

1 Development of Phyto-cosmetic Herbal Sindoor from *Rosa indica* L. and its 2 Characterization

3 ABSTRACT

4 **Aim:** The objective of this research study was to develop a liquid herbal sindoor containing flower
5 petal extract of *Rosa indica* L., and to evaluate the stability, antioxidant and antibacterial activities of
6 the developed formulation. **Place and duration of study:** The formulation was prepared in the
7 phytochemistry division of CSIR- National Botanical Research Institute. **The duration** of the study
8 period **was 90 days.**

9 **Methodology:** The crude ethanolic extract of *R. indica* flowers was incorporated into a base for the
10 preparation of liquid herbal sindoor. The antioxidant activity of this herbal product was evaluated
11 according to the DPPH radical scavenging method, and the stability study of the formulation was
12 evaluated for 90 days. The antibacterial activity was performed on those strains of bacteria, which are
13 responsible for several skin infections using Disc Diffusion Assay. The HPLC- DAD analysis
14 identified the anthocyanins cyanidin 3,5-diglucoside and pelargonidin 3,5-diglucoside in *R. indica*,
15 flower extract. **Results and Discussions:** The formulation showed antioxidant activity and its stability
16 was assessed in terms of pH values and color stability of the flower extract. It was observed that both
17 these properties did not show any significant changes. **Conclusion:** The prepared herbal liquid
18 sindoor was found safe to use having a potential to substitute the toxic lead tetraoxide (Pb_3O_4)
19 containing sindoor available in the market.

20 **Key words-** *Rosa indica*; flower extract; formulation; antioxidant; antibacterial; pH value and color stability;
21 lead tetraoxide

22

23 INTRODUCTION

24 **The use of phyto-cosmetics or herbal cosmetics has increased in personal care products.** There is a
25 high demand for cosmetics, having additional raw plant-derived ingredients as they are safe, non-
26 toxic, and environment friendly. On the other hand, **the overuse** of products made of synthetic
27 materials has put human health at risk and led to a number of health disorders. Sindoor is a traditional
28 vermilion red or orange-red coloured cosmetic powder, usually worn by married women along the
29 part of their **hairline in the Indian subcontinent.** Sindoor is a crucial cosmetics item in India for
30 married ladies, worship, and other uses. Long-term usage of synthetic sindoor containing lead
31 tetraoxide (Pb_3O_4) has shown symptoms of skin cancer, edema, hair loss, and greying of the hair **(1).**
32 Lead is one of the neurotoxins that has been the subject of the most recent and intensive research.
33 Over time, it accumulates like a typical poison (2). **In light of the situation mentioned above,** a process
34 technique has been developed to produce an alternative, safe, non- toxic, ecofriendly, natural dye
35 based Sindoor. The process provides an opportunity for the preparation of red color liquid herbal
36 Sindoor using ***Rosa indica*** as a raw material.

37 *Rosa indica* L. is a perennial flower shrub under the Rosaceae family, which contains herbs, shrubs or
38 trees that are thorny, rhizomatous or climbing (3). Rose is used in a wide range of ways. It is
39 traditionally been used in folk medicines, particularly in the field of skincare. It is appropriate for all
40 skin types, although it is particularly beneficial for dry, sensitive, or aging skin. It helps to reduce the
41 redness brought on by swollen capillaries since it has a tonic and astringent impact on the capillaries
42 right below the skin's surface (4). There are many important groups of compounds of rose that play an
43 important role in the antibacterial activity of rose. Among them, the groups of compounds with
44 antibacterial activity are flavonoids, terpenes, anthocyanins. Flavonoids are classified as natural plant
45 compounds, generally secondary metabolites with different phenolic structures (5). Rose flower is one
46 of the best sources for antibacterial activity against microorganisms. Strong antibacterial activity
47 against test pathogens such as *E. coli*, *P. aeruginosa*, and *S. aureus* is shown by ethanolic extracts of
48 rose petals, leaves, and stems (6). In addition to their antibacterial and antifungal properties,
49 flavonoids are also responsible for the aroma and color of flowers (5).

50 Due to the presence of antibacterial and antioxidant activities *R. indica* extract may be useful for the
51 synthesis of herbal drugs in the future (7). Natural sources with high anthocyanin contents are
52 becoming more and more popular in the food and medical industries for making dietary supplements
53 for therapeutic and nutritional purposes (8).

54 To find safe cosmetic alternatives in place of toxic traditional products, this work aims to develop an
55 antioxidant formulation containing ethanolic extract of the petals of *R. indica* and, evaluate its stability
56 and antibacterial activity.

57

58 MATERIAL AND METHODS

59 Plant material and preparation of crude ethanolic extract

60 Flowers of *R. indica* were collected from the temples of Lucknow, Uttar Pradesh, India, in the month
61 of July, 2021. The raw material was authenticated in the Herbarium of CSIR- NBRI, Lucknow
62 (LWG). Petals were separated from the flowers and weighed. The petals were macerated with
63 ethanol(100%) for 24 hours. The ethanolic extract obtained was filtered and concentrated under
64 reduced pressure on rotary evaporator.

65 Extract yield (%) = $(W_1/W_2) \times 100$

66 Where, W_1 is the weight of extracted plant residues in grams, and W_2 is the weight of rose petals in
67 grams. The yield of the extract obtained was 15.6 %.

68

69 Anthocyanin characterization using HPLC- DAD analysis

70 HPLC analysis of ethanolic extract of *R. indica* was performed on Shimadzu prominence system
71 equipped with an autosampler (SIL-20 AC), LC Solution 1.0 software and DAD detector. The plant
72 extract was analyzed in triplicate at the concentration of 1mg/ml as described by Huihua Wan et.al (9)
73 under the following conditions: stationary phase: C_{18} column (dimensions 250mm x 4.6mm and

74 particle size of 5 μm); mobile phase: solution A: water + 0.5% (v/v) formic acid and solution B:
 75 Acetonitrile (ACN), considering the elution gradient as shown in **Table I**. The injection volume and
 76 the flow rate were set at 10 μl and 0.5ml/min respectively at the constant temperature of 25°C. Before,
 77 analysis the solutions were degassed and filtered through 0.22 μm filter membrane. The wavelength
 78 was set at 254 nm for analysis. All the reagents used for chromatographic analysis were of HPLC
 79 grade.

80

81 **Table 1:** Gradient elution method for HPLC analysis of anthocyanin in *Rosa indica*

82

| Time (min) | Solution A: water+ 0.5% (v/v) formic acid (% v/v) | Solution B: Acetonitrile (ACN) (% v/v) |
|-------------------|--|---|
| 0 | 95 | 5 |
| 5 | 90 | 10 |
| 30 | 81 | 19 |
| 50 | 60 | 40 |
| 50.01- 60 | 95 | 5 |

83

84 **Development of formulation of liquid herbal sindoor**

85 For the development of the formulation the compatibility between the components and the flower
 86 extract has to be studied. The components used were as follows: bees wax, sunflower wax, castor oil,
 87 coconut oil and flower extract. These components were mixed together in petridish, kept on a hot
 88 plate at the temperature of 50°C. Then the herbal formulation was subjected to stirring on magnetic
 89 stirrer for 7 to 8 hours at 45°C.

90

Table.2 Ingredients with their composition (in percentage) in formulation

| S.No. | Ingredients | Composition (in %) |
|--------------|--------------------|---------------------------|
| 1 | Bees wax | 25 |
| 2 | Sunflower wax | 20 |
| 3 | Castor oil | 05 |
| 4 | Coconut oil | 10 |
| 5 | Flower extract | 40 |

91

92 **Stability studies**

93 The stability of the colour of the formulation was analyzed on 0, 30, 60 and 90 days using a
 94 colorimeter (High- Quality Spectrophotometer NS800). Using a colorimeter, the colours are expressed
 95 numerically in accordance with international standards. Anyone may comprehend what colour is

96 being conveyed in this way. In order to more precisely note the changes in colour, the data from the
97 colorimeter could enable the identification of different samples by any single quality (lightness,
98 chroma, or hue). The informational value of the data is substantially enhanced by the capacity to
99 appropriately define different colours (10). The following parameters—L*, a*b*, C*, and h°—were
100 monitored in order to accurately assess the formulation's colour.

101 A (previously calibrated) digital pH meter of LABMAN (Model no: LMPH- 12) was used for
102 measuring pH of the sample. Before measurement, the samples were diluted to 10% (w/v) in distilled
103 water (20).

104 **Antioxidant activity**

105 The DPPH radical-scavenging activity of the samples was measured by the method described by Blois
106 (11) with a slight modification using gallic acid and ascorbic acid as reference. The formulation
107 prepared from ethanolic extract of petals of *R. indica* was analyzed for radical-scavenging activities.
108 Briefly, the stock solution of formulation having concentration 1mg/ml was prepared. Various
109 concentrations (20µl, 40µl, 60µl, 80µl, 100µl) of formulation were prepared by adding both methanol
110 (80µl, 60µl, 40µl, 20µl) and 1mL of a DPPH ethanolic solutions. The mixtures were vortexed and
111 incubated for 30 min at room temperature in the dark. The absorbance of mixtures was determined by
112 using a spectrophotometer at 517 nm. UV–Vis absorption spectra were measured on a Perkin Elmer
113 Lambda 35 UV/VIS spectrophotometer (WIN LAB V6 STD Software). The free radical scavenging
114 activity was expressed in percentage using the following formula:

115 DPPH radical scavenging activity(%) = (absorbance control - absorbance of sample/absorbance
116 control) x 100

117

118 **Antibacterial activity**

119 *Preparation of inoculums of microorganism for testing*

120 Three pathogens *Escherichia coli*, *Bacillus pumilis*, and *Staphylococcus aureus* were used as test
121 subjects to determine the antibacterial activity of rosepetals. These three pre-isolated bacterial cultures
122 were obtained from KGMU, Lucknow, Microbiology Division. The cultures were subcultured on NA
123 slants and stored at 4°C until needed. For testing of the sample, the inoculums were prepared from the
124 stock culture and subcultured into the nutrient broth (30ml) using a sterilized wire loop. The inoculate
125 was further incubated overnight at 37°C in a rotary shaker. These inoculates were further stored at
126 4°C until usage.

127 *Activity analysis (Disc Diffusion Assay)*

128 The antibacterial activity of formulation and extract was done by Disc Diffusion Assay (12). The
129 antibacterial activity of formulation prepared from *R. indica* petals, was evaluated against three
130 pathogenic bacterial strains by using agar well diffusion method under sterilized conditions. For this
131 three NA plates were prepared for formulation and petal extract. 400µl inoculum of each selected
132 bacterium was uniformly spread over agar plates with the help of sterilized glass spreader.

133 After five minutes three wells, approximately 7mm in diameter were bored with the help of borer. The
 134 equal volume (50 µl) of antibiotic Tetracycline (100 mg/ml), formulation and petal extract (200mg/ml)
 135 were poured into the wells. The plates were incubated at 37 °C for 24 hrs in incubator. Next day, the
 136 results were observed and the antibacterial potential was measured in terms of diameter of zone of
 137 inhibition.

138

139 RESULTS AND DISCUSSIONS

140 With the purpose of obtaining the crude ethanolic extract, the fresh petals of *R.indica* were subjected
 141 to maceration with ethanol. The final solution, after being concentrated on a rotary evaporator, was
 142 subjected to HPLC analysis thereafter, a formulation was developed for preparation of liquid herbal
 143 sindoor. The prepared formulation was evaluated for various physical parameters as described in table
 144 3.

145

Table 3: Evaluation of physical parameters of herbal formulation

146

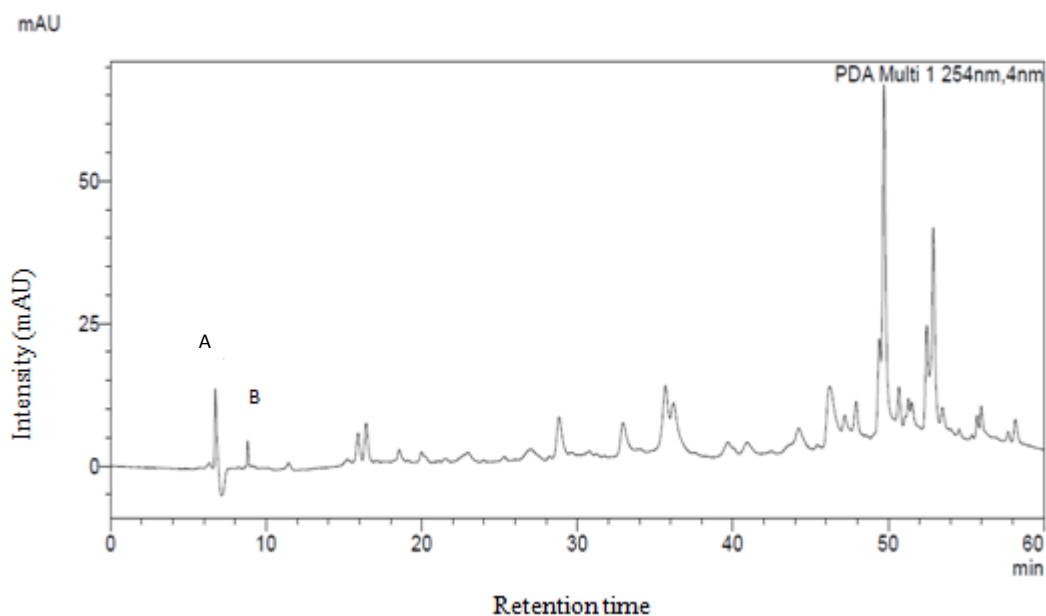
| S. No. | Parameters | Observation |
|--------|-------------------|--------------|
| 1 | Appearance | Thick liquid |
| 2 | Color | Dark red |
| 3 | Texture | Smooth |
| 4 | Odour | Pleasant |
| 5 | Water washability | Washable |

147

148 The analysis, of the flower extract of *R. indica* using high-performance liquid chromatography
 149 coupled to a DAD detector (HPLC–DAD), enabled the identification and quantification of cyanidin
 150 3,5-diglucoside and pelargonidin 3,5-diglucoside, the two analytical markers previously reported in
 151 this plant species (13). The retention time (Rt) of the two anthocyanins, 3,5-diglucoside and
 152 pelargonidin 3,5-diglucoside were 6.72 and 8.80 min. respectively. The analysis of the peak area
 153 percentage quantified the content of **these anthocyanins** in the crude ethanolic extract, the amount of
 154 3,5-diglucoside and pelargonidin 3,5-diglucoside was 0.0255 and 0.0047 mg/ml, respectively. **Figure**
 155 **1 presents the chromatographic profile of the crude ethanolic extract indicating the identified markers.**

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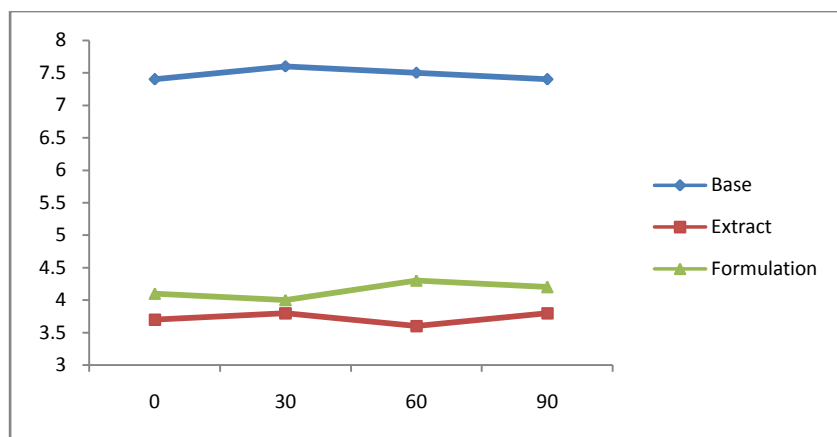
160 **Figure 1:** HPLC Chromatogram of ethanolic extract of *R. indica* at the wavelength of 254 nm. (A =
161 cyanidin 3,5-diglucoside, B = pelargonidin 3,5-diglucoside)
162

163 **On a fresh weight** basis the total anthocyanin content ranged from 0.24 to 578.10 mg/100g.

164 The environmental factors such as temperature and light strongly influence the anthocyanin content
165 of the rose petals (14). Lin et al. (2014) reported 0.025 to 18.69 mg/g of total content of anthocyanin
166 in rose petals. Anthocyanins give plants a variety of physiological health benefits and disease
167 preventive actions in addition to giving natural beautiful colors(15).

168 The pH analysis of the extract, base (castor oil, coconut oil, sunflower and bees wax) and formulation
169 was performed during the study period for 0, 30, 60 and 90 days and no significant variation was
170 observed .The observed pH values were around 4.2 showing biocompatibility with the skin, being
171 potentially non- irritating. **The pH gradient for the skin** surface ranges from 4 to 6 (16). The pH value
172 of the used base ranged from 7.4 to 7.6. Finally, the pH value of the formulation ranged from 4.0 to
173 4.3 over the period of study (90 days).

174
175



176
177

178 **Figure 2:** pH values of base, extract and formulation during the study period (0, 30, 60 and 90 days)

179

180 The color of the formulation was evaluated at the time interval of 0, 30, 60 and 90 days to observe any
181 significant change in color as described in table 4 . The color parameters such as L* (indicating
182 lightness), a*(redness), b*(yellowness), C*(chroma), h^o (hue) were measured using colorimeter. While
183 the parameter b* is positive for yellowish colors and negative for bluish hues, the parameter a* is
184 positive for reddish colors and negative for greenish colors. Each hue can be thought of as being
185 equivalent to a member of the greyscale that is in between black and white according to L*, an
186 approximation of brightness. Thus, each scale's L value represents the degree of lightness or darkness.
187 The delta values (ΔL , Δa , and Δb) reflect the degree of variation between a standard and sample for L,
188 a, and b respectively, and ΔE is the total color difference. L*, a*, and b* all have delta values i.e. ΔL ,
189 Δa , and Δb respectively, that can be either positive (+) or negative(-). The sign of the delta value
190 indicates whether the sample is redder or greener than the standard. Delta E (ΔE), is always positive
191 (17). The value of a* and ΔE were always found to be positive (+) in each measurement. No
192 significant change in color of the formulation was observed during the study period.

193 **Table 4:** The values of color parameters L* (indicating lightness), a*(redness), b*(yellowness),
194 C*(chroma), h^o(hue) measured using colorimeter

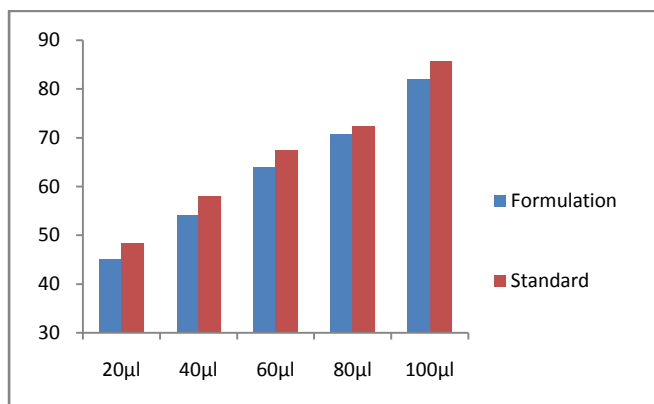
| ΔT^* | Standard | | | | | Formulation | | | | | Difference between standard and formulation | | | | | |
|--------------|----------|------|-------|------|----------------|-------------|------|-------|------|----------------|---|------------|------------|------------|------------|------------|
| | L* | a* | b* | C* | h ^o | L* | a* | b* | C* | h ^o | ΔL | Δa | Δb | ΔC | Δh | ΔE |
| 0 | 0.24 | 0.08 | 0.07 | 0.11 | 42.42 | 0.23 | 0.07 | -0.05 | 0.09 | 324.51 | -0.01 | -0.01 | -0.12 | -0.02 | -0.12 | 6.12 |
| 30 | 0.60 | 0.27 | 0.32 | 0.42 | 50.34 | 0.52 | 0.12 | 0.54 | 0.55 | 77.82 | -0.08 | -0.15 | 0.22 | 0.14 | 0.23 | 0.28 |
| 60 | 0.33 | 0.39 | -0.95 | 1.03 | 292.3 | 0.31 | 0.35 | -0.44 | 0.56 | 308.07 | -0.01 | -0.04 | 0.51 | -0.47 | 0.21 | 0.51 |
| 90 | 0.29 | 0.09 | -0.03 | 0.03 | 268.9 | 0.60 | 0.99 | -1.18 | 1.54 | 509.89 | 0.31 | 0.90 | -1.15 | 1.15 | 0.15 | 1.55 |

195 ΔT^* = Time interval in days

196 The antioxidant potential, of the formulation was tested according to the DDPH radical scavenging
197 assay. The concentration of the sample necessary to scavenge 50% of the DPPH free radicals, or the
198 IC₅₀ value, was computed from the graph. The term "IC₅₀" refers to the quantity or concentration of
199 extracts required to neutralise 50% of free radicals (8). Polyphenols present in the extract have a
200 major role, in the antioxidant activity. The antioxidant potential of the formulation according to DPPH

201 assay has an IC_{50} value of $27 \pm 05 \mu\text{g/ml}$ in case where gallic acid was used as reference standard as
 202 described in figure 3.

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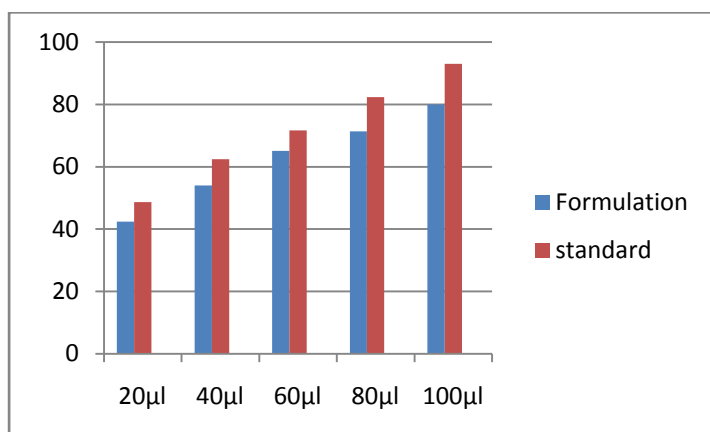


205
 206

207 **Figure 3:** Analysis of Free Radical Scavenging Activity of herbal formulation
 208 at different concentration using Gallic acid as standard

209

210 The antioxidant activity of formulation was also determined using ascorbic acid as reference
 211 standard (figure 4) and the IC_{50} value was $32.82 \pm 03 \mu\text{g/ml}$.



212

213

214 **Figure 4:** Analysis of Free Radical Scavenging Activity of herbal formulation at different
 215 concentration using Ascorbic acid as standard

216

217 Results of the test for antibiotic susceptibility were expressed as the diameter of the inhibition zone (in
 218 mm) of ethanolic extract and formulation against each bacterial strain employed. The zone of
 219 inhibition for the standard antibiotic (Tetracycline) against each strain was also recorded. Data of
 220 antibacterial activity of ethanolic extract and formulation are demonstrated in Table 5:

221

222

223

224

225 **Table 5:** Antibacterial susceptibility assay of *R. indica* petals extracts and herbal formulation

| Tested Bacteria | Zone of Inhibition (in mm) | | |
|-------------------|----------------------------|--------------------|--------------|
| | Ethanollic Extract | Herbal formulation | Tetracycline |
| <i>E. coli</i> | 22.0 ± 1.0 | 23.0 ± 1.0 | 31.0 ± 2.0 |
| <i>S. aureus</i> | 18 ± 1.0 | 17 ± 0.0 | 30.0 ± 1.0 |
| <i>B. pumilis</i> | 20 ± 0.0 | 21.0 ± 1.0 | 34.0 ± 1.0 |

226

227 **CONCLUSION**

228 In this research efforts were made to illustrate the compatibility between the base used and the crude
 229 ethanolic extract of petals of *R. indica*. The stability study of the formulation at room temperature and
 230 pressure showed no significant change in the pH and colour of the sample. The formulation also
 231 showed antioxidant activities carried out by DPPH radical scavenging method using gallic acid and
 232 ascorbic acid as standard . The anti-bacterial activity on various strains of bacteria (*S. aureus*, *B.*
 233 *pumilis*, and *E. coli*) indicated the efficacy of the formulation against such bacterial strains and
 234 demonstrated the antibacterial stability of the formulation. All these studies confirmed the anti-
 235 irritant, anti- allergic and antibacterial properties of the prepared formulation (liquid herbal sindoor).
 236 Thus this herbal product with no side effects may be used as substitute of lead tetraoxide (Pb₃O₄)
 237 containing sindoor available in the market. . However, in future, further studies are still required to
 238 develop such herbal based products using *R.indica* and to discover their beneficial health effects.

239

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247 **REFERENCES**

- 248 (1) Shivam Samariya, Sumeet Dwivedi, Shweta Patil, Debadash Panigrahi and Hemant Joshi:
 249 Formulation and Evaluation of Herbal Sindoor Using Different Natural/Herbal Ingredients.
 250 *International Journal of Pharmacy Teaching &Practices*.2013; 3: 752-754
 251 (2) Durgesh Wadhwa, and Gopal Arora: Identification of Lead from Sindur Samples. *International*
 252 *Journal of Innovative Research in Engineering & Management (IJIREM)*.2022; 9: 357-360
 253 (3) W. L. Crepet, K. C. Nixon, and M. A. Gandolfo: Fossil evidence and phylogeny: The age of
 254 major angiosperm clades based on mesofossil and macrofossil evidence from cretaceous deposits.
 255 *American Journal of Botany*.2004; 91: 1666–1682

- 256 (4) Thakare Priya, Ashok Rao, Deshbhratar Kiran, Suryawanshi M.N: A brief review on therapeutic
257 effects of – “ormamental plant” rose. *International Journal of Ayurveda and Pharma Research*.
258 2017;5: 46-52
- 259 (5) Yusra Safdar and Taqdees Malik: Antibacterial activity of the rose extract. *Open Access Journal of*
260 *Complementary and Alternative Medicine*. 2020; 2: 194-201
- 261 (6) R P Mishra, Mohammad Arshad and Abdul Sami. Antibacterial Properties of *Rosa indica*
262 (L.)Stem, Leaves and Flowers. *Journal of Pharmaceutical and Biomedical Sciences (JPBMS)*.2011;
263 12: 1-3
- 264 (7)Prerna , Prince sharma ,Sanjana Chaudhary, Dr. JyotiTyag : A review on antimicrobial and
265 antioxidant activity of *rosa damascene* against different species of microbes. *International Journal of*
266 *Advance Research and Innovative Ideas in Education*.2021; 7: 3555-3568
- 267 (8) Jin Hwan Lee, Hyeon-Jin Lee, Myoung-Gun Choung: *Food Chemistry*. 2011; 129: 272-278
- 268 (9) Huihua Wan, Chao Yu, Yu Han, Xuelian Guo, Sagheer Ahmad, Aoying Tang, Jia Wang, Tangren
269 Cheng, Huitang Pan, Qixiang Zhang. Flavonols and carotenoids in yellow petals of rose cultivar
270 (*Rosa* 'Sun City'): A possible rich source of bioactive compounds. *Journal of Agricultural and Food*
271 *Chemistry*.2018;66: 4171–4181
- 272 (10) Mohammed Azhar Bintory ,Seetharamu GK, Munikrishnappa PM, Ramegowda GK and
273 Basavaraj G. Evaluation of the Colour of Dried Dutch Rose Flowers Using a Colorimeter. *Journal of*
274 *Horticulture*.2015; 2
- 275 (11)Blois, M. A. Antioxidant determination by the use of a stable free radical.*Nature*.1958; 181:
276 1199–1200
- 277 (12) Amit Alexander Charan and Prerak Gupta.Comparative Analysis of Antibacterial, Antioxidant
278 and Photosynthetic activity of *Azadirachta indica*, *Rosa indica* and *Moringa oliefera* cultivars.
279 *International Journal of Current Research*.2013; 5: 556-561
- 280 (13)Huihua Wan, Chao Yu, Yu Han, Xuelian Guo, Le Luo, Huitang Pan, Tangchun Zheng, Jia Wang,
281 Tangren Cheng and Qixiang Zhang: Determination of Flavonoids and Carotenoids and Their
282 Contributions to Various Colors of Rose Cultivars (*Rosa* spp.). *Frontiers in Plant Science*.2019;10:
283 Article 123
- 284 (14) Poonam Kumari, D V S Raju, K V Prasad, Kanwar Pal Singh, Supradip Saha, Ajay Arora and
285 Firoz Hossain. Quantification and correlation of anthocyanin pigments and their antioxidant activities
286 in rose (*Rosa hybrida*) varieties.*Indian Journal of Agricultural Sciences*.2017; 87: 76-82
- 287 (15) Lin L, Hyeonmi H, JeehyeS, Younghwa K and Heon-Sang J L. Antioxidant activities of
288 methanolic extracts from four different rose cultivars. *Journal of Food and Nutrition Research*.2014;
289 2: 69–73
- 290 (16) Serup, J.; Jemec, G.B.E.; Grove, G.L. (Eds.) Handbook of Non-Invasive Methods and the Skin,
291 2nd ed.; *Taylor & Francis*: Boca Raton, FL, USA, 2006; 1056p.

- 292 (17) Deepali Singhee and Adrija Sarkar. Colorimetric Measurement and Functional Analysis of
293 Selective Natural Colorants Applicable for Food and Textile Products. *IntechOpen*. 2022: 1-33
- 294 (18) Akshada Kakade, Divya Yadav, Pranoti Bhonjale, Pallavi Velapure, Akshay Vedpathak and
295 Prajyoti Hiware. A New Approach of Formulation and Evaluation of Herbal Sindoor
296 Powder and Stick from Herbal Ingredients. *European Journal of Biomedical and*
297 *Pharmaceutical Sciences*. 2016; 3: 251-253
- 298 (19) Sheema Bai, Leena Seasotiya, Anupma Malik, Pooja Bharti and Sunita Dalal: Bioactive
299 compounds and pharmacological potential of *Rosa indica* L. and *Psidium guajava* L. methanol extracts
300 as antiurease and anticollagenase agents. *Der Pharmacia Lettre*. 2015;7: 179-184
- 301 (20) Rafaela Santos de Melo , Silvio Alan Gonçalves Bomfim Reis, Amanda Leite Guimarães ,
302 Naiane Darklei dos Santos Silva, Joao Miguel Rocha, Nouredine El Aouad, and Jackson Roberto
303 Guedes da Silva Almeida : Phytocosmetic Emulsion Containing Extract of *Morus nigra* L. (Moraceae):
304 Development, Stability Study, Antioxidant and Antibacterial Activities. *Cosmetics*. 2022; 9: 1-12