



1 Characteristic Change in Activity of *Lantana camara* Triptene

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5 The bioactive molecule triptene of *Lantana camara* exhibited therapeutic applications. It was separated by vacuum distillation followed by
6 gel chromatography procured 16-18 h extraction time. Exhibited single of pure triptene at 4.693 and yield of triptene was 0.13, 0.27 and
7 0.24 mg for the year 2013, 2014 and 2015, respectively. Characteristic change of eluted fractions assessed through TLC, HPLC, UV and
8 FTIR spectrophotometer observe triptene approach for further application.

9 **Keywords:** *Lantana camara*, Triterptene, Pharmacological activity, Behavioural activity.

INTRODUCTION

10 A renowned notorious weed; *Lantana camara* L.
11 (Verbanaceae) also known as wild or red sage. It is a perennial
12 shrub, brought to India some 80 years ago from South America.
13 Popular ornamental garden *Lantana camara* has become exotic
14 and spread to different regions of the country [1]. It's consti-
15 tuent numerous secondary bioactive metabolites that exhibited
16 therapeutic and traditional significance [2]. The leaf extract
17 of *Lantana camara* is rich in different types of plant secondary
18 metabolites such as terpenoids, steroids and polyphenols inclu-
19 ding flavonoids, tannin [2,3]. Bioactive metabolites isolated
20 from *Lantana camara* leaves namely triptene/triterptenoids
21 possess, anti-inflammatory [4,18], antipyretic activity [4], anti-
22 filarial activity [5], antitumor activity [6] antithrombin activity
23 [7], antimicrobial activity [10], oxidizing activity [11] anti-
24 cancer activity [12,16]. Researchers have studied the potential
25 of bioactive triterptenes in alcoholic extract [14-17]. The main
26 chemical constituent; Triptene is acts as promoting agent [9]
27 and used for the treatment of chicken pox, measles, asthma,
28 ulcers, swellings, eczema and high blood pressure [2]. Triterptene
29 also exhibits the anticoagulant and anti-inflammatory activity
30 and has potential to inhibit the aggregation of proteins [17,18].
31 Therefore, the present study focused on the extraction of triptene
32 from *Lantana camara* leaves by conventional methods and to
33 observe the change in behaviour of active molecule/triptene
34 throughout the methodology for the perspectives of it in further
35 industrial application as well chemotherapy of life. Quanti-
36 fication of triptene was carried out by spectrophotometric
37 method and monitored tread by TLC, UV-visible and FTIR
38 spectrophotometer without any further treatment. The activity

of triptene was optimized with the aid of different factors such 39
as extraction time. Optimal conditions of triptene provided good 40
alternatives for the further pharmacological applications. 41

EXPERIMENTAL

42 **Collection of plant leaves:** A *Lantana camara* healthy 42
fresh leaves was collected from the campus of Vidya Prasarak 43
Mandal's B.N. Bandodkar College of Science, Thane (W) India 44
in month of September 2015, September 2014 and 2013 to 45
compare yield. Segregated leaves were washed, dried. Pulve- 46
rized powder stored in self sealing bags until used. 47

48 **Isolation of triptene from *Lantana camara* Linn leaves:** 48
Lantana camara leaves powder (100 g) was treated with increase 49
in volume of methanol and serially refluxed for 0.5 to 18 h. 50
Solvent was removed under vacuum (13-14 mm/Hg and disti- 51
llation temperature up to 58 °C). Obtained residue was suspen- 52
ded distilled water after filtration, the residue was dissolve in 53
a methanol:water (1:7) mixture and extracted with ethyl acetate 54
(2 × 25 cm³) and with *n*-butanol ((2 × 25 cm³). The ethyl acetate 55
layer was concentrated under reduced pressure to get crude 56
triptene. Each step was monitored timely with TLC and quali- 57
tative test. Formulated triptene was measured from 0.5 to 18 h 58
to find out extraction time. The whole process is repeated to 59
observe behaviour/yield active compound/triptene collected in 60
month of September 2013, September 2014 and September 61
2015. 62

63 **Column chromatography of *Lantana camara* triptene:** 63
Crude triptene was loaded over silica gel (30 g) column (60- 64
120 mesh) and 32 elutions were collected using chloroform: 65
methanol (9:1) as eluting solvent. The active fraction 12th was 66

67 rechromatogram since it showed five spot on TLC. After
68 rechromatography active fractions (19, 22 and 23) were pulled,
69 concentrated and rechromatography was carried out using *n*-
70 hexane with successive increasing amount of acetone. Different
71 fractions were monitored with TLC using ethyl acetate:methanol:
72 water (9:3:1) throughout the process. Purity of active com-
73 pound (50 μ L) checked on HPLC using C-18 column having
74 mobile phase A-0.1 % trifluoroacetic acid in water (80:20)
75 and Mobile phase B-acetonitrile on Alliance 2695 model.

76 Characterization of *Lantana camara* tripene

77 **Qualitative test for triterpene:** Active fraction from silica
78 gel column (50 μ L) was treated with 0.5 cm³ chloroform and
79 warmed for about 0.5 h 2-3 drops of concentrated H₂SO₄
80 was added and mixed well. The presence of triterpenoid was
81 observed by change in colour [9].

82 **Spectrophotometric study:** The active fractions from
83 silica gel column chromatography were explored to UV-1800
84 Shimadzu spectrophotometer to observed exact λ_{\max} . The
85 observe spot on TLC were dissolved in diethyl ether. UV and
86 FTIR (Thermo scientific Nicolet iS5) were carried out for each
87 fractions monitored from 190 to 1100 nm wavelength.

RESULTS AND DISCUSSION

88 *Lantana camara* Linn. is renowned weed for therapeutic
89 activity and economical wellness. Hence, role of its secondary
90 metabolites need to facilitate in drug delivery system. Attention
91 of present research especially on the change in behaviour
92 of active molecule throughout the processing. Behavioural
93 activity is depends on seasonal changes and attribution of time,
94 volume of solvents and effective concentration of reactant used
95 to achieved the tripene yield (Tables 1 and 2). Organic solvents
96 contributed exceptionally low yield showing very little diffe-
97 rence in physicochemical properties of pentacyclic tripene
98 [10,11]. Problems with the separation of the complex triterpene
99 mixtures result in a very small quantities, usually a few milli-
100 grams of individual compounds being isolated. Thus, for the
101 structure elucidation non-degradative techniques are preferred.
102 The compound is being a new structure or to confirm its simi-
103 larity to the known standards. TLC analysis may be of a great
104 help along with UV, IR.

105 Ethyl acetate does not able to extract tripene completely
106 consequently to achieve tripene the methanol-water mixture
107 was used. Extracted yield (10.34 g) showing its λ_{\max} 228 nm
108 (Fig. 1A) that was treated with butanol and ethyl acetate pro-
109 cured producible results (0.95 mg) of tripene. The qualitative
110 test was conducted for crude compound showing red colour
111 after adding of H₂SO₄.

112 The obtained crude tripene is 0.95 mg from 100 g of dried
113 leaves powder in 18 h refluxed time. Serially refluxed and
114 process 100 g of leaves dried powder treated with as per process
115 shown increase in yield of product till 16 and 18 h (Table-1).

TABLE-1
EXTRACTION TIME OF *Lantana camara* Linn.

Time (h)	Colour	Weight (mg)
0.5	Greenish colour liquid	–
1	Green brown colour	–
2	Green brown colour	0.12
4	Faint brown colour	0.16
8	Brown colour	0.23
10	Brown colour	0.23
12	Brown colour	0.24
14	Dark brown colour	0.48
16	Dark brown colour	0.53
18	Dark brown colour	0.95
36	Black colour	0.54
40	Black colour	0.54
48	Mask clear solution	0.51

116 There is no change in product and colour of the compound up
117 to 40 h after the product is drops its quantity and characteristics
118 including colour. At 48 h obtained product is very dark and
119 sticky mask can removed out from the dish. The maximum
120 extraction time is 16 to 18 h was preferred for characterization.

121 The process of separation of tripene was carried out for
122 the collected in the year serially September 2013, 2014 and
123 2015 to ensure yield that is shown in Table-2. Ratio of crude
124 to pure compound isolated consistently shown negligible diffe-
125 rence, which may be due to geographical activity and human
126 analytical error. HPLC of fraction 19 showed single peaks at
127 4.693' and 4.980' min (Fig. 2).

128 After crude extract (0.95 mg) was loaded on column
129 chromatography and rechromatogram. The active fraction 12th
130 was monitored by TLC. Five yellowish coloured spot observed
131 for fraction 12th having R_f values are 0.35, 0.54, 0.76, 0.85,
132 0.93 and respective UV spectra was 294, 416, 236, 235 and
133 211 nm. After rechromatography of Fraction 12th; procured
134 three active fractions (19, 22 and 23). Fraction 19 and 23 pro-
135 cured the single spot (R_f 0.97 and 0.45) and fraction 22 showed
136 three spot (R_f 0.36, 0.50, 0.84) on TLC. The fraction 12th was
137 segregated in three active fractions on rechromatography may
138 be due to the matrix may functionalized affinity support works
139 on spacer and support matrix which eliminates toxic reagents
140 which used during isolation and reaches occurs if any.

141 Crude tripene showed λ_{\max} 228 nm while after purification
142 it showed fraction 19 (λ_{\max} 409 nm), fraction 22 (λ_{\max} 235 nm)
143 and fraction 23 (λ_{\max} 232 nm), respectively. The fraction 22
144 and 23 may be the bifurcation of elution. Hence, the observed
145 λ_{\max} 235 nm and λ_{\max} 232 nm. This may be due to ($\pi \rightarrow \pi^*$)
146 transition which is stabilized by H-bonding in diethyl ether. It
147 lowers the energy and shift wavelength from 232 nm to 235nm
148 (Fig. 1B). Crude tripene showed after purification it showed
149 fraction 12th (λ_{\max} 416 nm) and rechromatography fraction
150 19 (λ_{\max} 409 nm) showed the divergence of elutions. The
151 observed λ_{\max} 416 nm and λ_{\max} 409 nm it showed the π^* level
152 will be higher and $n \rightarrow \pi^*$ transition will be higher energy.

TABLE-2
SEASON WISE EXTRACTION YIELD OF CRUDE AND PURE ACTIVE COMPOUND FROM 100 g OF *Lantana camara* Linn.

Year	Crude (mg)	Pure (mg)	Ratio	R _f value	UV (nm)
September 2013	0.83	0.24	0.29	0.72, 0.73, 0.83	435
September 2014	0.95	0.27	0.28	0.76 0.85	409
September 2015	0.53	0.13	0.25	0.36, 0.45, 0.50, 0.84, 0.97	409, 235, 232

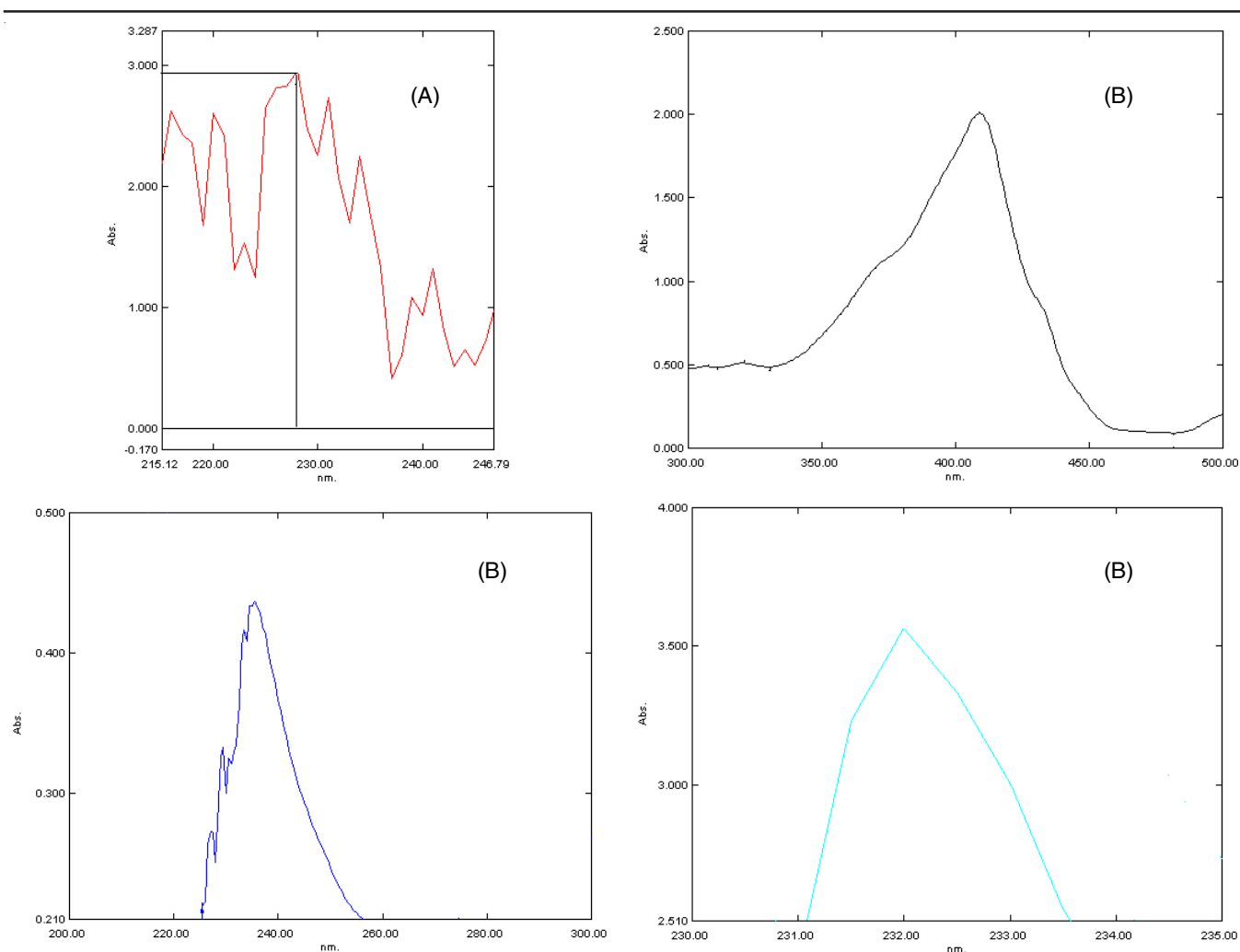


Fig. 1. UV maxima of *Lantana camara* leaves (A) crude active fraction/tripene before column chromatography showed λ_{\max} 228 nm (B) after rechromatography showed fraction 19 (λ_{\max} 409 nm, fraction 22 (λ_{\max} 235 nm) and fraction 23 (λ_{\max} 232 nm)

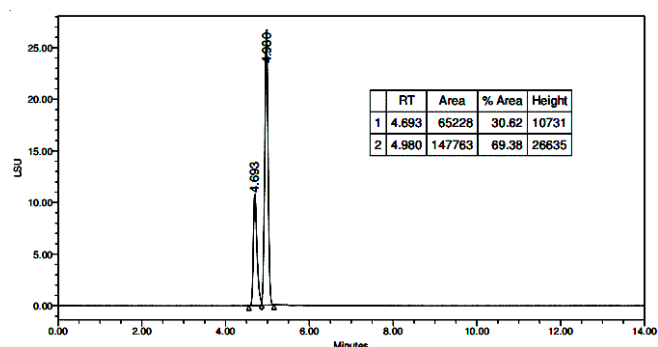


Fig. 2. Pure tripene compound (fraction 19 50 μ L) provoked single peak on HPLC

153 Hence, shift lower energy (red shift) to higher energy (blue
154 shift). It is higher energy and shift wavelength from 409 nm to
155 416 nm. When neutral solvent was used as solvent system,
156 tripene has not exhibited any identifiable spot in iodine and also
157 under long wavelength and short wavelength in UV chamber
158 while ethyl acetate:methanol:water (9:3:1) was procured
159 yellowish colour spot. The colour observed in visible light
160 and the ultraviolet fluoresces found more distinguishing. After
161 vacuum distillation, the fraction 22 observed Brown greenish
162 in colour having R_f value 0.85.

FTIR after and before column chromatography: The
163 intermolecular H bond does not take place as the molecules
164 are widely separated. Increasing concentration of solvent for
165 extraction was produce sharp band at frequency 2927 cm^{-1} ;
166 due to O-H stretching in vacuum distillation fraction. The
167 bending vibration of methyl group occur at 1462 cm^{-1} and 1467
168 cm^{-1} (fraction 22). The experiment reveals that the band 1462
169 cm^{-1} , 1467 cm^{-1} is due to α -methylene group 1456 cm^{-1}
170 represent β -methylene group (Fraction 19). Confirming the
171 compound is aromatic in nature substitution pattern of the ring
172 C-H bond in the region 668 cm^{-1} . Strong absorption band at
173 the region around 1700 cm^{-1} . Consistently, all fractions due to
174 C=O stretching indicates that compound may be ketone,
175 aldehyde, ester *etc.* At frequency 1716 cm^{-1} pentanone shown
176 C=O stretching vibration in conjugated ketone. The C-O
177 stretching in vacuum distilled fraction at 1232 cm^{-1} increases
178 the resonance character of double bond in the ring, which is
179 disappearance the frequency in fraction spectra since tripene/
180 triterpenoids have diverse chemical structures and bioactivities
181 [11].
182

FTIR spectrum before column chromatography: 3695
183 cm^{-1} (O-H stretching (acid free)), 3683 cm^{-1} (O-H str), 3654
184 cm^{-1} (O-H str), 2927 cm^{-1} (CH_2 , CH_3 str, =CH stretching), 1710
185 cm^{-1} (C=O stretching), 1606 cm^{-1} (C=C str, NH_2 bending), 1513
186

187 cm^{-1} (C=C aromatic, NO_2 gr), 1462 cm^{-1} (CH_2 , CH_3 bend),
 188 1378 cm^{-1} (CH_2 , CH_3 bend, C-O-C stretching ether), 1232 cm^{-1}
 189 (C-OH stretching acid).

190 The FTIR spectra were carried out identify the possible
 191 biomolecules in selected plant *Lantana camara* Linn. The IR
 192 spectrum of the lantana tripene peak was observed as 1750
 193 cm^{-1} frequency of a C=O observed. Alkane group observed in
 194 2900 cm^{-1} frequency and ether group 1378 cm^{-1} O-H stretching
 195 acid free 3695 cm^{-1} , =CH str 2927 cm^{-1} , C=C 1600 cm^{-1} and
 196 OH, -C-O stretching acid is a 1232 cm^{-1} frequency observed.

197 Disappearance of strong intensity absorption 1378 cm^{-1}
 198 band in the rechromatogram. However, these bands are rarely
 199 useful they overlap other stronger absorption.

200 FTIR of after column chromatography

201 **Fraction 19:** 3504 cm^{-1} (aromatic O-H), 2925 cm^{-1} (alkanes
 202 strong intensity aldehyde weak intensity), 2360 cm^{-1} (alkanes
 203 m-w intensity, nitrile), 2341 cm^{-1} (C=C), 1733 cm^{-1} (conjugated
 204 aldehyde, C=O), 1699 cm^{-1} (aldehyde, amide), 1652 cm^{-1} (C=O
 205 ester, alkenes), 1635 cm^{-1} (alkenes C=C, amide), 1558 cm^{-1}
 206 (C=C, N=O), 1540 cm^{-1} (R- NO_2 group strong intensity), 1507
 207 cm^{-1} (C-Cl, C-N, C-C, C-O R- NO_2 group strong intensity),
 208 1457 cm^{-1} (CH_3 - CH_2 bend), 668 cm^{-1} (C-Cl, C-O, C-N, C-C).

209 **Fraction 22:** 3773 cm^{-1} (O-H), 3457 cm^{-1} (C-H, N-H, O-
 210 H bonding), 2924 cm^{-1} (C-H str), 2359 cm^{-1} (alkanes m
 211 intensity, nitrile), 1738 cm^{-1} (aldehyde, ketone, C=O g), 1659
 212 cm^{-1} (C=O amide, C=C alkenes isolated), 1640 cm^{-1} (C=O,
 213 alkanes isolated), 1534 cm^{-1} (C-C, C-N, C-O, N-H str vib),
 214 1502 cm^{-1} (C-C, C-N, C-O, amine), 1467 cm^{-1} (CH_2 bend),
 215 1414 cm^{-1} (CH_3 bend), 1351 cm^{-1} (C-C, C-N, C-O, C-Cl), 1022
 216 cm^{-1} (C-O alcoholic, ester, ether, C=O anhydride, C-N amine),
 217 669 cm^{-1} (C-C-C medium).

218 **Fraction 23:** 3441 cm^{-1} (C=O band, overtone C=O str),
 219 2952 cm^{-1} (C-H str), 1716 cm^{-1} (ketone), 1699 cm^{-1} (ketone),
 220 1653 cm^{-1} (C=C str), 1558 cm^{-1} (N=N, C=O), 1540 cm^{-1} (N=N,
 221 aliphatic N-O stronger), 1507 cm^{-1} (C-Cl, C-N, C-C, C-O R-
 222 NO_2 group strong intensity), 1489 cm^{-1} (C=C aromatic), 1457
 223 cm^{-1} (carbonate ion), 1378 cm^{-1} (CH_2 , CH_3 bend sp^3), 1233
 224 cm^{-1} (C-O-C strong, C-O-H), 1180 cm^{-1} (C-N str, O-H
 225 bending), 1038 cm^{-1} (C-O ester, ether, alco), 679 cm^{-1} (=C-H
 226 bending alkenes strong), 662 cm^{-1} (CO_2 medium, NH_2), 473
 227 cm^{-1} (CHO, C-N).

228 Infrared region 4000-650 cm^{-1} large number of bands. The
 229 possibility that two different compound shows same spectrum.
 230 The pure sample of TLC gives different infrared spectra must
 231 be different. If they passes supreme spectra there they represent
 232 same spectra. The region to the right of 1500-600 cm^{-1} is usually
 233 complex showing stretching and bending band. Hence,
 234 correlation of an individual bend with specific functional group
 235 of offends difficult. (*Lantana camara* active fraction after vacuum
 236 distillation followed by column chromatography shows bands
 237 1539, 1463, 1378, 1075, 797, 721 and 465 cm^{-1}).

Conclusion

Yearly assessed bioactive molecule/tripene from *Lantana* 239
camara leaves produced similar yield with negligible diffe- 240
 rence. Change in extraction time in relation to colour and R_f 241
 values, UV spectra, arrival and departure of FTIR exhibits 242
 richness of tripene that assists the diversified pharmacological 243
 application of active fraction/tripenes in processing. 244

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