	16.7	0.24	20.3	1.02	6.9	0.68
2	LuL	63.7	0.75	20.1	0.95	59.0	0.98	47.8	0.31	46.1	0.34

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263 LuA contained secondary metabolites as follows: cardiac glycosides 22.3±0.17 (mg digoxin/g extract),
264 flavonoids 20.3±1.02 (mg rutin/g extract), phenolics 16.7±0.24 (mg gallic acid/g extract), alkaloids
265 10.7±0.59 (mg berberine/g extract), terpenoids 6.9±0.68 (mg ursolic acid/g extract). LuL contained
266 secondary metabolites as follows: cardiac glycosides 63.7±0.75 (mg digoxin/g extract), phenolics
267 59.0±0.98 (mg gallic acid/g extract), flavonoids 47.8±0.31 (mg rutin/g extract), triterpenoids 46.1±0.34
268 (mg ursolic acid/g extract), alkaloids 20.1±0.95 mg (berberine/g extract). The results show that LuA still
269 contained all active compounds but in a lower quantity than sample LuL. This could be explained by the
270 fact that the sample was burned out into ash incompletely.

271 3.4 Antioxidant activity of LuA and LuL extract

272 The DPPH radical scavenging activities of LuL extract showed an IC₅₀ value of 87.63 ± 10.00 mg/ml. or
273 LuL 87.63 mg/ml can remove 50 % of free radical, DPPH. Whereas LuA presented % inhibition
274 3.17±0.69 (mg/mL) or LuA at a concentration of 1 mg/ml reduce can remove 3.7 % of DPPH (*p* < 0.05).
275 This phenomenon showed the importance of phenolic compounds in DPPH's radical scavenging
276 activities.

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278 3.5 Antibacterial Activity of *Luffa* Leaf (LuL) Extract and *Luffa* Ash (LuA) Extract

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280 In this study, the inhibitory activities of leaf extract were investigated for the minimal inhibitory
281 concentration (MIC) and minimal bactericidal concentration (MBC) against pathogenic bacteria,
282 *Staphylococcus aureus*, *Pseudomonas aeruginosa*, and *Escherichia coli*. by broth dilution
283 method. The result demonstrated that LuL extract revealed an inhibitory effect on all tested
284 bacteria. Moreover, LuL extract showed the lowest MIC and MBC values of 125 mg/ml against
285 *Staphylococcus aureus* and *Pseudomonas aeruginosa*, followed by *Escherichia coli* with MIC
286 and MBC values of 250 mg/ml (Table 5). But LuA extract showed the lowest MIC and MBC
287 values of 125 mg/ml against *Staphylococcus aureus*, *Pseudomonas aeruginosa* and *Escherichia*
288 *coli*. This revealed that LuA extract possessed more effective antibacterial effect on *E. coli* than
289 LuL.

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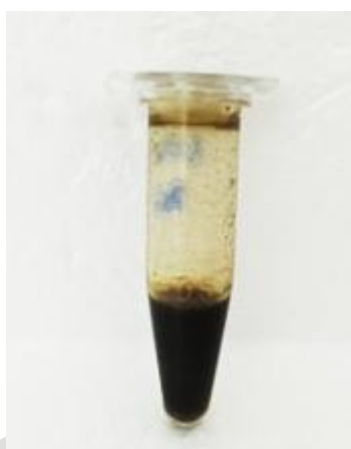
291 Table 5 MIC and MBC values of LuL extract and LuA extract against pathogenic bacteria

sample	MIC and MBC (mg/ml)					
	<i>S. aureus</i>		<i>Ps. aeruginosa</i>		<i>E. coli</i>	
	MIC	MBC	MIC	MBC	MIC	MBC
LuL extract	125± 0.0	125± 0.0	125± 0.0	125± 0.0	250± 0.0	250± 0.0
LuA	125± 0.0	125± 0.0	125± 0.0	125± 0.0	125± 0.0	125± 0.0

292 Data represents mean values of three replicates ± SD

293 Therefore, LuL and LuA extracts showed strong antimicrobial activity against pathogenic
 294 bacteria, which are usually present on the skin. In addition, the extract demonstrated a high content of
 295 phenolics and flavonoids that served the antimicrobial activity. Flavonoids are effective both in directly
 296 damaging the envelope of Gram-negative and Gram-positive bacteria [30]. Thus, this study implied that
 297 the LuL and LuA extracts contained both bacteriostatic and bactericidal effects.

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302 **Fig. 6** LuA extract at 500 mg/ml in DMSO

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306 **Fig. 7** LuL extract at 500 mg/ml dissolve in DMSO

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310 **3.6** Metal ion in the LuA and LuL

311 The results of the metals in the sample were as presented in Table 6. The minerals that were
 312 higher in LuL were high in iron, zinc, and copper, but also they were in larger quantities in LuA.
 313 Interestingly, a high-level supplement of trace metal not only improved growth performance but
 314 also reduced footpad lesions by improving the wound healing process via promotion of collagen

315 synthesis, decomposition and organization, cell migration, matrix remodeling, angiogenesis, and
316 regulation of inflammation [31]. The role of each mineral in wound healing should be the subject
317 of future studies.

318 **Table 6** Metals in LuL and LuA by Inhouse Method Based on EPA 3052, by ICP-OES
319 Technique

Test Item	Result		Unit
	LuL	LuA	
Arsenic (As)	0.9	2.04	mg/kg
Cadmium (Cd)	0.15	0.29	mg/kg
Copper (Cu)	7.25	25.34	mg/kg
Iron (Fe)	148	1332	mg/kg
Lead (Pb)	1.56	8.83	mg/kg
Mercury (Hg)	Not detected	Not detected	mg/kg
Zinc (Zn)	43.28	189	mg/kg

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322 Discussion

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324 Herbal medicines have become a popular form of therapy in developing countries. They are
325 believed to be nontoxic, with little side effects compared to modern drugs. This is in accordance
326 with the common use of *Luffa cylindrica* (L.) Roem in several countries worldwide for the
327 traditional management of diseases. There was evidence from a few pharmacological
328 investigations that *Luffa cylindrica* possessed anti-inflammatory, analgesic, antipyretic,
329 hypoglycemic, antibacterial, antifungal, antiviral, anthelmintic, antioxidant, anticancer,
330 hepatoprotective, antiemetic, wound healing, immunological, bronchodilation, reproductive effect,
331 and in the treatment of cataract [13], [20], [22], [25].

332 The aim of the present study was to compare the chemical composition, antibacterial, antioxidant,
333 and wound healing activity of extract from dry leaves (LuL) and the ash dry leaves (LuA) of *Luffa*
334 *cylindrica* (L.) Roem, through the determination of the physico-chemical effect, phytochemical effect,
335 antioxidant activity, antibacterial activity, and metal ion contents.

336 The present study show that LaH and LuL can be candidate for therapeutic treating of pathogenic
337 skin infection and antioxidation. The determination of the antibacterial activity of *Luffa* Leaf (LuL)
338 extract and *Luffa* ash (LuA) extract revealed that LuA extract was more effective in antibacterial
339 activity than LuL. This result was in accordance with with studies by [17], [19], [21], and may
340 be explained by the high content of phenolics and flavonoids of the extract that served the
341 antimicrobial activity. Flavonoids and phenolic compound presented in this study, are
342 component in a variety of medicinal, pharmaceutical, nutraceutical and cosmetic applications.
343 This is attributed to their antioxidative, anti inflammatory, antimutagenic, anticarcinogenic
344 properties and capacity to modulate key cellular enzyme functions [35]

345 In LuA, more metal ion items were detected, such as arsenic (As), cadmium (Cd), copper
346 (Cu), iron (Fe), lead (Pb), mercury (Hg), and zinc (Zn). This is in accordance with the preview
347 study by [24], [31], [33] The role of each mineral in wound healing should be the subject of
348 future studies.

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351 Conclusion

352 The results of the present study clearly indicated that the crude methanol extract of *Luffa cylindrica*
353 did produce strong antimicrobial activity against pathogenic bacteria on the skin. In addition, the extract

354 demonstrated a high content of phenolics and flavonoids that may serve as antimicrobial agents. Thus,
355 this study implied that the activity of extracts from dry leaves (LuL) and ash-dry leaves (LuA) contained
356 both bacteriostatic and bactericidal effects.

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