



Lawsonia inermis Linn: A Breakthrough in Cosmeceuticals

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Abstract: Herbal cosmetics are formulated using different cosmetic ingredients to form the base in which one or more herbal ingredients are used to cure various skin ailments. The name suggests that herbal cosmetics are natural and free from all the harmful synthetic chemicals that otherwise may be toxic to the skin. Compared to other beauty products, natural cosmetics are safe to use. Cosmeceuticals are cosmetic-pharmaceutical hybrid products intended to improve the health and beauty of the skin by providing a specific result. There are numerous herbs available naturally that have different uses in cosmetic preparations for skincare, hair care, and as antioxidants. The current study included a review and authentication of the various aspects of the plant *Lawsonia inermis*. *L. inermis*, commonly known as henna, has been cultivated for thousands of years for its leaves, which contain a natural dye molecule called lawsone that is commonly used to dye hair, skin, and fabrics. Henna has a long history of use in traditional medicine, where it has been used to treat a variety of ailments. In addition to its medicinal and cosmetic uses, henna has cultural and religious significance in many parts of the world and is commonly used to decorate the skin for weddings, festivals, and other special occasions. Because of these properties, the *L. inermis* plant can be used as a medicine against a wide range of pathogenic organisms and diseases. This review covers the phytochemistry, pharmacological properties, and traditional uses of the plant.

Introduction

The most recent fashion and beauty trend are herbal cosmetics. Since natural products provide the body with nutrients, improve health, and provide satisfaction because they are free from synthetic chemicals and have comparatively fewer side effects than synthetic cosmetics, most women today choose natural products over chemicals for their care to enhance their beauty (1).

The word "cosmetic" originates from the Greek word "kosmos" which means power, arrangement, and skill in decoration. As cosmetics evolved throughout human history, a consistent narrative about their beginnings emerged. These topical corrective pharmaceutical combinations are intended to enhance beauty by using their constituents' appropriate characteristics needed for skin and hair care (2, 3).

The area of cosmetics market is expanding the fastest. Cosmeceuticals are cosmetic-pharmaceutical products that target a specific issue, such as acne control, anti-aging properties, or sun protection, to enhance the health and beauty of the skin. According to the theory put forth by Dr. Albert Klingman, "Cosmeceuticals are topical agents that are distributed across a broad spectrum of materials, lying somewhere between pure cosmetics (lipstick and rouge) and pure drugs" (antibiotics, corticosteroids) (4).

Methods

The search engines PubMed, Scopus, and Google Scholar were used to conduct a thorough literature review using the keywords *Lawsonia inermis* with cosmeceuticals, henna, pharmacological, antioxidant, and antifungal up until April 2023. Additionally, the predicted objective data was assembled.

The regulatory status of cosmeceuticals

Cosmeceuticals - cosmetics or drugs?

A product's intended use determines whether it is considered a cosmetic or a drug for legal purposes. The line between a cosmetic product and a drug is not clearly defined under the current concept, and different rules and laws are applied to various product categories. According to The Drugs and Cosmetics Act, of 1940, drugs are defined as "All medicines for internal or external use of human beings or animals and all substances intended to be used for; or in the diagnosis, treatment, mitigation or prevention of any disease or disorder in humans or animals" (5). And cosmetics are defined as "Any article intended to be rubbed, poured, sprinkled or sprayed on or introduced into or applied to any part of the human body for cleansing, beautifying, promoting attractiveness or altering the appearance and includes any article intended for use as a component of cosmetic" (6).

Cosmetics and drugs

Some products fall under both the cosmetics and drug definitions. When a product has multiple intended uses, this might emerge. Shampoo, for instance, is a cosmetic because its main purpose is to clean hair. As a treatment for dandruff, an anti-dandruff shampoo considers a drug. Among the cosmetic/drug combinations are moisturizers with sun protection

claims, deodorants that are antiperspirants, and toothpaste that contains fluoride (7). The Food and Drug Administration (FDA) review and approval process is required for claims made about drugs but not for claims made about cosmetics. Even though there isn't a specific legal category for cosmeceuticals, the term has come to be used to describe products that fall somewhere between cosmetics and pharmaceuticals (8, 9).

The term itself is not recognized by the Federal Food, Drug, and Cosmetics Act. Consumers frequently find it challenging to verify "claims" made about the actions or efficacy of cosmeceuticals in the absence of the FDA or another reputable regulatory body's approval. Some nations have product categories that fall between the two classifications of drugs and cosmetics, such as 'controlled cosmetics' in Thailand, 'Quasi-drugs' in Japan, and 'cosmetic-type drugs' in Hong Kong. The laws governing cosmeceuticals are not uniform across the USA, Europe, Asia, and other nations (10).

Herbs Used in Cosmetics/Cosmeceuticals

Numerous herbs are found in nature and have a variety of uses, including antioxidants, fragrances, and preparations for skin and hair care. Table 1 contains some crucial examples.

Table 1. Various herbs are used in different herbal formulations.

Action	Ingredients	References
Skincare	Coconut Oil, Sunflower Oil, Jojoba Oil, Olive Oil, Aloe	(11-14)
Anti-aging	<i>Rhodiola rosea</i> , Carrot, Ginkgo, Neem	(15-19)
Skin Protection	Green Tea, Calendula, Turmeric, Aloe, Jojoba Oil, Carrot Seed Oil, Wheat Germ Oil, Witch Hazel, Arnica	(20)
Dandruff Treatment	Henna, Shikakai	(21, 22)
Haircare	Amla, Eucalyptus Oil, Snake Grass, Coconut Oil, Shikakai, Reetha, Neem, Sandalwood Oil, Brahmi, Castor Oil, Babchi, Carrot Seed Oil, Witch Hazel Extract, Almond Oil, Lemmon Grass Oil	(23-26)
Antioxidant	Tamarind, Vitamin C, Vitamin E, Beta-Carotene, Retinol, Ginkgo, Green Tea, Flavonoids, Turmeric Oil, Glutathione, Coenzyme Q 10,	(27-31)
Oral Hygiene Product	Nutmeg, Clove oil, Neem oil, Miswak Extract, Eucalyptus Oil, Myrrh Extract, Tea Tree Oil, Piper Betel Extract, Tulsi Oil, Peppermint Oil, Babhul, Vajradanti, Majuphal, Wintergreen oil	(32-36)
Natural Colorants	Red poppy petals, Annatto, Red beetroot, Blue Tansy, Butterfly Pea, Flavoxanthin, Beta carotene pigment, Chamomile, saffron, curcumin, carthamin, red sandalwood, henna	(37-40)
Fixed oils	Coconut oil, almond oil, olive oil, sesame oil, castor oil,	(41-44)
Waxes	Beewax, carnauba wax,	(45-47)
Gums	Agar, tragacanth	(48, 49)
Perfumes	Rose oil, lavender oil, immortelle oil, German chamomile oil, neroli oil, rosemary essential oil, tea tree oil,	(50-53)
Bleaching agent	Licorice extract, kojic acid, <i>Phyllanthus emblica</i> , Giga white, willow bark extract	(54-60)



Figure 1. A picture of *L. inermis* (henna plant).

Lawsonia inermis Linn.

Until the 19th century, natural plant dyes served as the foundation for the cosmetic and food industries (Figure 1). In addition to their traditional medicinal uses, many plants in Iraq were used as natural dyes (61). Henna, also known as *L. inermis* Linn (Family: Lythraceae), is primarily found in subtropical and tropical regions and is used all over the world. It has been used as a dye for cosmetic purposes for more than 9,000 years. *L. inermis* contain phenolic, flavonoids, saponins, proteins, alkaloids, terpenoids, quinones, coumarins, xanthenes, fat, resin, and tannins, according to phytochemical analysis. Additionally, 2-hydroxy-1,4 naphthoquinone is present (lawsone). Numerous alkaloids, naphthoquinone derivatives, phenolics, and flavonoids were discovered in *L. inermis*' various parts.

The results of the pharmacological studies indicated that *L. inermis* exhibited a wide range of

pharmacological effects, like an antibacterial, antifungal, antiparasitic, molluscicidal, antioxidant, hepatoprotective, central nervous, analgesic, anti-inflammatory, antipyretic, wound and burn healing, immunomodulatory, antidiabetic hypolipidemic, antiulcer, antidiarrhoeal. The pharmacological properties and chemical properties of *L. inermis* will be highlighted in the current review (62).

Plant Profile

Synonyms

Alcanna spinosa, *Casearia multiflora*, *Lawsonia alba*, *Lawsonia speciosa*, *Lawsonia spinosa*, and *Lawsonia Rotantha combretoides* (63). Table 2 contains common/popular names of henna plants in different languages.

Historic Perspective

Issac Lawsone, a Scottish army doctor in the 18th century and a close friend of Carl Linnaeus, inspired the Latin name "Lawsonia" for the plant. The plant is commonly used in a wide range of religious and ritualistic ceremonies of the Hindu and Muslim communities and is a symbol of auspiciousness, prosperity, and happiness in South Asian nations like India, Pakistan, Iran, and the United Arab Emirates. Muslim men dyed their hair and beard with henna because they believed it to be a sunnah or an honorable practice of the Prophet Muhammad. Muslim women in the Middle East were urged to use henna to color their nails to show off their femininity and distinguish their hands from men (64).

Table 2. Common names of henna plants in different language(s).

Languages	Name
English	Henna, Samphire, Cypress shrub
Hindi & Urdu	Mehendi, Mehndi, Hinna
Sanskrit	Rangani, Mendika, Madayanti, Timir
Tamil	Marithondi, Marithonali, Alvanam, Aivani
Gujarati	Medi, mendi
Bengali	Mendi, Mehadi
Turkish	Kenaag
German	Agyptische
French	Alcana d' orient, Henne
Malayalam	Mailanchi
Italian	Enne, Cipro
Kannada	Mayilanchi
Telugu	Gorintaku, Kormmi
Greek	Kypros
Marathi	Mendhi

Source: <http://www.health-fundas/henna...synonyms/42060>. The Ayurvedic Pharmacopoeia of India. Part-I, vol. IV, pp. 64.

Historical treasures and traditional use of henna show that women have long painted their hands, feet, nails, and hair with henna. The earliest piece of art depicting actual living women with henna stains on their hands, feet, and nails can be found in "The House of the Ladies," Room 1, East Section, North Wall in Akrotiri. Both women in this wall painting have henna stains on their fingernails and soles (65, 66). Bronze Age texts attest to women's ceremonial henna use. These confirm the usage of henna in the eastern Mediterranean before the Santorini eruption, which occurred around 1627 BCE (67).

Habitat

L. inermis is primarily grown for cosmetics and traditional medicine throughout the world, but it is native to Africa and Asia. It was distributed in Africa: Egypt, Ethiopia, Somalia, Sudan, Zaire, Niger, Benin, Burkina Faso, Cote D'Ivoire, Gambia, Ghana, Guinea, Guinea-Bissau, Liberia, Mali, Nigeria, Senegal, Sierra Leone, Togo, South Africa, Comoros, Seychelles; Asia: India, Pakistan, Sri Lanka. It is widely cultivated in tropical regions of the world, North and East Africa, the Arabian Peninsula, the Southern areas of the Middle East, and South Asia (68, 69).

Taxonomic Classification

Kingdom: Plantae, Subkingdom: Viridiplantae, Infrakingdom: Streptophyta, Superdivision: Embryophyta, Division: Tracheophyta, Subdivision: Spermatophytina, Class: Magnoliopsida, Superorder: Rosanae, Order: Myrtales, Family: Lythraceae, Genus: *Lawsonia*, Species: *Lawsonia inermis* (70)

Botanical Description

It grows to a height of 2.4-5 m and is a densely branched, globous, deciduous, occasionally spinescent shrub or small tree with greyish-brown bark. It is grown as a commercial crop for its dye in some Indian states and as a hedge plant throughout the country (71). The leaves are 1.3 -3.2 by 0.6-1.6 cm, elliptic or broadly lanceolate, acute, or obtuse, frequently micronucleate, and have a tapering base. Petioles are very short. Flowers are numerous, 1.3 cm across, fragrant, white, or rose-colored, and borne in long, slender pedicels in large terminal pyramidal cymes. The lobes of the calyx, which is 3-5 mm long and broadly campanulate, are suborbicular or sub reniform and undulate. Eight stamens are inserted into the calyx tube in pairs. 6 mm-diameter capsules that are slightly veined on the outside, supported by the persistent calyx, and have the style (72). The red, globose, pea-sized seed capsules contain numerous tiny, pyramid-shaped, brown-pitted seeds (71).

Traditional Uses

The leaves of *L. inermis* are a major source of cosmetic dye. For centuries, the Middle East, the Far East, and

Northern Africa have used henna leaves extensively as a dye for textiles, nails, hands, and hair. Additionally, henna was used to treat skin conditions, headaches, jaundice, amebiasis, and spleen enlargement (73, 74).

In the Charaka Samhita, this plant is mentioned as a remedy for jaundice, and epilepsy, and for coloring grey hair. It has been suggested as a treatment for malignant ulcers in Sushruta Samhita (71). The Ayurvedic Pharmacopoeia of India recommended using leaves for pruritus, dysuria, bleeding disorders, and other stubborn skin conditions (77). The leaf has a bitter taste and is used to treat spleen, vulnerary, headache, lumbago, bronchitis, boils, ophthalmia, syphilis, sores, and other conditions as well as to promote hair growth. An infusion of the flowers relieves headaches. Flowers are used to treat insomnia and as a refrigerant (76). The bark is prescribed for jaundice and spleen enlargement, as well as for calculous affections, leprosy, and stubborn skin conditions (72). Due to its antibacterial, antifungal, anti-amoebiasis, astringent, antihemorrhagic, hypotensive, and sedative effects, it is used as a medicinal plant (75).

It was used as a folk remedy in parts of south India to treat skin conditions and ringworm infections (78-80). Traditional Yemeni healers used the henna plant to treat burn wounds and bacterial infections (81, 82). Additionally, it was mentioned in the Ebers Papyrus medical texts during the Roman Empire in Rome (83). Medicine of the Prophet listed specific applications for henna as a treatment for tumors, migraines, leprosy, ulcers, smallpox, and other conditions. (84) Initial research using purified henna plant constituents and extracts identified antibacterial (85), antifungal (86), antioxidant, immunomodulatory (86, 90), protein glycation inhibition (87), anti-sickling (88, 91), macrophage-stimulating (92), hepatoprotective (93), analgesic, anti-inflammatory, antipyretic (94), anticomplementary (95), and cytotoxic activities (96) in various fractions. There were no allergic or cancerous side effects associated with the bactericidal and fungicidal actions, which were attributed to the tanning effect of the plant (97, 98).

Medically Useful Plant Parts

Different diseases are treated with the entire plant, including the stem, leaves, roots, fruits, inflorescence, rhizomes, bulbs, latex, seeds, and flowers (99).

Leaf

Henna leaf has an orange-red dye, and leaf paste or powder is frequently used to add designs to hands, feet, and nails. It can also be used to dye hair. Jaundice, skin conditions, sexually transmitted diseases, smallpox, and spermatorrhoea are treated with it (100-102).

Table 3. Ethnopharmacological uses of *L. inermis*.

Plant parts	Ethnopharmacological uses	References
Bark	Antifungal, Antibacterial, Anti-parasitic, Virucidal	(188)
Leaves	Bitter, Astringent, Acrid, Diuretic, Oedema, Expectorant, Anti-inflammatory, Depurative, Liver Tonic, Jaundice haematinic, Styptic, Febrifuge, Trichogenous, Wound healing, Strangury, Cough, Bronchitis, Soothing agent, Cephalalgia, Hemicranias, Lumbago, Rheumatalgia, Diarrhoea, Dysentery, Leprosy, Leucoderma, Scabies, Hepatopathy, Splenopathy, Anti-haemorrhagic, Hemoptysis, Fever, Ophthalmia, Amenorrhoea, to stop hair fall, Hair dye,	(189, 190)
Flowers	Cardiotonic, Refrigerant, Sedatives, Febrifuge, Liver tonic, Cephalalgia, Soothing agent, Amentia, Insomnia, Fever	(191, 192)
Roots	Bitter, Depurative, Diuretic, Emmenagogue, Abortifacient, Soothing agent, Leprosy, Amenorrhoea, Hair dye	(192)
Seeds	Antipyretic, Memory enhancer, Recurrent fevers, Insanity, Diarrhoea, Dysentery and to improve the gastric flow	(193)

Table 4. Traditional medicinal uses of henna plant parts in various countries.

Country/region	Medicinal use	Form applied	Administration	References
Africa	Induce abortion	Decoction of the whole plant	Oral	(197)
Cambodia	Diuretic, gonorrhoea, bronchitis	Roots	Oral	(198)
Egypt	Pain, skin infections, intestinal amoebiasis	Leaf paste, Decoction of leaf	Topical, Oral	(199)
Europe and America	Aromatherapy	Essential oils from aerial parts	Topical	(200)
India	Jaundice and other liver disorders, Itching, and other skin disorders	Decoction of stem bark, Leaf powder alone or mixed with other herbs, Leaves, Whole plant, seeds, leaf paste	Topical, Oral	(201, 110)
Iran	Recurrent unilateral headache	Leaf paste	Topical	(110)
Jordan	Hair loss, hair dyeing, and skin diseases	Leaf paste	Topical	(202)
Lebanon	Antirheumatic and antineuralgic	Leaves	Topical	(203)
Nigeria	Trypanosomiasis	Leaves, Whole plant	Oral	(204, 205)
Pakistan	Jaundice, as diuretic and blood purifier, skin diseases such as irritation of hands and feet, demulcent and resolvent, Hair, and skin problems	Decoction of Leaves, and leaf paste	Oral, Topical	(196)
Sudan	skin diseases	Whole plant	Topical	(194)
Saudi Arabia	Diabetic foot disorders	Whole plant	Topical	(195)

In the past, henna leaf paste was used to treat kidney lithiasis (104), diarrheal infusion, and skin inflammation (103) while leaf decoction was effective for cleaning and healing wounds (105). The tribes of the Andhra Pradesh region in India frequently used *L. inermis* leaves, *Hibiscus rosa-sinensis*, *Eclipta prostrata*, and *Abrus precatorius* seeds in equal quantities ground into a paste and soaked in sesame oil for 5 days as hair oil to prevent dandruff and hair fall (106). Tribes in Nigeria used leaves as a blood tonic (108), poliomyelitis treatment (107), and measles treatment (108). In Nigeria and Ivory Coast, *L.*

inermis leaves were also recommended as a treatment for African trypanosomiasis (109).

Flowers

Flowers have a strong fragrance, and they are used to make perfume. Flower infusion is a useful treatment for bruises. The flower decoction is referred to as an emmenagogue. Seeds have a deodorizing effect.

Seeds

For liver disorders and related issues, seeds in powdered form are an effective treatment. Dysentery

can be treated with powered seeds infused with real ghee (clarified butter) (100-102).

Bark

Burns and scalds are treated with a decoction of the bark. It is administered internally for a variety of ailments, including spleen enlargement, jaundice, and leprosy as an alternative to external treatments for persistent skin conditions (100-102).

Root

The root is regarded as a powerful treatment for herpes and gonorrhea. The root is astringent and can be pulped and applied topically to sore eyes. The pulped root can also be used to treat boils on children's heads. A decoction is consumed as a diuretic in Cambodia. The root is typically decocted as an effective abortifacient along with prepared indigo. The root is thought to be effective in the treatment of nervous disorders and hysteria (100-102). Ancient tribes of the Bhoja community of India frequently used a half-teaspoonful decoction made from *L. inermis* root taken orally twice daily for 10-15 days to treat jaundice (110). Table 3 contains Ethnopharmacological uses of *L. inermis*.

Modern-Day Use of Henna for Medicinal Purposes

L. inermis is used as traditional or folk medicine around the world to treat a variety of ailments that may seem unrelated to one another (Table 4). But in many instances, a pattern emerges that suggests similar uses by various cultural groups.

Cultivation and Production

Although henna trees can reach heights of six meters and can live for fifty years, they are frequently cut back to just one meter or less and their valuable leaves are harvested (111). Henna is grown in hedgerows because of its durability and tenacious roots, which protect the home garden from desert winds and soil erosion (112). Henna is primarily grown by family women because it doesn't require specialized tools or labor and can be found close to smallholder household compounds. When droughts kill off other crops, henna is a dependable source of income. A smallholder's henna tree is a valuable source of natural cures for minor illnesses. Henna can speed up the healing of wounds and treat ringworm and other fungal infections in children, adults, and livestock (113, 97). Henna's fungicidal, anti-inflammatory, and analgesic properties help nursing mothers with thrush-infected nipples (*Candida albicans*) (114, 115). The analgesic and antimicrobial properties are beneficial in household burn (116). For efficient dental self-care, cleaning the teeth, and preventing oral microbes, henna twigs are applied to the teeth (117). The belief that henna is "women's work" and that it is "old-fashioned" may have hindered efforts to develop, improve, and research the medicinal and economic potential of

henna. Currently, men predominate in the commercial henna export and processing industries. Even though henna is a product of North Africa and the Middle East, it is grown commercially in Pakistan, India, Egypt, Somalia, Sudan, Morocco, Iran, Yemen, and Somalia (67). Present commercial cultivation of henna employs annual or semi-annual pruning to harvest twigs and leaves (118). The cuttings are dried, and the leaves are stripped off and sorted, pulverized, and sieved for use as hair dye and skin stain.

Globalization of Henna as Body Art and Product

In the last fifteen years, henna body art has popularized, spreading globally from its areas of origin, and changing from being a traditional bridal and festival adornment to an exotic fashion accessory (Figure. 2A). Although henna's traditional uses for beauty and wellness have not gained as much popularity as body art and hair dye, researchers are increasingly looking into these practices. They are not doing this because henna is "blessed" or has the power to prevent diseases brought on by the "evil eye," but rather because it is a proven therapeutic for skin care. To prevent the oxidation of lawsone, the substance that gives henna its color, a mildly acidic liquid is combined with the powder before use (114). Henna is applied with a variety of implements that all bring the wet paste into contact with skin, hair, or nails. Without using heat or a mordant, the colorant penetrates, stains, and binds to the keratin in the skin, nails, and hair. After that, the skin is scraped or rinsed off the paste. About 1% of the lawsone migrates into the blood-bearing, living layers of the skin, where it eventually gets carried away by the urine. The majority of the lawsone stays in the skin's outermost, dead layers (119).

Lawsone undergoes oxidation after attaching to keratin (120). The stains are lighter in color and only last for a week or two in regions where the skin is thin and has a high lipid content. Although torso stains are light, they provide a complete sun-blocking effect that lasts beyond the duration of the stain and can provide protection for up to a year at the location of the stain; potentially an effective proactive treatment for melanoma (121). Henna is ideal for body inscriptions that need to be durable, but not permanent (122). Topical applications of henna are usually harmless (123). Henna body art is most seen on palms and soles, where the stains last three to four weeks. For brides and social gatherings, ornamental henna body art is typically limited to the hands, forearms, feet, and legs. Henna body art is currently associated with Hindus, Christians, Jews, Zoroastrians, Sikhs, Animists, and Muslims. Islamic culture plays the largest role in the widespread adoption of henna bridal traditions and

women's health. From the Atlantic coast of Africa to Malaysia and the Philippines, from South Africa to the farthest extent that trade routes carried henna into the Ottoman Empire, Persia, and Central Asia, Muslim women have used henna as part of their celebrations, fitness, and weddings (Figure. 2B). Although henna is now exported to Europe and North America for use as hair dye and body art, it was never native to the Western Hemisphere.

Processing and Preparation of Henna

One of the main sources of natural dye is *L. inermis*, a plant whose leaves give off an orange color. Before the development of synthetic dyes, henna was frequently used to dye clothing in addition to hair and skin. *L. inermis*, which is non-grazable and is a successful cash crop for farmers in arid and semi-arid tropical regions, is grown as a hedge to protect crops and orchards from castles (124). Since ancient times, henna, also known as mehndi, has been used to dye hair, skin, and nails. It has also come to have special significance in both Islamic and Hindu cultures (125).

Henna is traditionally made in India, Pakistan, Bangladesh, and the United Arab Emirates by chopping freshly collected leaves into a paste, which is then ground in a kitchen mixer or with an old stone pistol. Because naphthoquinone, glycosides are present in henna and effectively split off when applied to the skin with water, henna gives hair and skin color. No color is produced by isolated lawsone for the skin or hair (126).

Factors such as the method of cultivation, harvesting, processing, and storage affect the dying properties of henna. Therefore, for better quality and to stop color loss and deterioration, the right method of collection and processing is required. Although shade

drying can keep the leaf's green color, it is impractical for large-scale collection in December due to the low temperatures and rain. Maximum lawsone content (2.49%) is produced by using a herbal dryer for 3 hours at 50°C, whereas sun dry takes longer (72 hours) and produces minimum content (1.69%) (126). Therefore, simple, and inexpensive processing must be used to maintain post-harvest high quality and get excellent returns. It also becomes crucial to adhere to good agricultural and manufacturing practices for henna production with the utmost attention to preserve its dying properties. For best results, it is advised to collect henna leaf during the hot summer as lawsone content has been reported to be maximum during this season. Henna leaves must be ground with water to create a paste. Henna leaves cannot be used for cosmetic purposes (hair and skin dye) in their intact form. In Yazd, Iran, and many other places, henna leaves were ground into powder using henna millstones that were driven by powerful men, camels, and other animals. These henna millstones are made of limestone and have grooves cut into their surfaces to make grinding easier (127). Henna is produced commercially on a large scale for industrial uses in many nations, including India, Pakistan, Bangladesh, and Sudan. India is a significant producer of the highest quality henna among them and has contributed significantly to the development of all major henna-exporting and cultivating nations. The market has expanded significantly because of its domestic utility and symbolic significance in culture. Henna produced in India is used for a variety of other purposes, including domestic products, and is exported to the USA, UAE, Turkey, and the Middle East in about 31% of cases (128). According to a report by the Directorate General of Commercial Intelligence and Statistics (DGCIS), Government of India, henna also contributes to about 8% of the total amount of medicinal plants and herbs exported from India.

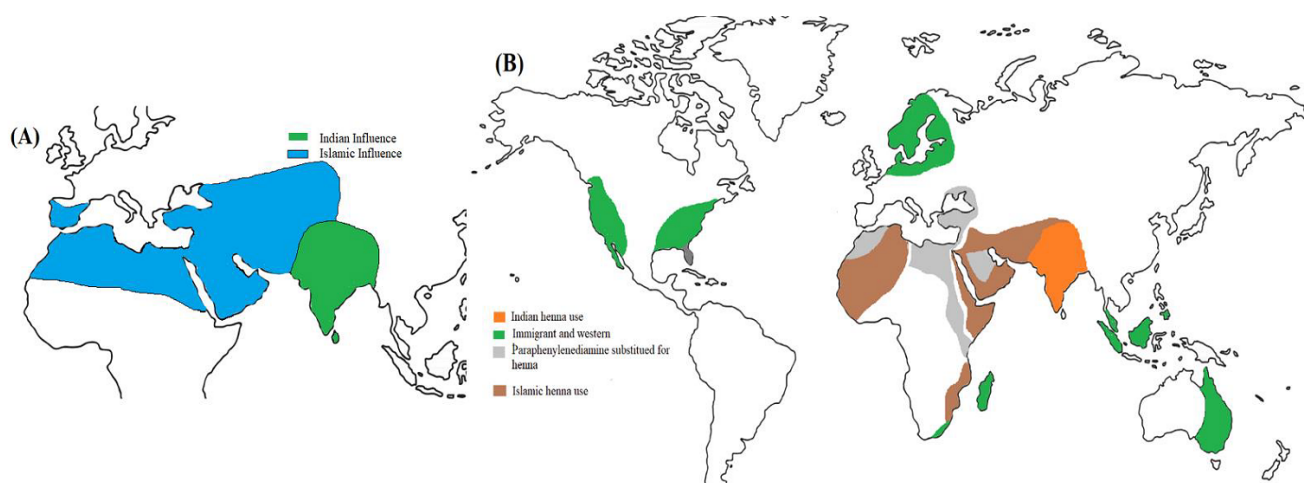


Figure 2. (A) Locations of artifacts with body markings consistent with henna or text mentioning henna between 700 and 1250 CE; (B) Regions where henna body art was practiced between 2000 and 2006.

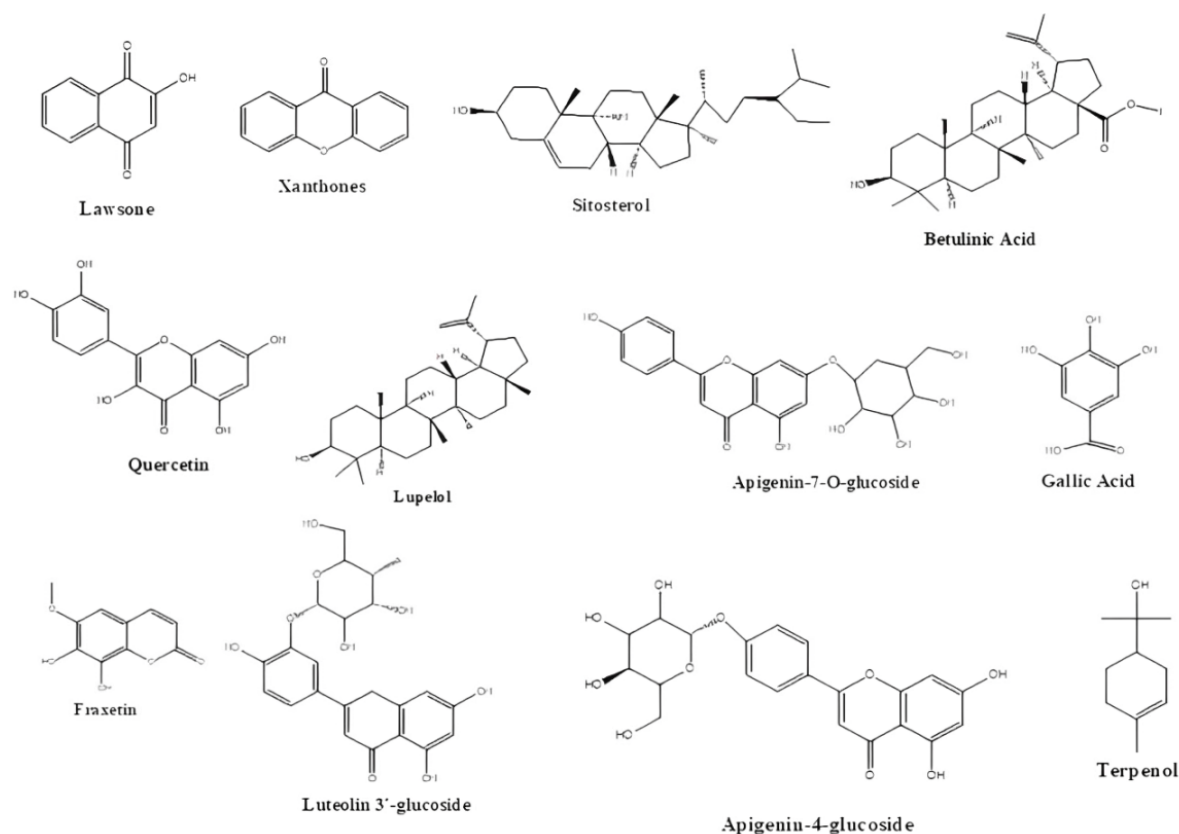


Figure 3. Chemical structures of a variety of compounds isolated from *L. inermis*.

Table 5. Some reported bioactive constituents of *L. inermis*.

Class of Compounds	Bioactive Constituents	Major Site of Occurrence	References
Napthoquinone	Lawsone (2-hydroxy 1,4-naphthoquinone) also called Hennotannic acid, 1, 3-dihydroxy naphthalene, 1, 4-naphthoquinone, 1, 2-dihydroxy-4-glucosyl naphthalene, Lawsoniaside (1,3,4-trihydroxynaphthalene 1,4-di- β -d-glucopyranoside), 5-hydroxy-2-methyl-1,4-naphthoquinone	Leaves, Stem, Bark	(109)
Polyphenolic components	Lalioside (2,3,4,6-tetrahydroxyacetoxy-2- β -d-glucopyranoside) Lawsoniaside B (3-(4-O- α -d-glucopyranosyl-3,5-dimethoxy) phenyl-2E-propanol), syringinoside, daphneside, daphnorin, agrimonolide 6-O- β -d-glucopyranoside, (+)-syringaresinol O- β -d-glucopyranoside, syringaresinol di-O- β -d-glucopyranoside, isoscutellarin, gallic acid	Stem, Bark, and Leaves	(206)
Terpenes and terpenoids	3 β , 30-dihydroxylup-20(29)-ene (hennadiol), (the 20S)-3 β , 30-dihydroxylupane, Lupeol, 30-nor-lupin-3- β -ol-20-one, Betulinic acid, Lawnermis acid (3 β -28 β -hydroxy-urs-12,20-diene-28-oic acid) and its methyl ester, -(Z)-2-hexenol, linalool, α -ionone, β -ionone, α -terpineol, terpinolene, δ -3-carene and γ -terpineol	Stem Bark, Seeds, and Essential Oil from Leaf and Flower	(206, 207)
Phytosterols and aliphatic compounds	Lawsaritol, Stigmasterol and β -sitosterol, 3-methyl-nonacosan-1-ol, n-tricontyl n-tridecanoate	Seeds, Stem, Root	(208)
Xanthones	Laxanthone I (1,3 dihydroxy-6,7 dimethoxy xanthone), Laxanthone II (1-hydroxy-3,6 diacetoxy-7-methoxyxanthone), Laxanthone III (1-hydroxy-6-acetoxy xanthone)	Whole Plant	(209)
Flavonoids	Quercetin, Apigenin, apigenin-7-glucoside, apigenin-4-glycoside, luteoline, luteolin-7-glucoside, luteolin-3-glucoside	Whole Plant, Leaves	(210, 211)
Miscellaneous chemical constituents	(+)-pinoselinol di-O- β -d-glucopyranoside Carbohydrate, proteins, fibres, and Trace metal-(Cu, Ni, Mo, V, Mn, Sr, Ba, Fe and Al) Minerals-Na ₂ O, CaO and K ₂ O	Whole Plant	(212)

Chemical Constituents

Aqueous extract *L. inermis* was subjected to a preliminary phytochemical analysis, which identified the following substances: 6% fat, 2-3% resin, 7-8% tannins, phenolic compounds, flavonoids, saponins, proteins, alkaloids, terpenoids, quinones, coumarins, and saponins (129-135). There was 2-hydroxy-1,4-naphthoquinone in *L. inermis* (lawsone). Lawsone concentrations in *L. inermis* flower, leaf, and branch extracts were determined by HPLC to be 116.7, 486.2, and 5.4 µg/g, respectively (136). Table 5 and Figure 3 contain some reported bioactive constituents of *L. inermis*.

Physicochemical Characteristics

Various physicochemical analyses of the leaf revealed that its total ash content was 14.60%, along with acid-insoluble ash of 4.50%, water-soluble ash of 3.0%, loss on drying of 4.5%, the alcohol-soluble extractive value of 3.8% w/w, and aqueous extractive value of 5.0% w/w (137).

Pharmacological Effects

Hypoglycaemic Activity

A study was done in 2008 by Syamsudin *et al.*, (2008) to see how ethanolic extracts of *L. inermis* leaves affected the glucose levels of rats with artificially induced diabetes (138). Extract from ethanol plant leaves significantly reduced glucose levels, demonstrating hypoglycemic activity. Additionally, they noted that this extract has hypolipidemic properties. Arayne *et al.*, (2007) showed a significant *in vitro* hypoglycaemic activity of *L. inermis* methanolic leaf extract (139).

Antimicrobial Activity

To assess the antimicrobial potential, *L. inermis* leaf samples were procured from the Dammar region, north of Sudan. Six human pathogenic fungi and four different types of bacteria were grown more slowly when crude extracts of water, methanol, and chloroform were obtained and bio-assayed *in vitro*. The three types of extracts' variations in bioactivity were examined. The water extract was superior despite the extreme activity variations. Anthraquinones, which are a major component of plant leaves and are frequently known to have antimicrobial activity, were discovered through phytochemical analyses (140, 141).

Antibacterial Activity

Twenty different plant species were used by Yemeni traditional healers to treat pathogenic illnesses. The antibacterial screening of various plant species uses both gram-positive and gram-negative bacteria. The ethyl acetate extract of *L. inermis* demonstrated the highest antibacterial activity of all the tested plant species (116). Quinone compounds from *L. inermis*

were examined for their antimicrobial properties *in vitro* by Dama *et al.*, (1999) (142). Lawsone was the subject of genotoxic studies by Kirkland and Marzin in 2003, who postulated that strain TA2637 of *Salmonella typhimurium* was more clearly mutagenic than strain TA98 and that the latter was a weaker bacterial mutagen (143). Overall, it appears that there is no genotoxic risk associated with *L. inermis* for consumers. The aqueous extract of leaves from *L. inermis* was also reported to have an antibacterial effect (144).

L. inermis leaf crude extracts in aqueous, methanol, and chloroform demonstrated *in vitro* antimicrobial activity by preventing the development of various strains of pathogenic bacteria (145-147). Studies on the tuberculostatic activity of *L. inermis* were reported by Sharma *et al.* (1990) in both *in vitro* and *in vivo* settings. He reported that 6 g/ml of the herb inhibits the growth of *Mycobacterium tuberculosis* H37Rv and Tubercle bacilli from sputum on the Lowenstein Jensen medium in studies on the *in vitro* tuberculostatic activity of henna. He also stated that in his *in-vivo* studies, guinea pigs and mice treated with *Mycobacterium tuberculosis* H37Rv infection experienced a significant resolution of experimental tuberculosis at a dose of 5 mg/kg body weight (148). *L. inermis* leaf ethanol extract demonstrated antibacterial activity in 1973, according to Abd-el-Malek *et al.* (1973) (149).

Antifungal Activity

Khan and Nasreen (2010) tested 10 phytopathogenic fungi and *Candida albicans* B017 for antifungal activity in methanolic extracts of five different plants. The target fungi's mycelial growth was most significantly inhibited by *L. inermis* among all the extracts tested (76.47-87.77%). In comparison to the nonprotein fractions, the protein fractions of *L. inermis* showed a four to five times greater percentage inhibition of the mycelial growth of *Bipolaris oryzae* and *Colletotrichum lindemuthianum* (150). According to Khan and Nasreen (2010), the active compounds responsible for the effectiveness against plant pathogens were proteinaceous in nature or proteins. Aqueous, methanol, and chloroform crude extracts of *L. inermis* leaves showed *in vitro* antimicrobial activity by inhibiting the growth of different strains of pathogenic fungi (145, 147).

Trypsin Inhibitory Activity

Lawsone (naphthoquinone), sugars, and tannins were detected in the preliminary phytochemical screening of the ethanolic extract. Trypsin inhibition was observed by *L. inermis* alcoholic extract of Lawsone (151).

Wound Healing Activity

L. inermis ethanolic extract was used to test the ability

to heal wounds in rats. The animals were given three sets of six in the excision designs and two sets of six in the dead space and incision designs. The excision wound group received topical treatment, whereas the dead space and incision wound group received oral treatment. When compared to the control group, animals treated with extracts had a 71% lower in the wound site. The use of *L. inermis* in wound healing management is supported by increased wound contraction, hydroxyproline, skin-breaking strength, and histological findings (98). According to the findings of the current study, henna leaf extracts can prevent the development of microorganisms that induce burn wound infections. As a result, this finding lends support to the use of henna in the treatment of burn wound infections. The primary intruders of burnt injuries were tested with water and chloroform extracts of leaves (152).

Anti-Cancer Activity

The anticancer activity of *L. inermis* chloroform extract using an MTT-based cytotoxic assay carried out by Endrini et al. (2002) (153). The mitochondrial dehydrogenase enzyme in these viable tumor cells converts the soluble tetrazolium salt into the insoluble colored formazone. A spectrophotometer is used to measure formazone after it has been dissolved. This extract was tested on normal liver cell lines as well as liver cancer cell lines. The IC₅₀ value explains cell inhibition or cell killing. Cytotoxicity against liver and human breast cancer cell lines was determined using IC₅₀ values of 0.3 and 24.85g/ml, respectively. Mice given *L. inermis* extract were compared to control mice given only water on the 12th day. Control mice had larger diameters of the gluteal solid tumor mass than *L. inermis* treated mice. It was also discovered that extract-treated mice had higher pH levels and lower levels of glutathione lipid peroxidation than control mice. It suggested that the extract could inhibit cancer cell metabolism (154). A similar study found that *L. inermis* extracts inhibited the multiplication of DLA-induced tumor cells in mice. It also increased mice's average survival time and life span. These findings suggest that *L. inermis* could be used as a novel drug in cancer treatment (155).

Antioxidant Activity

L. inermis, also known as henna, has been found to possess significant antioxidant activity. Antioxidants are molecules that help to protect the body's cells from damage caused by free radicals, which are unstable molecules that can cause oxidative stress and lead to various diseases. Studies have shown that henna contains high levels of phenolic compounds, which are potent antioxidants. These compounds are believed to scavenge free radicals and prevent cellular damage caused by oxidative stress. In addition, henna has been found to exhibit strong radical scavenging activity,

which suggests that it may be effective in protecting the body against oxidative damage (156).

Philip et al. (2011) investigated the antioxidant as well as free radical scavenging properties of *L. inermis* seeds. The flavonoid and total phenolic content and antioxidant activity of four different extracts of *L. inermis* seeds are compared with an aqueous extract, ethanol extract, dichloromethane extract, and petroleum ether extract. They discovered that the *L. inermis* ethanolic extract is a more potent antioxidant than the aqueous, petroleum ether, and dichloromethane extracts because it contains a higher concentration of flavonoid and phenolic compounds (157).

Anticorrosion Activity

In a 1 molar HCL solution, henna extracts were used to investigate the inhibitory effect on mild steel corrosion using electrochemical techniques and surface analysis (SEM/EDS). All the tested compounds behave as mixed inhibitors, according to polarisation measurements, with inhibition efficiency rising with inhibitor concentration. The highest level of inhibition efficiency (92.06%) is attained at 1.2 g/l of henna extract. Inhibition becomes more effective in the following order: -D lawsone Tannic acid > glucose. Thermodynamic parameters and inhibition mechanisms are also covered (158).

Analgesic, Anti-Inflammatory, and Antipyretic Activity

In rats, a crude extract in ethanol had significant analgesic, anti-inflammatory, and antipyretic effects. The liquid-liquid extraction method separated the extract into butanol, chloroform, and water fractions, which were then tested for the mentioned activities. The analgesic, anti-inflammatory, and antipyretic effects of the butanol and chloroform fractions were stronger than those of the crude extracts, while the aqueous extract had a significantly lower effect. When compared to the other extracts, butanol extract was the most effective in the analgesic test. A pure compound was isolated from the chloroform extract and identified as 2-hydroxy-1,4- naphthoquinone using chromatographic and spectroscopic techniques (lawsone). The isolated compound was discovered to have an analgesic, anti-inflammatory, and antipyretic effect (159).

L. inermis leaves, which are used in indigenous medicine, were discovered to have anti-inflammatory activity (160).

Gupta et al. (1986) isolated and identified seven crystalline compounds from the chromatographic fraction of an alcoholic extract of *L. inermis* leaves. The fraction yielded luteolin (m.p. 237°C), with a yield of

0.95%. After concentration, the mother liquor yielded traces of lawsone. After removing laxanthone I and lawsone, the ethyl acetate extract was extracted with a saturated sodium carbonate solution (100ml). The alkaline layer was neutralized with concentrated sulphuric acid and extracted with 130ml. of ethyl acetate, yielding laxanthone II (m.p. 180°C), yield 0.47%. The concentration fraction yielded crystals of 3-Oglucoside of-sitosterol (m.p.285°C), yielding 1.87% (161).

Antiparasitic Activity

During an ethnopharmacological study of antiparasitic medicinal herbs used in the Ivory Coast, 17 medicinal plants were recognized and collected. Alkaloid, polar, non-polar, and extracts from different parts of these plants were tested *in vitro* for antiparasitic activity. The activities of antimalarial, leishmanicidal, trypanocidal, anti-helminthics, and anti-scabies drugs were determined. Among the plants studied, *L. inermis* demonstrated promising trypanocide properties (162).

Protein Glycation Inhibition

Protein glycation is a non-enzymatic reaction between proteins and reducing sugars, such as glucose or fructose, resulting in the formation of advanced glycation end products (AGEs). AGEs can accumulate in various tissues and have been implicated in several chronic diseases, including diabetes, neurodegenerative diseases, and cardiovascular diseases. Therefore, inhibiting protein glycation and reducing the formation of AGEs is a potential therapeutic approach to prevent or mitigate these diseases.

Research on *L. inermis* has suggested that it possesses protein glycation inhibitory activity. Studies have shown that the extracts of *L. inermis*, particularly its leaves, contain compounds that can inhibit protein glycation by blocking the formation of AGEs or scavenging reactive carbonyl species, which are involved in the glycation process. These compounds may include phenolic compounds, flavonoids, and other bioactive compounds present in *L. inermis* (163).

Hepatoprotective Activity

Hepatoprotective activity was found in a 90% ethanol extract of *L. inermis* and its ethyl acetate fraction by Chaudhary et al., (2012). Carbon tetrachloride caused hepatotoxicity in rats (CCl₄). Ethanol extract and its ethyl acetate fractions of 200 and 400 mg/kg b.wt. reduced alkaline phosphatase (ALP), serum transaminases (AST and ALT), and total bilirubin significantly (TB). As a result, it has been suggested that *L. inermis* seeds be used to treat liver disorders. Against CCl₄ (0.5 mL/kg, i.p.) induced mice, this extract significantly increases albumin and total

protein levels ($p < 0.01$) in a dose-dependent manner. The seeds extract and its fraction also reduced hepatic malondialdehyde levels by inhibiting free radical production and prevented CCl₄-induced oxidative stress by significantly increasing reducing glutathione levels. 90% ethanol extract of *L. inermis* and its ethyl acetate fraction showed these biochemical parameters were supplemented by histopathological examination of liver sections, which revealed that the ethyl acetate fraction has a more significant ($p < 0.05$) hepatoprotective effect against CCl₄-induced hepatotoxicity in rats (164).

Tapas et al. (2008) discovered that the presence of flavonoids causes hepatoprotective and lipid peroxidation inhibitory properties (165).

Immunomodulatory Effect

According to Mikhail et al. (2004), a 1 mg/ml methanolic extract of henna leaves exhibits immunomodulatory activity as evidenced by the stimulation of T-lymphocyte proliferative responses (166).

As per Dikshit et al. (2000), the Naphthoquinone fraction shows a significant immunomodulatory effect, obtained from leaves *L. inermis* (167).

Ant Sickling Activity

Aqueous extract of *L. inermis* leaves was discovered to inhibit sickling and increase the oxygen affinity of HbSS blood (168).

Enzyme Inhibiting Activities

The IC₅₀ values for trypsin inhibitory activity of ethanol extracts of *L. inermis* leaves and lawsone were 64.87 and 48.6g/ml, respectively (169).

Memory and Behavior Effectiveness

L. inermis influences memory and behavior that is mediated by monoamine neurotransmitters. Using elevated plus maze and passive shock avoidance paradigms, the impact of *L. inermis* extract in acetone-soluble pet. ether on memory was evaluated by Iyer et al., (1998). How clonidine affects hypothermia it causes to investigate the impact on noradrenaline, serotonin, and dopamine-mediated behavior, respectively, the effects of lithium on head twitches and the effects of haloperidol on catalepsy were observed. The pet. ether extract's acetone fraction showed pronounced nootropic activity. The portion that altered the behavior was mediated by 5-HT and NA. The leaves of *L. inermis* have the potential to explore a nootropic principle, it has been determined (140).

Nematicidal Effect

L. inermis has a suppressive effect on Meloidogyne incognita development. When tomato and henna were grown together, henna reduced the number of tomato

root galls, the number of egg-laying females, and the rate of nematode reproduction. Also, when tomato plants were grown in soil containing henna root exudates, a reduction in nematode biological processes was observed. When henna was grown alone, the root gall index and nematode production rate were reduced by 75% and 99%, respectively, when compared to tomatoes grown alone (170).

Anticoagulant Effect

Lawson and its oxazine derivatives isolated from *L. inermis* leave shown to be potential anticoagulant agents (171).

Gingivitis Healing Activity

The efficacy of *L. inermis* leaves methanol extracts (62.500, 31.250, and 15.625 g/ml) in healing gingivitis was investigated in Sprague Dawley rats with mandibular labial gingiva inflammation induced by 10% H₂O₂. There was no difference in healing between the three concentrations of *L. inermis* leaves methanol extract and povidone-iodine, but there were differences between the three concentrations. A higher concentration (62.500 g/ml) can hasten inflammatory cell reduction and epithelial connective tissue repair (172).

Clinical trials were conducted to investigate the effect of *L. inermis* leaves infusion on gingivitis healing. Sixty-three gingivitis patients were instructed to rinse with *L. inermis* leaves infusion at three concentrations (50000, 10000, and 5000 g/ml), 0.1% hexetidine solution, and placebo as a control. *L. inermis* leaves infusion at 10000 g/ml concentration (80%) reduced bleeding index more than hexetidine 0.1% (76%) (173).

Anti-Urolithiasis Activity

L. inermis plants have traditionally been used to treat urolithiasis (kidney stones). Several studies have investigated the potential anti-urolithiasis action of *L. inermis*. One study evaluated the impact of *L. inermis* leaf extract in ethanol on calcium oxalate-induced urolithiasis in rats. The findings revealed that the extract significantly reduced the number and size of calcium oxalate crystals in the urine and kidneys of the rats, indicating a potential anti-urolithiasis activity (174).

Another study looked into the impact of *L. inermis* leaf extract in methanol on forming of calcium oxalate crystals *in vitro*. The extract significantly reduced the formation of calcium oxalate crystals, implying a potential anti-urolithiasis effect (175).

Antidiarrheal Effects

The castor oil-induced diarrhea model in mice was used to test the anti-diarrheal properties of an ethanol extract of the leaf of *L. inermis*. In comparison to the

control group, the ethanol extract at a dose of 500 mg/kg had antidiarrheal activity and provided approximately 1.398 of the mean latent periods for the diarrhoeal episode ($p < 0.002$) (176).

Diuretic Activity

Several studies have been conducted to evaluate the diuretic function of *L. inermis*. In one study, aqueous *L. inermis* extract was tested for diuretic activity in rats. The extract increased urine output and electrolyte excretion significantly more than the control group, indicating potential diuretic activity. In rats, *L. inermis* leaf extract in ethanol significantly increased urine output and electrolyte excretion, indicating a potential diuretic effect (177).

Anticataleptic Activity

An aqueous extract of henna was found to be effective in treating haloperidol-induced catalepsy in mice. At a dose of 400 mg/kg, there was a reduction in cataleptic scores and an increase in superoxide dismutase activity (178).

Synergistic Effect

According to Bhuvaneswari et al. (2002), *Staphylococcus aureus*, *Klebsiella pneumonia*, *Pseudomonas aeruginosa*, and *Proteus mirabilis* are the primary causes of urinary tract infections. *S. aureus* causes boils, pimples, and other skin conditions that were previously treated with plant leaves (179).

Compared to medications (Gentamycin, Erythromycin, Tetracycline, Chloramphenicol, and Streptomycin) that inhibit protein synthesis, two classes of antibiotics—that inhibit cell walls and those that inhibit nucleic acids—are less effective. Since many organisms today are resistant to antibiotics, this study uses the synergistic effects of plant extracts to create drugs that treat various diseases (180).

Abortifacient Activity

The extract of *L. inermis* roots in methanol was studied for its abortifacient properties by Aguwa (1987), who found that it induces abortion in rats, mice, and guinea pigs in a dose-dependent manner. The results were confirmed by its ethnomedical use in some regions of Nigeria to induce human abortion (181).

Toxicity Studies in Henna

Most toxicological studies claim that hepatotoxicity is connected to toxic effects brought on using herbal medicine. Medical journals have also reported on mutagenicity, carcinogenicity, and other toxic effects on the kidney, nervous system, blood, and cardiovascular system. To conduct any cutting-edge biological experimental techniques have been employed to conduct standard safety tests before the efficacy study to the literature, *L. inermis* has been

shown to have significant analgesic, antioxidant, anti-inflammatory, antibacterial, hepatoprotective, and adaptogenic effects demonstrating that it is a substance that can be used as a drug regularly without causing harm (182).

Prospects of Henna Research

A few workers are looking into the many different uses of henna leaf, extracts, and powder. It's been proposed that the cold and hot aqueous leaf extracts of henna have good staining qualities and make a good substitute for the counter stains used in the Gram staining procedure (183). Henna extract in ethyl acetate demonstrated superior corrosion inhibition and may be used as a green inhibitor to prevent corrosion from aluminum alloy (184). Some of Lawsone's biological activities have been attributed in part to its capacity for redox cycling and the chelation of trace metal ions (114, 185). It has also been demonstrated that silver nanoparticles made from leaf aqueous extract have loculicidal activity (186, 187). Another study discovered that henna-derived molluscicides, either alone or in combination with acetogenins, reduced the fertility, and survival of young snails significantly (188).

Conclusions

Today, the cosmetics industry occupies a unique position. Cosmetic companies are free to develop and market products that are known to affect the structure and function of skin, with little oversight. Most consumers believe that cosmeceuticals are regulated and tested in the same way that drugs are. Consumers believe that ingredients have been tested for safety and that advertising claims are true. Consumers also believe that claims such as "natural," "cruelty-free," and "hypoallergenic" are true and substantiated. The truth is that active cosmeceutical and pharmaceutical ingredients have never been more closely related. When lawmakers mandated the regulatory structure for cosmetics decades ago, they could not have imagined this evolving field.

The data presented here indicated the use of *L. inermis* plant as herbal medicines and bioactive compounds for cosmetic purposes and treatment of various diseases, and it is based on the product's correcting redient requirements and superiority. Herbal cosmetics must be subjected to stringent quality control measures to ensure their safety. Exploration of various types of the literature reveals that the *L. inermis* plant has a broad spectrum of pharmacological activities, and these activities allow it to be used as a remedy in herbal medicines. Because this plant contains a wide range of phytoconstituents, it can treat a broad range of ailments. This plant has antibacterial, antiviral, antimycotic, antimicrobial, and antifungal properties, among others. Because this plant has a

wide range of therapeutic properties, scientists and researchers should take it into account when developing a game-changing drug right now. However, more research is needed to uncover *L. inermis*' hidden potential and its therapeutic uses for the benefit of humans.

Declarations

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References

1. Joshi LS, Pawar HA. Herbal cosmetics and cosmeceuticals: An overview. *Nat Prod Chem Res.* (2015) 3(2):170.
2. Aburjai T, Natsheh FM. Plants used in cosmetics.

Phytother Res. (2003) 17(9):987-1000.

3. Biswas P, Anand U, Saha SC, Kant N, Mishra T, Masih H, et al. Betelvine (*Piper betle* L.): A comprehensive insight into its ethnopharmacology, phytochemistry, and pharmacological, biomedical, and therapeutic attributes. *J Cell Mol Med.* (2022) 26(11):3083-119.

4. Gediya SK, Mistry RB, Patel UK, Blessy M, Jain HN. Herbal plants: used as cosmetics. *J Nat Prod Plant Resour.* (2011) 1(1):24-32.

5. Kuchekar BS, (2008 Jan 8) *Pharmaceutical Jurisprudence*, Pragati Books Pvt. Ltd., 9788185790282, 1st Edition.

6. Jain NK. (2007), *A textbook of Forensic pharmacy*, MK Jain Vallabh Prakashan, 818573187X, 7th Edition.

7. Udupa N, Popli H, (2010), *Pharmaceuticals, cosmeceuticals, and nutraceuticals: an overview of regulations*. Career Publications; 9788188739837, 1st Edition.

8. Dureja H, Kaushik D, Gupta M, Kumar V, Lather V. Cosmeceuticals: An emerging concept. *Indian J Pharmacol.* (2005) 37(3):155.

9. Martin KI, Glaser DA. Cosmeceuticals: the new medicine of beauty. *Mo Med.* (2011) 108(1):60-3.

10. Trüeb RM. The Value of Hair Cosmetics and Pharmaceuticals. *Dermatology.* (2001) 202(4):275-82.

11. Datta HS, Paramesh R. Trends in aging and skin care: Ayurvedic concepts. *J Ayurveda Integr Med.* (2010) 1(2):110-3.

12. Rathod S, Mali S, Shinde N, Aloorkar N. Cosmeceuticals and Beauty Care Products: Current trends with prospects. *Res J Top Cosmet Sci.* (2020) 11(1):45.

13. Tabassum N, Hamdani M. Plants used to treat skin diseases. *Pharmacogn Rev.* (2014) 8(15):52.

14. Saraf S, Kaur C. Phytoconstituents as photoprotective novel cosmetic formulations. *Pharmacogn Rev.* (2010) 4(7):1.

15. Furmanowa M, Skopińska-Rozewska E, Rogala E, Hartwich M. *Rhodiola rosea* in vitro culture - phytochemical analysis and antioxidant action. *Acta Soc Bot Pol.* (2014) 67(1):69-73.

16. Eroglu A, Hruszkewycz DP, Dela Sena C, Narayanasamy S, Riedl KM, Kopec RE, et al. Naturally Occurring Eccentric Cleavage Products of Provitamin A β -Carotene Function as Antagonists of Retinoic Acid Receptors. *J Biol Chem.* (2012) 287(19):15886-95.

17. Gupta RK, Soni P, Shrivastava J, Rajput P, Parashar S. Cosmeceutical role of Medicinal plants / Herbs: A Review on commercially available Cosmetic

ingredients. *Int J Innov Sci Technol.* (2018) 9:70-3.

18. Mohammed Rahmoun N, Boucherit-Atmani Z, Benabdallah M, Boucherit K, Villemin D, Choukchou-Braham N. Antimicrobial Activities of the Henna Extract and Some Synthetic Naphthoquinones Derivatives. *Am J Med Biol Res.* (2013) 1(1):16-22.

19. Agrawal S, Bablani Popli D, Sircar K, Chowdhry A. A review of the anticancer activity of *Azadirachta indica* (Neem) in oral cancer. *J Oral Biol Craniofacial Res.* (2020) 10(2):206-9.

20. Draelos ZD. Cosmeceuticals: undefined, unclassified, and unregulated. *Clin Dermatol.* (2009) 27(5):431-4.

21. Chaudhary G, Goyal S, Poonia P. *Lawsonia inermis* Linnaeus: a phytopharmacological review. *Int J Pharm Sci Drug Res.* (2010) 2(2):91-8.

22. Khanpara K, Renuka V, Harisha C. A detailed investigation on shikakai (*Acacia concinna* Linn.) fruit. *J. Curr. Pharm. Res.* (2012) 9:6-10.

23. Adelman MJ, Bedford LM, Potts GA. Clinical efficacy of popular oral hair growth supplement ingredients. *Int J Dermatol.* (2021) 60(10):1199-210.

24. Thoidingjam S, Tiku AB. Therapeutic efficacy of *Phyllanthus emblica*-coated iron oxide nanoparticles in A549 lung cancer cell line. *Nanomed.* (2019) 14(17):2355-71.

25. Al Badi K, Khan SA. Formulation, evaluation, and comparison of the herbal shampoo with the commercial shampoos. *Beni-Suef Univ J Basic Appl Sci.* (2014) 3(4):301-5.

26. Thorén S, Yazar K. Contact allergens in 'natural' hair dyes: CONTACT ALLERGENS IN 'NATURAL' HAIR DYES. *Contact Dermatitis.* (2016) 74(5):302-4.

27. Rendon MI, Gaviria JI. Review of skin-lightening agents. *Dermatol Surg.* (2005) 31:886-90.

28. Choi CM, Berson DS. Cosmeceuticals. *Semin Cutan Med Surg.* (2006) 25(3):163-8.

29. Boonsong P, Laohakunjit N, Kerdchoechuen O, Matta FB. Detection of Pigments and Natural Colorants from Thai Herbal Plants for Possible Use as Coloring Dyes. *HortScience.* (2011) 46(2):265-72.

30. Valadas LA, Oliveira Filho RD, Rodrigues Neto EM, Bandeira MA, Fonteles MM, Passos VF, Fiallos AC, Lotif MA, Sena NJ, Dantas TC, Lima Soares I. *Camellia sinensis* in dentistry: Technological prospection and scientific evidence. *Evid Based Complement Alternat Med.* (2021) 31:2021.

31. Watanabe F, Hashizume E, Chan GP, Kamimura A. Skin-whitening and skin-condition-improving effects of topical oxidized glutathione: a double-blind and

- placebo-controlled clinical trial in healthy women. *Clin Cosmet Investig dermatol.* (2014) 17:267-74.
32. Varoni E, Iriti M. *Odontonutraceuticals: Pleiotropic Phytotherapeutic Agents for Oral Health.* Pharmaceuticals (Basel). (2016) 9(1):10.
 33. Goyal A, Sharma A, Kaur J, Kumari S, Garg M, Sindhu RK, et al. *Bioactive-Based Cosmeceuticals: An Update on Emerging Trends.* Mol Basel Switz. (2022) 27(3):828.
 34. Majeed M, Majeed S, Nagabhushanam K, Mundkur L, Neupane P, Shah K. *Clinical Study to Evaluate the Efficacy and Safety of a Hair Serum Product in Healthy Adult Male and Female Volunteers with Hair Fall.* Clin Cosmet Investig Dermatol. (2020) 13:691-700.
 35. Boelsma E, Hendriks HF, Roza L. *Nutritional skin care: health effects of micronutrients and fatty acids.* Am J Clin Nutr. (2001) 73(5):853-64.
 36. Oh JY, Park MA, Kim YC. *Peppermint Oil Promotes Hair Growth without Toxic Signs.* Toxicol Res. (2014) 30(4):297-304.
 37. Rodriguez-Amaya DB. *Natural food pigments and colorants.* Curr Opin Food Sci. (2016) 7:20-6.
 38. Ichihashi M, Ueda M, Budiyo A, Bito T, Oka M, Fukunaga M, et al. *UV-induced skin damage.* Toxicology. (2003) 189(1-2):21-39.
 39. Larsson SC, Bergkvist L, Näslund I, Rutegård J, Wolk A. *Vitamin A, retinol, and carotenoids and the risk of gastric cancer: a prospective cohort study.* Am J Clin Nutr. (2007) 85(2):497-503.
 40. Colvard MD, Cordell GA, Villalobos R, Sancho G, Soejarto DD, Pestle W, et al. *Survey of medical ethnobotanicals for dental and oral medicine conditions and pathologies.* J Ethnopharmacol. (2006) 107(1):134-42.
 41. Newburger AE. *Cosmeceuticals: myths and misconceptions.* Clin Dermatol. (2009) 27(5):446-52.
 42. Deen A, Visvanathan R, Wickramarachchi D, Marikkar N, Nammi S, Jayawardana BC, et al. *Chemical composition and health benefits of coconut oil: an overview.* J Sci Food Agric. (2021) 101(6):2182-93.
 43. Cowan MM. *Plant Products as Antimicrobial Agents.* Clin Microbiol Rev. (1999) 12(4):564-82.
 44. Soler L, Canellas J, Saura-Calixto F. *Oil content and fatty acid composition of developing almond seeds.* J Agric Food Chem. (1988) 36(4):695-7.
 45. Tan QG, Cai XH, Du ZZ, Luo XD. *Three Terpenoids and a Tocopherol-Related Compound from Ricinus communis.* Helv Chim Acta. (2009) 92(12):2762-8.
 46. Al-Waili NS. *Topical application of natural honey, beeswax, and olive oil mixture for atopic dermatitis or psoriasis: partially controlled, single-blinded study.* Complement Ther Med. (2003) 11(4):226-34.
 47. Fratini F, Cilia G, Turchi B, Felicioli A. *Beeswax: A minireview of its antimicrobial activity and its application in medicine.* Asian Pac J Trop Med. (2016) 9(9):839-43.
 48. Chowdhury MMU. *Allergic contact dermatitis from prime yellow carnauba wax and coathylene in mascara: CONTACT POINT.* Contact Dermatitis. (2002) 46(4):244-244.
 49. Yousuf B, Mir NA, Bhardwaj M, Gul K, Wani AA (2019) *Introduction to Food Hydrocolloids.* In: Food Hydrocolloids as Encapsulating Agents in Delivery Systems, Editor: Gani A, Masoodi FA, Shah U, Shah A, CRC Press, 978-1461360599, 1st edition
 50. B Suchithra A, Jeganath S, Jeevitha E. *Pharmaceutical Gels and Recent Trends-A Review.* Res J Pharm Technol. (2019) 12(12):6181.
 51. Wińska K, Mączka W, Łyczko J, Grabarczyk M, Czubaśzek A, Szumny A. *Essential oils as antimicrobial agents—myth or real alternative.* Molecules. (2019) 24(11):2130.
 52. Agarwal A, Chaudhary B. *Clinical and microbiological effects of 1% Matricaria chamomilla mouth rinse on chronic periodontitis: A double-blind randomized placebo-controlled trial.* J Indian Soc Periodontol. (2020) 24(4):354-61.
 53. Silvestre WP, Medeiros FR, Agostini F, Toss D, Pauletti GF. *Fractionation of rosemary (Rosmarinus officinalis L.) essential oil using vacuum fractional distillation.* J Food Sci Technol. (2019) 56(12):5422-34.
 54. Fischer A, Brodziak-Dopierała B, Loska K, Stojko J. *The Assessment of Toxic Metals in Plants Used in Cosmetics and Cosmetology.* Int J Environ Res Public Health. (2017) 14(10):1280.
 55. Fu B, Li H, Wang X, Lee FSC, Cui S. *Isolation and Identification of Flavonoids in Licorice and a Study of Their Inhibitory Effects on Tyrosinase.* J Agric Food Chem. (2005) 53(19):7408-14.
 56. Rendon MI, Gaviria JL. *Review of skin-lightening agents.* Dermatol surg. (2005) 31:886-90.
 57. Bilia AR, Isacchi B, Righeschi C, Guccione C, Bergonzi MC. *Flavonoids Loaded in Nanocarriers: An Opportunity to Increase Oral Bioavailability and Bioefficacy.* Food Nutr Sci. (2014) 05(13):1212-327.
 58. Nakagawa M, Kawai K, Kawai K. *Contact allergy to kojic acid in skin care products.* Contact dermatitis. (1995) 32(1):9-13.
 59. Fuchs J. *Potentials and limitations of the natural*

- antioxidants RRR- α -tocopherol, L-ascorbic acid, and β -carotene in cutaneous photoprotection. *Free Radic Biol Med.* (1998) 25(7):848-73.
60. Kaul S, Gulati N, Verma D, Mukherjee S, Nagaich U. Role of Nanotechnology in Cosmeceuticals: A Review of Recent Advances. *J Pharm.* (2018) 2018:3420204.
61. Sumbul S, Ahmad MA, Asif M, Akhtar M, Saud I. Physicochemical and phytochemical standardization of berries of *Myrtus communis* Linn. *J Pharm Bioallied Sci.* (2012) 4(4):322-6.
62. Khristi V, Patel VH. Therapeutic potential of *Hibiscus rosa Sinensis*: A review. *Int J Nutr Diet.* (2017) 4(2):105-23.
63. Rahman MM, Uddin MJ, Reza ASMA, Tareq AM, Emran TB, Simal-Gandara J. Ethnomedicinal Value of Antidiabetic Plants in Bangladesh: A Comprehensive Review. *Plants Basel Switz.* (2021) 10(4):729.
64. Kalwij JM. Review of 'The Plant List, a working list of all plant species.' Palmer M, editor. *J Veg Sci.* (2012) 23(5):998-1002.
65. Gallo FR, Multari G, Giambenedetti M, Federici E. Chemical fingerprinting of *Lawsonia inermis* L. using HPLC, HPTLC, and densitometry. *Phytochem Anal.* (2008) 19(6):550-9.
66. Moro P, Morina M, Milani F, Pandolfi M, Guerriero F, Bernardo L. Sensitization and Clinically Relevant Allergy to Hair Dyes and Clothes from Black Henna Tattoos: Do People Know the Risk? An Uncommon Serious Case and a Review of the Literature. *Cosmetics.* (2016) 3(3):23.
67. Moor, Johannes Cornelis (1971) The Seasonal Pattern in the Ugaritic Myth of Ba'lu according to the Version of Ilmilku In *Alter Orient und Altes Testament*, Editor: Degen R, Butzon & Bercker, 3788702931, 1st Edition
68. S H Hook (2021) *Middle Eastern Mythology*, HASSELL STREET Press, 1013640543, 1st Edition
69. Jallad KN, Espada-Jallad C. Lead exposure from the use of *Lawsonia inermis* (Henna) in temporary paint-on-tattooing and hair dying. *Sci total environ.* (2008) 397(1-3):244-50.
70. Badoni Semwal R, Semwal DK, Combrinck S, Cartwright-Jones C, Viljoen A. *Lawsonia inermis* L. (henna): Ethnobotanical, phytochemical and pharmacological aspects. *J Ethnopharmacol.* (2014) 155(1):80-103.
71. Ahmad Supian FN, Osman NI. Phytochemical and Pharmacological Activities of Natural Dye Plant, *Lawsonia inermis* L. (Henna). *J Young Pharm.* (2023) 15(2):201-11.
72. Nishteswar K. Pharmacological expression of *Rasayanakarma*. *Ayu.* (2013) 34(4):337-8.
73. Singh DK, Luqman S, Mathur AK. *Lawsonia inermis* L. – A commercially important primeval dying and medicinal plant with diverse pharmacological activity: A review. *Ind Crops Prod.* (2015) 65:269-86.
74. Zumrutdal E, Ozaslan M. A Miracle Plant for the Herbal Pharmacy; Henna (*Lawsonia inermis*). *Int J Pharmacol.* (2012) 8(6):483-9.
75. Nayak BS, Isitor G, Davis EM, Pillai GK. The evidence-based wound healing activity of *Lawsonia inermis* Linn. *Phytother Res.* (2007) 21(9):827-31.
76. Al-Rubiay KK, Jaber NN, Al-Mhaawe BH, Alrubaiy LK. Antimicrobial efficacy of henna extracts. *Oman Med J.* (2008) 23(4):253-6.
77. Dixit S, Ali H. Antioxidant Potential Some Medicinal Plants of Central India. *J Cancer Ther.* (2010) 01(02):87-90.
78. Hizli H. A study on the use of the henna plant (*Lawsonia inermis* Linn) for the treatment of fungal disease (*Trichophyton verrucosum*) in calves. *J Hell Vet Med Soc.* (2021) 71(4):2483.
79. Ponnusamy K, Petchiammal C, Mohankumar R, Hopper W. In vitro antifungal activity of indirubin isolated from a South Indian ethnomedicinal plant *Wrightia tinctoria* R. Br. *J Ethnopharmacol.* (2010) 132(1):349-54.
80. Velayutham K, Rahuman AA, Rajakumar G, Roopan SM, Elango G, Kamaraj C, et al. Larvicidal activity of green synthesized silver nanoparticles using bark aqueous extract of *Ficus racemosa* against *Culex quinquefasciatus* and *Culex gelidus*. *Asian Pac J Trop Med.* (2013) 6(2):95-101.
81. Kumar M, Kaur P, Chandel M, Singh AP, Jain A, Kaur S. Antioxidant and hepatoprotective potential of *Lawsonia inermis* L. leaves against 2-acetylaminofluorene induced hepatic damage in male Wistar rats. *BMC Complement Altern Med.* (2017) 17(1):56.
82. Ali NAA, Jülich WD, Kusnick C, Lindequist U. Screening of Yemeni medicinal plants for antibacterial and cytotoxic activities. *J Ethnopharmacol.* (2001) 74(2):173-9.
83. Smith GE (1930) *Ancient Egyptian medicine: the papyrus ebers*. Ares Publishers, 1684225221, 5th Edition
84. Holtzman L. Accused of Anthropomorphism: Ibn Taymiyya's Miḥan as Reflected in Ibn Qayyim al-Jawziyya's al-Kāfiya al-Shāfiya: Accused of Anthropomorphism. *Muslim World.* (2016) 106(3):561-87.

85. Malekzadeh F. Antimicrobial activity of *Lawsonia inermis* L. *Appl Microbiol.* (1968) 16(4):663-4.
86. Mikhaeil BR, Badria FA, Maatooq GT, Amer MMA. Antioxidant and Immunomodulatory Constituents of Henna Leaves. *Z Naturforsch C J Biosci.* (2004) 59(7-8):468-76.
87. Sultana N, Choudhary MI, Khan A. Protein glycation inhibitory activities of *Lawsonia inermis* and its active principles. *J Enzyme Inhib Med Chem.* (2009) 24(1):257-61.
88. Chang H, Suzuka SE. Lawsone (2-OH-1,4-naphthoquinone) derived from the henna plant increases the oxygen affinity of sickle cell blood. *Biochem Biophys Res Commun.* (1982) 107(2):602-8.
89. Tripathi RD, Srivastava HS, Dixit SN. A fungitoxic principle from the leaves of *Lawsonia inermis* Lam. *Experientia.* (1978) 34(1):51-2.
90. Hsouna AB, Trigui M, Culioli G, Blache Y, Jaoua S. Antioxidant constituents from *Lawsonia inermis* leaves Isolation, structure elucidation, and antioxidative capacity. *Food Chem.* (2011) 125(1):193-200.
91. Clarke DT, Jones GR, Martin MM. The anti-sickling drug lawsone (2-OH-1, 4-naphthoquinone) protects sickled cells against membrane damage. *Biochem Biophys Res Commun.* (1986) 139(2):780-6.
92. Wagner H, Kreher B, Jurcic K. In vitro stimulation of human granulocytes and lymphocytes by pico- and femtogram quantities of cytostatic agents. *Arzneimittelforschung.* (1988) 38(2):273-5.
93. Anand KK, Singh B, Chand D, Chandan BK. An evaluation of *Lawsonia alba* extracts as a hepatoprotective agent. *Planta med.* (1992) 58(01):22-5.
94. Alia BH, Bashir AK, Tanira MO. Anti-inflammatory, antipyretic, and analgesic effects of *Lawsonia inermis* L.(henna) in rats. *Pharmacology.* (1995) 51(6):356-63.
95. Ignacimuthu S, Shanmugam N. Antimycobacterial activity of two natural alkaloids, vasicine acetate and 2-acetyl benzylamine, isolated from Indian shrub *Adhatoda vasica* Ness. leaves. *J Biosci.* (2010) 35(4):565-70.
96. Gupta S, Ali M, Alam MS. A naphthoquinone from *Lawsonia inermis* stem bark. *Phytochemistry.* (1993) 33(3):723-4.
97. Singh DK, Luqman S. *Lawsonia inermis* (L.): A perspective on anticancer potential of Mehndi/Henna. *Biomed Res Ther.* (2014) 1(4):18.
98. Kiraz N, Metintas S, Oz Y, Koc F, Koku Aksu EA, Kalyoncu C, et al. The prevalence of tinea pedis and tinea manuum in adults in rural areas in Turkey. *Int J Environ Health Res.* (2010) 20(5):379-86.
99. Jacob PP, Saral AM. Two Harmala Alkaloids from *Lawsonia inermis* Seeds. *Chem Nat Compd.* (2013) 49(4):780-780.
100. Charoensup R, Duangyod T, Palanuvej C, Ruangrunsi N. Pharmacognostic specifications and lawsone content of *Lawsonia inermis* leaves. *Pharmacogn Res.* (2017) 9(1):60.
101. Chopra RN, Nayar SL, Chopra IC (1956), Glossary of Indian medicinal plants, Council of Scientific & Industrial Research, 8172360487, 15th Edition
102. Sharma J, Gairola S, Gaur RD, Painuli RM. The treatment of jaundice with medicinal plants in indigenous communities of the Sub-Himalayan region of Uttarakhand, India. *J Ethnopharmacol.* (2012) 143(1):262-91.
103. Vimalanathan S, Ignacimuthu S, Hudson JB. Medicinal plants of Tamil Nadu (Southern India) are a rich source of antiviral activities. *Pharm Biol.* (2009) 47(5):422-9.
104. Bhandarkar M, Khan A. Protective effect of *Lawsonia alba* Lam., against CCl₄ induced hepatic damage in albino rats. *Indian J Exp Biol.* (2003) 41(1):85-7.
105. Joshi R, Satyal P, Setzer W. Himalayan Aromatic Medicinal Plants: A Review of their Ethnopharmacology, Volatile Phytochemistry, and Biological Activities. *Medicines.* (2016) 3(1):6.
106. Patel S, Sharma V, Chauhan N, Thakur M, Dixit VK. Hair Growth: Focus on Herbal Therapeutic Agent. *Curr Drug Discov Technol.* (2015) 12(1):21-42.
107. Oladunmoye, M K, Kehinde, F Y. Ethnobotanical survey of medicinal plants used in treating viral infections among Yoruba tribe of South Western Nigeria. *Afr J Microbiol Res.* (2011) 5(19):2991-3004.
108. Olorunnisola OS, Adetutu A, Balogun EA, Afolayan A J. Ethnobotanical survey of medicinal plants used in the treatment of malarial in Ogbomoso, Southwest Nigeria. *J Ethnopharmacol.* (2013) 150(1):71-8.
109. Atawodi SE, Ameh DA, Ibrahim S, Andrew JN, Nzelibe HC, Onyike EO, et al. Indigenous knowledge system for treatment of trypanosomiasis in Kaduna state of Nigeria. *J Ethnopharmacol.* (2002) 79(2):279-82.
110. Raghuvanshi D, Dhalaria R, Sharma A, Kumar D, Kumar H, Valis M, et al. Ethnomedicinal Plants Traditionally Used for the Treatment of Jaundice (Icterus) in Himachal Pradesh in Western Himalaya—A Review. *Plants.* (2021) 10(2):232.
111. Gupta DK, Arumugam K, Mohamed MN, Bhatt RK,

- Kumar P, Shukla AK, et al. Dynamics of biomass and soil carbon sequestration across an age-sequence of *Lawsonia inermis* plantation in semi-arid Region, Rajasthan, India. *Indian J Agric Sci.* (2021) 92(6):705–10.
112. Mohamed M, Eldin I, Mohammed A, Hassan H. Effects of *Lawsonia inermis* L. (Henna) Leaves' Methanolic Extract on CCl₄-induced Hepatotoxicity in Rats. *J Intercult Ethnopharmacol.* (2016) 5(1):22.
113. Oda Y, Nakashima S, Kondo E, Nakamura S, Yano M, Kubota C, et al. Comparison of lawsone contents among *Lawsonia inermis* plant parts and neurite outgrowth accelerators from branches. *J Nat Med.* (2018) 72(4):890–6.
114. Pradhan R, Dandawate P, Vyas A, Padhye S, Biersack B, Schobert R, et al. From Body Art to Anticancer Activities: Perspectives on Medicinal Properties of Henna. *Curr Drug Targets.* (2012) 13(14):1777–98.
115. Rahmoun N, Boucherit-Otmani Z, Boucherit K, Benabdallah M, Choukchou-Braham N. Antifungal activity of the Algerian *Lawsonia inermis* (henna). *Pharm Biol.* (2013) 51(1):131–5.
116. Aldayarov N, Tulobaev A, Salykov R, Jumabekova J, Kydyralieva B, Omurzakova N, et al. An ethnoveterinary study of wild medicinal plants used by the Kyrgyz farmers. *J Ethnopharmacol.* (2022) 285:114842.
117. Volpato G, Kourková P, Zelený V. Healing war wounds and perfuming exile: the use of vegetal, animal, and mineral products for perfumes, cosmetics, and skin healing among Sahrawi refugees of Western Sahara. *J Ethnobiol Ethnomed.* (2012) 8(1):49.
118. Hafiz H, Chukwu O, Nura S. The potentials of henna (*Lawsonia inermis* L.) leaves extracts as counter stain in gram staining reaction. *Bayero J Pure Appl Sci.* (2013) 5(2):56–60.
119. Kraeling MEK, Bronaugh RL, Jung CT. Absorption of Lawsone Through Human Skin. *Cutan Ocul Toxicol.* (2007) 26(1):45–56.
120. Kapadia G, Rao G, Sridhar R, Ichiishi E, Takasaki M, Suzuki N, et al. Chemoprevention of Skin Cancer: Effect of *Lawsonia inermis* L. (Henna) Leaf Powder and its Pigment Artifact, Lawsone in the Epstein- Barr Virus Early Antigen Activation Assay and in Two-Stage Mouse Skin Carcinogenesis Models. *Anticancer Agents Med Chem.* (2013) 13(10):1500–7.
121. Mehendale VG, Chaudhari NC, Shenoy SN, Mehendale AV. Henna as a Durable Preoperative Skin Marker. *World J Surg.* (2011) 35(2):311–5.
122. Marzin D, Kirkland D. 2-Hydroxy-1,4-naphthoquinone, the natural dye of Henna, is non-genotoxic in the mouse bone marrow micronucleus test and does not produce oxidative DNA damage in Chinese hamster ovary cells. *Mutat Res Toxicol Environ Mutagen.* (2004) 560(1):41–7.
123. Vančo J, Trávníček Z, Hošek J, Suchý P. In vitro and in vivo anti-inflammatory active copper(II)-lawsone complexes. Shahid M, editor. *PLOS ONE.* (2017) 12(7):e0181822.
124. Opretzka LCF, Espírito-Santo RFD, Nascimento OA, Abreu LS, Alves IM, Döring E, et al. Natural chromones as potential anti-inflammatory agents: Pharmacological properties and related mechanisms. *Int Immunopharmacol.* (2019) 72:31–9.
125. Reich E, Widmer V. Plant Analysis 2008 – Planar Chromatography. *Planta Med.* (2009) 75(07):711–8.
126. Munawar T M, Rao CK, Rajesh AV, Sharma UN, Gupta I, Surya Prakash DV. Studies on Cytotoxic and Genotoxic potential of Ethanolic extract of *Lawsonia inermis* leaves. *Int J Health Sci.* (2022) 11:4271–9.
127. Hassan Wagini N. Phytochemical Analysis of Nigerian and Egyptian Henna (*Lawsonia inermis* L.) Leaves using TLC, FTIR, and GCMS. *Plant.* (2014) 2(3):27.
128. Al-Snafi AE, Talab TA, Alfuraiji N. The analgesic and anti-inflammatory effect of Lawsone isolated from *Lawsonia inermis*. *Sci Pharm Sci.* (2022) 1(35):77–84.
129. Patel KM, Patel PR. Review on *Lawsonia inermis* Linn.: An Update. *Asian J Pharm Technol.* (2017) 7(4):237.
130. Thalkari AB, Karwa PN, Shinde PS, Gawli CS, Chopane PS. Pharmacological actions of *Tridax procumbens* L.: A Scientific Review. *Res J Pharmacogn Phytochem.* (2020) 12(1):27.
131. Neela F, Khan MSI, Islam M, Alam M, Akter A. Screening of ethanol, petroleum ether and chloroform extracts of medicinal plants, *Lawsonia inermis* L. and *Mimosa pudica* L. for antibacterial activity. *Indian J Pharm Sci.* (2010) 72(3):388.
132. Habbal O, Hasson S, El-Hag A, Al-Mahrooqi Z, Al-Hashmi N, Al-Bimani Z, et al. Antibacterial activity of *Lawsonia inermis* Linn (Henna) against *Pseudomonas aeruginosa*. *Asian Pac J Trop Biomed.* (2011) 1(3):173–6.
133. Habbal OA, Al-Jabri AA, El-Hag AH, Al-Mahrooqi ZH, Al-Hashmi NA. In-vitro antimicrobial activity of *Lawsonia inermis* Linn (henna). A pilot study on the Omani henna. *Saudi Med J.* (2005) 26(1):69–72.
134. Al-Waili NS. Topical application of natural honey, beeswax, and olive oil mixture for atopic dermatitis or psoriasis: partially controlled, single-blinded study. *Complement Ther Med.* (2003) 11(4):226–34.

135. Manuja A, Rathore N, Choudhary S, Kumar B. Phytochemical Screening, Cytotoxicity and Anti-inflammatory Activities of the Leaf Extracts from *Lawsonia inermis* of Indian Origin to Explore their Potential for Medicinal Uses. *Med Chem.* (2021) 17(6):576–86.
136. Khantamat O, Dukaew N, Karinchai J, Chewonarin T, Pitchakarn P, Temviriyankul P. Safety and bioactivity assessment of aqueous extract of Thai Henna (*Lawsonia inermis* Linn.) Leaf. *J Toxicol Environ Health A.* (2021) 84(7):298–312.
137. Dasgupta T, Rao AR, Yadava PK. Modulatory effect of the henna leaf (*Lawsonia inermis*) on drug metabolizing phase I and phase II enzymes, antioxidant enzymes, lipid peroxidation, and chemically induced skin and forestomach papilloma genesis in mice. *Mol Cell Biochem.* (2003) 245(1-2):11–22.
138. Itoigawa M, Ito C, Tan HT, Okuda M, Tokuda H, Nishino H, et al. Cancer chemopreventive activity of naphthoquinones and their analogs from *Avicennia* plants. *Cancer Lett.* (2001) 174(2):135–9.
139. Hsouna AB, Mongi S, Culioli G, Blache Y, Ghlissi Z, Chaabane R, et al. Protective effects of ethyl acetate fraction of *Lawsonia inermis* fruits extract against carbon tetrachloride-induced oxidative damage in rat liver. *Toxicol Ind Health.* (2016) 32(4):694–706.
140. Akhtar N, Khan BA, Majid A, Khan HMS, Mahmood T, Gulfshan null, et al. Pharmaceutical and biopharmaceutical evaluation of extracts from different plant parts of indigenous origin for their hypoglycemic responses in rabbits. *Acta Pol Pharm.* (2011) 68(6):919–25.
141. Farahbakhsh H, Pasandi Pour A, Reiahi N. Physiological response of henna (*Lawsonia inermis* L.) to salicylic acid and salinity. *Plant Prod Sci.* (2017) 20(2):237–47.
142. Zannat KE, Ahmed AU, Tanzim SM, Ahmed SM, Afrin A, Saha BC, et al. Antibacterial Effects of Henna (*Lawsonia inermis*) Leaf Extracts (Chloroform) Against Two Food Borne and Nosocomial Infection Causing Pathogens: *Staphylococcus aureus* and *Escherichia coli*. *Mymensingh Med J MMJ.* (2023) 32(1):83–9.
143. Baker RA, Tatum JH, Nemec S. Antimicrobial activity of naphthoquinones from *Fusaria*. *Mycopathologia.* (1990) 111(1):9–15.
144. Kirkland D, Marzin D. An assessment of the genotoxicity of 2-hydroxy-1,4-naphthoquinone, the natural dye ingredient of Henna. *Mutat Res Toxicol Environ Mutagen.* (2003) 537(2):183–99.
145. Kirkland DJ, Henderson L, Marzin D, Müller L, Parry JM, Speit G, et al. Testing strategies in mutagenicity and genetic toxicology: an appraisal of the guidelines of the European Scientific Committee for Cosmetics and Non-Food Products for the evaluation of hair dyes. *Mutat Res.* (2005) 588(2):88–105.
146. Zannat KE, Tanzim SM, Afrin A, Saha BC, Joynal JB, Khanam TA, et al. Antibacterial Effects of Methanolic Leaf Extracts of Henna (*Lawsonia inermis*) Against Two Most Common Pathogenic Organisms: Gram-Positive *Staphylococcus aureus* and Gram-Negative *Escherichia coli*. *Mymensingh Med J MMJ.* (2023) 32(2):296–302.
147. Abd-el-Malek YA, el-Leithy MA, Reda FA, Khalil M. Antimicrobial principles in leaves of *Lawsonia inermis* L. *Zentralblatt Bakteriell Parasitenkd Infekt Hyg Zweite Naturwissenschaftliche Abt Allg Landwirtsch Tech Mikrobiol.* (1973) 128(1):61–7.
148. Tadege H, Mohammed E, Asres K, Gebre-Mariam T. Antimicrobial activities of some selected traditional Ethiopian medicinal plants used in the treatment of skin disorders. *J Ethnopharmacol.* (2005) 100(1-2):168–75.
149. Fatahi Bafghi M, Salary S, Mirzaei F, Mahmoodian H, Meftahizade H, Zareshahi R. Antibacterial and anti-trichomonas characteristics of local landraces of *Lawsonia inermis* L. *BMC Complement Med Ther.* (2022) 22(1):203.
150. Lavari N, Ostadrahimi N, Rahimi R, Raei M, Abbassian A. The effect of a topical formulation from *Lawsonia inermis* L. (henna) on pain intensity in patients with chronic sciatica: A randomized double-blind clinical trial. *J Ethnopharmacol.* (2023) 313:116519.
151. Donkor MN, Donkor AM, Mosobil R. Combination therapy: synergism among three plant extracts against selected pathogens. *BMC Res Notes.* (2023) 16(1):83.
152. Yogisha S, Samiulla DS, Prashanth D, Padmaja R, Amit A. Trypsin inhibitory activity of *Lawsonia inermis*. *Fitoterapia.* (2002) 73(7-8):690–1.
153. Moreira CS, Silva ACJA, Novais JS, Sá Figueiredo AM, Ferreira VF, da Rocha DR, et al. Searching for a potential antibacterial lead structure against bacterial biofilms among new naphthoquinone compounds. *J Appl Microbiol.* (2017) 122(3):651–62.
154. Endrini S, AR, PI, TYYH. Anticarcinogenic Properties and Antioxidant Activity of Henna (*Lawsonia inermis*). *J Med Sci.* (2002) 2(4):194–7.
155. Li Q, Gao WQ, Zhao YQ. [Advances in studies on chemical constituents and biological activities of *Lawsonia inermis*]. *Zhongguo Zhong Yao Za Zhi Zhongguo Zhongyao Zazhi China J Chin Mater Medica.* (2013) 38(6):795–9.
156. Priya R, Ilavenil S, Kaleeswaran B, Srigopalram S, Ravikumar S. Effect of *Lawsonia inermis* on tumor expression induced by Dalton's lymphoma ascites in

- Swiss albino mice. *Saudi J Biol Sci.* (2011) 18(4):353-9.
157. Esteki R, Miraj S. The Abortifacient Effects of Hydroalcoholic Extract of *Lawsonia Inermis* on BALB/c Mice. *Electron Physician.* (2016) 8(6):2568-75.
158. Al-Assar NB, Khattak MNK, Mashwani Z ur R, Kanan S, Ullah I, Ali U, et al. Phytochemical profile and antiproliferative activities of acetone extracts of *Asplenium polypodioides* Blume. and *A. dalhousiae* Hook. in MDA-MB-231 breast cancer cells. *Saudi J Biol Sci.* (2021) 28(11):6324-31.
159. Philip JP, Madhumitha G, Mary SA. Free radical scavenging and reducing power of *Lawsonia inermis* L. seeds. *Asian Pac J Trop Med.* (2011) 4(6):457-61.
160. Ostovari A, Hoseinie SM, Peikari M, Shadizadeh SR, Hashemi SJ. Corrosion inhibition of mild steel in 1M HCl solution by henna extract: A comparative study of the inhibition by henna and its constituents (Lawson, Gallic acid, α -D-Glucose, and Tannic acid). *Corros Sci.* (2009) 51(9):1935-49.
161. Yang JY, Lee HS. Antimicrobial activities of active component isolated from *Lawsonia inermis* leaves and structure-activity relationships of its analogs against food-borne bacteria. *J Food Sci Technol.* (2015) 52(4):2446-51.
162. Cuéllar MJ, Giner RM, Recio MC, Máñez S, Ríos JL. Topical anti-inflammatory activity of some Asian medicinal plants used in dermatological disorders. *Fitoterapia.* (2001) 72(3):221-9.
163. Gad MM, Al-Sunni M, Al-Shayeb A, Al-Namsy R, Al-Naser Z, Q Khan S. The in-vitro effects of white henna addition on the *Candida albicans* adhesion and physical properties of denture base resin. *Eur Oral Res.* (2021) 55(2):86-93.
164. Okpekon T, Yolou S, Gleye C, Roblot F, Loiseau P, Bories C, et al. Antiparasitic activities of medicinal plants used in Ivory Coast. *J Ethnopharmacol.* (2004) 90(1):91-7.
165. Sarang H, Rajani P, Vasanthakumari MM, Kumara PM, Siva R, Ravikanth G, et al. An endophytic fungus, *Gibberella moniliformis* from *Lawsonia inermis* L. produces lawson, an orange-red pigment. *Antonie Van Leeuwenhoek.* (2017) 110(7):853-62.
166. Jain A, Katewa SS, Galav PK, Sharma P. Medicinal plant diversity of Sitamata wildlife sanctuary, Rajasthan, India. *J Ethnopharmacol.* (2005) 102(2):143-57.
167. Tapas A, Sakarkar D, Kakde R. Flavonoids as Nutraceuticals: A Review. *Trop J Pharm Res.* (2008) 7(3):1089-99.
168. Dhouafli Z, Ben Jannet H, Mahjoub B, Leri M, Guillard J, Saidani Tounsi M, et al. 1,2,4-trihydroxynaphthalene-2-O- β -D-glucopyranoside: A new powerful antioxidant and inhibitor of A β 42 aggregation isolated from the leaves of *Lawsonia inermis*. *Nat Prod Res.* (2019) 33(10):1406-14.
169. Sharma VK. The tuberculostatic activity of henna (*Lawsonia inermis* Linn.). *Tubercle.* (1990) 71(4):293-5.
170. Aqil F, Khan MSA, Owais M, Ahmad I. Effect of certain bioactive plant extracts on clinical isolates of β -lactamase producing methicillin-resistant *Staphylococcus aureus*. *J Basic Microbiol.* (2005) 45(2):106-14.
171. N A, Parambil RP, George N, Meethal KV. Toxicity of plant extracts containing trypsin inhibitor to the larvae of *Aedes aegypti*. *Int J Mosq Res.* (2021) 8(3):22-7.
172. Nasiou E, Giannakou IO. Nematicidal Potential of Thymol against *Meloidogyne javanica* (Treub) Chitwood. *Plants.* (2023) 12(9):1851.
173. Bari MW, Islam A, Islam MM, Sultana MJ, Afroz R, Khan MMR, et al. Determination of in vitro antioxidant activity and in vivo antineoplastic effects against Ehrlich ascites carcinoma of methanolic extract of *Sphagneticola calendulacea* (L.) Pruski. *Heliyon.* (2021) 7(6):e07228.
174. Lies Z, Janti S. Effectiveness of *Lawsonia inermis* L. Leaves Methanol Extracts on Gingivitis Healing (In vivo Study on Sprague dawley Rats). *Br J Med Med Res.* (2016) 15(2):1-8.
175. Diehl MS, Atindehou KK, Téré H, Betschart B. Prospect for anthelmintic plants in the Ivory Coast using ethnobotanical criteria. *J Ethnopharmacol.* (2004) 95(2-3):277-84.
176. Oda Y, Nakashima S, Nakamura S, Yano M, Akiyama M, Imai K, et al. New potent accelerator of neurite outgrowth from *Lawsonia inermis* flower under non-fasting condition. *J Nat Med.* (2016) 70(3): 384-90.
177. Cuong NX, Nhiem NX, Thao NP, Nam NH, Dat NT, Anh HLT, et al. Inhibitors of osteoclastogenesis from *Lawsonia inermis* leaves. *Bioorg Med Chem Lett.* (2010) 20(16):4782-4.
178. Patel JR, Tripathi P, Sharma V, Chauhan NS, Dixit VK. *Phyllanthus amarus*: Ethnomedicinal uses, phytochemistry and pharmacology: A review. *J Ethnopharmacol.* (2011) 138(2):286-313.
179. Nićiforović N, Mihailović V, Masković P, Solujić S, Stojković A, Pavlović Muratspahić D. Antioxidant activity of selected plant species; potential new sources of natural antioxidants. *Food Chem Toxicol Int J Publ Br Ind Biol Res Assoc.* (2010) 48(11):3125-30.
180. Bornet B, Branchard M. Nonanchored Inter Simple Sequence Repeat (ISSR) markers: Reproducible and

specific tools for genome fingerprinting. *Plant Mol Biol Report.* (2001) 19(3):209-15.

181. Bhuvaneswari K, Gnana PS, Kuruvilla A, Appala RB. Inhibitory concentrations of *Lawsonia inermis* dry powder for urinary pathogens. *Indian j pharmacol.* (2002) 34(4):260.

182. Ajaiyeoba EO, Onocha PA, Olarenwaju OT. In vitro Anthelmintic Properties of *Buchholzia coriacea* and *Gynandropsis gynandra* Extracts. *Pharm Biol.* (2001) 39(3):217-20.

183. Jeyathilakan N, Murali K, Anandaraj A, Abdul Basith S. In vitro evaluation of anthelmintic property of ethno-veterinary plant extracts against the liver fluke *Fasciola gigantica*. *J Parasit Dis.* (2012) 36(1):26-30.

184. Rahmany E, Çakici A, Çakir E. Antioxidant Activity and Phenolic Compounds of *Lawsonia Molecule* Extracted From *Lawsonia Inermis* (Henna). *IJFER.* (2015) 7(1):1-17.

185. Jeyaseelan EC, Jenothiny S, Pathmanathan M, Jeyadevan J. Antibacterial activity of sequentially extracted organic solvent extracts of fruits, flowers, and leaves of *Lawsonia inermis* L. from Jaffna. *Asian Pac J Trop Biomed.* (2012) 2(10):798-802.

186. Kandil HH, Al-Ghanem MM, Sarwat MA, Al-Thallab FS. Henna (*Lawsonia inermis* Linn.) inducing hemolysis among G6PD-deficient newborns. A new clinical observation. *Ann Trop Paediatr.* (1996) 16(4):287-91.

187. Singha S, Chandra G. Mosquito larvicidal activity of some common spices and vegetable waste on *Culex quinquefasciatus* and *Anopheles stephensi*. *Asian Pac J Trop Med.* (2011) 4(4):288-93.

188. Subarani S, Sabhanayakam S, Kamaraj C, Elango G, Kadir MA. Efficacy of larvicidal and pupicidal activity of *Catharanthus roseus* aqueous and solvent extracts against *Anopheles stephensi* Liston and *Culex quinquefasciatus* Say (Diptera: Culicidae). *Asian Pac J Trop Med.* (2013) 6(8):625-30.

189. Ortiz G, Terron M, Bellido J. Contact Allergy to Henna. *Int Arch Allergy Immunol.* (1997) 114(3):298-9.

190. Singh SK, Yadav RP, Singh A. Molluscicides from some common medicinal plants of eastern Uttar Pradesh, India. *J Appl Toxicol JAT.* (2010) 30(1):1-7.

191. Kamal M, Jawaid T. PHARMACOLOGICAL ACTIVITIES OF *LAWSONIA INERMIS* LINN.: A REVIEW. *Int J Biomed Res.* (2011) 1(2):37-43.

192. Ahmed S, Rahman A, Alam A, Saleem M, Athar M, Sultana S. Evaluation of the efficacy of *Lawsonia alba* in the alleviation of carbon tetrachloride-induced oxidative stress. *J Ethnopharmacol.* (2000) 69(2):157-64.

193. Bodhankar S, Jain B, Bhardwaj S, Badole S, Patel N. Antihyperglycemic activity of aqueous extract of leaves of *Cocculus hirsutus* (L.) Diels in alloxan-induced diabetic mice. *Indian J Pharmacol.* (2006) 38(1):49.

194. Singh SP, Kumar S, Mathan SV, Tomar MS, Singh RK, Verma PK, et al. Therapeutic application of *Carica papaya* leaf extract in the management of human diseases. *Daru J Fac Pharm Tehran Univ Med Sci.* (2020) 28(2):735-44.

195. Pareek A, Godavarthi A, Issarani R, Nagori BP. Antioxidant and hepatoprotective activity of *Fagonia schweinfurthii* (Hadidi) Hadidi extract in carbon tetrachloride-induced hepatotoxicity in HepG2 cell line and rats. *J Ethnopharmacol.* (2013) 150(3):973-81.

196. Loew D, Kaszkin M. Approaching the problem of bioequivalence of herbal medicinal products. *Phytother Res.* (2002) 16(8):705-11.

197. Frykberg RG, Banks J. Management of Diabetic Foot Ulcers: A Review. *Fed Pract Health Care Prof VA DoD PHS.* (2016) 33(2):16-23.

198. Simeonova R, Bratkov VM, Kondeva-Burdina M, Vitcheva V, Manov V, Krasteva I. Experimental liver protection of n-butanolic extract of *Astragalus monspessulanus* L. on carbon tetrachloride model of toxicity in rat. *Redox Rep Commun Free Radic Res.* (2015) 20(4):145-53.

199. Perinet I, Lioson E, Tichadou L, Glaizal M, de Haro L. [Hemolytic anemia after voluntary ingestion of henna (*Lawsonia inermis*) decoction by a young girl with G6PD deficiency]. *Med Trop Rev Corps Sante Colon.* (2011) 71(3):292-4.

200. Mina B, Jeevani V, Revathy S, Pramod C, Ragav R, Manjula S, Mruthunjaya K. Phytochemical and microscopical investigations on *Lawsonia inermis* roots. *Int J Current Pharm Rev Res.* (2012) 3(3):54-9.

201. Hanke ME, Talaat SM. The Biochemistry and Physiology of Henna (*Lawsonia alba*): its use as a Remedy for Intestinal Amoebiasis. *Trans R Soc Trop Med Hyg.* (1961) 55(1):56-62.

202. Chatterjee S, Datta R, Bhattacharyya D, Bandopadhyay S. Emollient and antipruritic effect of Itch cream in dermatological disorders: A randomized controlled trial. *Indian J Pharmacol.* (2005) 37(4):253.

203. Gorji A. Pharmacological treatment of headache using traditional Persian medicine. *Trends Pharmacol Sci.* (2003) 24(7):331-4.

204. Alzweiri M, Sarhan AA, Mansi K, Hudaib M, Aburjai T. Ethnopharmacological survey of medicinal herbs in Jordan, the Northern Badia region. *J Ethnopharmacol.* (2011) 137(1):27-35.

205. Marc EB, Nelly A, Annick DD, Frederic D. Plants

used as remedies antirheumatic and antineuralgic in the traditional medicine of Lebanon. J Ethnopharmacol. (2008) 120(3):315-34.

206. Takeda Y, Fatope MO. New Phenolic Glucosides from *Lawsonia inermis*. J Nat Prod. (1988) 51(4):725-9.

207. Rout GR, Das G, Samantaray S, Das P. In vitro micropropagation of *Lawsonia inermis* (Lythraceae). Rev Biol Trop. (2001) 49(3-4):957-63.

208. Oyediji AO, Ekundayo O, Koenig WA. Essential Oil Composition of *Lawsonia inermis* L. Leaves from Nigeria. J Essent Oil Res. (2005) 17(4):403-4.

209. Siddiqui BS, Kardar MN. Triterpenoids from *Lawsonia alba*. Phytochemistry. (2001) 58(8):1195-8.

210. Oyediji AO, Ekundayo O, König WA. Constituents of the Essential Oil from the Leaves of *Leonotis*

nepetaefolia (L.) Ait. f. J Essent Oil Res. (1999) 11(6):716-8.

211. Xavier MR, Santos MMS, Queiroz MG, De Lima Silva MS, Goes AJS, De Moraes Jr MA. Lawsone, a 2-hydroxy-1,4-naphthoquinone from *Lawsonia inermis* (henna), produces mitochondrial dysfunctions and triggers mitophagy in *Saccharomyces cerevisiae*. Mol Biol Rep. (2020) 47(2):1173-85.

212. Scheen AJ. Drug Treatment of Non-Insulin-Dependent Diabetes Mellitus in the 1990s: Achievements and Future Developments. Drugs. (1997) 54(3):355-68.

213. Ag TR, Kumar MS, Shivannavar CT, Gaddad SM. Antibacterial and anti-biofilm activities of crude extracts of *Lawsonia inermis* against Methicillin-Resistant *Staphylococcus aureus*. Asian J Pharm Clin Res. (2016) 9(6):263-5.

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