A Review of Genetic Taxonomy, Biomolecules Chemistry and Bioactivities of *Citrus hystrix* DC

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**Citrus hystrix DC.** with common name makrut lime or kafir lemon, is a very popular traditional medicinal plant as well as an important spice in Asiatic countries. The plant is native of the Indonesian island Sumbawa, then, it is cultivated in Indonesia, Thailand, Malaysia and the tropical region of Asia. It mainly contains essential oil and phenolic compounds. The most intense odor compounds of kafir lemon are Citronellal, L-Linalool, 1,8-Cineole, á-Terpeneol and á-Cadinene. Such as Citrusosides-A and furanocoumarines, Makrut lime content also non-volatile compounds like alkaloids and glyceroglycolipids. **Citrus hystrix** DC has many biological activities due to its volatile and nonvolatile compounds, and it has been used in traditional medicine for treating various illnesses, particularly cold pain and stomach disorder. It is also used as a juice for its fruit or as spice for its aromatic leaves. This review covers data on the chemistry and biological effects of *Citrus hystrix* DC biomolecules, and aims to lay the foundation for further study on the extraction enhancement of these biomolecules and more useful formulations.

**Keywords:** Biomolecules chemistry, Biological activities, Citrus taxonomy, Citrus hystrix, Essential oils, Phenolic compounds.

Herbs and spices are extensively used in Asiatic countries for culinary purposes and for traditional medicine. Then, several organs such as the leaves and fruits of Citrus species are widely used to flavor foods as perfumes in ceremonial celebrations. Among these Citrus spices commonly planted in small garden plots of trees, C. *hystrix* D.C., C. *aurantijolia* Swingle, C. *microcarpa* Bunge, C. *limn* Burm. and C. *mim* Merr., are by far the most economically important plant. It is believed to have originated in south-eastern Asia, in an area that includes China, India and the Indochinese peninsula and nearby archipelagos1.
The genus *Citrus* belongs to the subfamily Aurantioideae and order of Sapindales of the Rutaceae family which comprises of about 140 genera and 1,300 species distributed between 7 subfamilies. The fruits and the leaves of the *Citrus* species contain a variety biologically-active compounds including essential oils with various distinct flavors which are important to human nutrition and diet, vitamin C, folic acid, potassium, flavonoids, coumarins, pectin, and dietary fibers.

The name *Mauritius papeda*, or Kaffir lime, is widely used in the Netherlands and Germany while in France, Italy and Spain, the lime is often called ‘combava’. Later in 1824, the name “*Mauritius papeda*” was introduced to Kaffir lime by De Candolle (DC), who brought the seeds from Mauritius to his botanical garden in Montpellier in southern France. Before that time, De Candolle had studied and classified *C. hystrix* as the first species of the Papeda sub-genus.

*Genus* *Citrus* is native to tropical southeast Asia, southern China and Malaysia. It has been introduced and cultivated elsewhere in the tropics and sub-tropics including in northern Australia. However, due to its aromatic, strong, unique and spicy flavor, both fruit and leaves of *C. hystrix* are popular used ingredient in Asian cooking. They are frequently used for instance in soups, curries, or to add flavour to rice. It can also be used as an infusion for both alcoholic and non-alcoholic drinks. In addition, the leaves can be used fresh or dried, and can be stored frozen. In Thailand, the fruit is used for seasoning and to prepare drinks teas such as *Citrus hystrix* flavonoid-rich sachet, which has been promoted to have great potential as a natural antioxidant health product. The essential oil is normally produced from fresh leaves by steam distillation and serves as a source of kaffir lime leaf flavours and essences in a large variety of internationally marketed products. The main producers of kaffir lime leaves are Thailand, Indonesia, Malaysia and India. Recently, Thai growers have developed and started growing a kaffir lime without wrinkles that is easier to pack and ship around the world.

In aromatic plants, the composition of essential oils usually varies considerably because of intrinsic (sexual, seasonal, ontogenetic, and genetic variations) and extrinsic (ecological and environmental aspects) factors.

Recent studies on the Malaysian *Citrus* plants have reported the identification and composition of essential oils of several *Citrus* species including *C. aurantifolia*, *C. grandis*, *C. hystrix*, and *C. microcarpa*. Due to its high content of phenolic and flavonoid compounds, studies showed that citrus peel could be, also, potential source for natural antioxidant. In general, three types of flavonoid compounds are found on citrus peel; flavanone (e.g. hesperidin, naringin, and hesperitin), polymethoxylated flavone (e.g. nobiletin and tangeretin), and flavonol (e.g. rutin).

In Europe, North America, Asia and Australia, dried kaffir lime leaves are available in most of the marts. Usually, a bag of dried leaves can be stored in a sealed airtight container for a couple of years with little physical change. However, the quality of the dried kaffir lime leaf product depends mostly on the drying methods that are industrially employed. The leaves can be harvested all year round, especially when the trees are small. In addition to these uses, the hesperidium is still used to wash hair (as was noted in the seventeenth century by Rumphius) and, in Sri Lanka, to keep off leeches, apparently reflecting its insecticidal compounds having a rather broad range of effects on invertebrates. The leaves can be used to treat stomach ache caused by dyspepsia and insect bites; also, the rind has been prescribed for treatment for worms and headache. In Peninsular Malaysia, the fruit and leaves have been also used for washing hair; the fruit is halved and the grated rind is rubbed on the head or the whole fruit is boiled and used as shampoo; In addition, the fruit juice is used in softening the skin and the mixture of the fruit juice with bath water can be used to eliminate body odor. In traditional Medicine, *C. hystrix* is also used to treat flu, fever, hypertension, abdominal pains, and diarrhea in infants.

The fruits are used as a digestive stimulant, blood purifier, and reduce high blood pressure. In Malaysia, the oils from the fruits and the leaves of *C. hystrix DC* are commercially used as flavors and fragrances, as well as in cooking, perfumery and medical treatments, especially in aromatherapy. This review aims to summarize the main studies reporting the chemical composition of essential oils and other extracts.
from the kaffir lime leaf, the used extraction processes, and discus their main biological activities. Finally, the formulation and safety level are considered.

**Botany and plant taxonomy**

*Citrus hystrix* is generally a small tree of 3–6 m high and a width of 2.5–3 m, often not straight, crooked with glabrous and spiny branches (Fig. 1). Leaves of kaffir lime are unique among the citrus varieties, they are alternate, unifoliolate, broadly ovate to ovate-oblong, 7.5–10 cm long, dark green on top, lighter on the bottom, very fragrant with long petiole expanded into prominent wings, 15 cm long by 5 cm wide, then, each leaf comes in two parts, seemingly a double leaf (Fig. 1.c). Leaf and expanded petiole appear to be a single “pinched” leaf. Leaf base is cuneate, or rounded, apex obtuse or slightly acuminate or notched8. Flowers (Fig. 1.d) are small, fragrant, white; calyx cuspidate 4-lobed, white with violet fringe; petals 4–5, ovate-oblong, yellowish white tinged with pink; stamens 24–30 free. Fruit is large, verrucose, warty or bumpy, globose, ovoid to elliptic, green turning yellowish-green when ripe, approximately 5–7 cm diameter, rind thick, pulp yellowish, very acid and bitter with wrinkle on the surface of fruit (Fig. 1.b). Seeds are numerous, ridged, ovoid-oblong, 1.5–1.8 by 1–1.2 cm, monoembryonic with white cotyledons (Fig. 1.c)6–8.

Kaffir lime grows well in a warm subtropical or tropical climate and prefers well-drained, neutral to slightly acid soil and direct sunlight with ample moisture during the growing season4. Citrus taxonomy is still controversial due to the large degree of morphological diversity found in the group, sexual compatibility between the species and apomixes in many genotypes6. Based on morphological and phenotypic data, the two major classification systems currently used are those of Swingle and Reece (1967)6 and Tanaka et al. (2001)21 have separated eight clusters as *C. hystrix* belong to *Micrantha* cluster with *C. micrantha*, *C. macroptera* and *C. latipes*, all belonging to subgenus *Papeda*. These results place *C. hystrix* in a cluster genetically distinct from the ‘citron’ cluster as showed in Fig. 2.21,22

Studying the genetic relationships among members of the *Aurantioidae*, especially of the genus *Citrus* and based on the chloroplast matK gene sequences analysis, Penjor et al. (2013)23 confirmed the results of Nicolosi et al. (2001)21 study that *C. hystrix* belongs to the *Pummelo* cluster which differs and stands out between the *Mandarin* cluster and the *Citron* Cluster as shown in Fig. 3.

However, Liu et al. (2015)24 reported the successful development of *DArT* microarrays and their applications in phylogenetic analysis of *Citrus* species, then, genetic relationships based on neighbor-joining method show that *C. hystrix DC*, *C. macroptera Montr.*, and *C. celebica Koord* are grouped into subcluster C1 which is sister to another subcluster C2 including grapefruit (*C. paradisi Macf*), pummelo (*C. grandis (L.) Osbeck*), lime (*C. aurantifolia*), Indian wild orange (*C. indica Tan*), citron (*C. medica L.*), and mandarin (*C. reticulate*) as shown in Fig. 4.

**Chemical composition**

Based on chemical analysis, *C. hystrix DC*, is riche on bioactive molecules such as essential oils, phenolic compounds, glycerolipides and others; The main compounds of *C. hystrix DC*, will be described in the following paragraphs.

**Essential oils**

According to the International Standard Organization on Essential Oils (ISO 9235: 2013) and the European Pharmacopoeia, the term “essential oil” is reserved for a product obtained from vegetable raw material, either by distillation with water or steam, or from the epicarp of citrus fruits by a mechanical process, or by dry distillation5. Essential oils are complex mixtures of volatile to semi-volatile compounds, usually with a strong odor, rarely colored, soluble in organic solvents, and insoluble in water. They comprise volatile compounds of terpenoid and non-terpenoid origin, synthetized through different biosynthetic routes and with distinct primary metabolic precursors. In nature, essential oils can be found in various plant organs (flowers, fruits, seeds, leaves, stems, and roots), and they play very important roles in plant defense and signaling processes25, 26. Also, essential oils are used as raw materials in many fields, such as pharmaceutical, agronomic, food,
The essential oil of C. hystrix is used in aromatherapy and as essential ingredient of various cosmetic and beauty products; furthermore, the essential oil of C. hystrix has been reported to have various bioactivities such as antioxidant, antibacterial, antileukimic, and antitussive\textsuperscript{28,29}. Limonene, a monoterpene hydrocarbon, is the major component in the essential oils from the peels of the Malaysian Citrus species\textsuperscript{30,31}. In general, hydro diffusion steam distillation system, steam distillation with induction heating system, and automated steam distillation process with optimized temperature were applied to extract kaffir lime peel essential oil\textsuperscript{32,33,34}. From the fresh leaves of C. hystrix, the essential oil extracted by the steam distillation and the Likens-Nikerson extraction methods, was found to be dominated by citronellal (61.0%–73.0%), ã-citronellol (10.0%–14.0%), and limonene (5.0%–7.0%) as major components.

In addition, citronellal (72.4%), ã-citronellol (6.7%), and citronellyl acetate (4.1%) were reported to be the major components in kaffir lime leaves, followed by ã-pinene (1.9%) and limonene (0.1%) as minor components\textsuperscript{3}. Studying the chemical composition and antimicrobial activity of the essential oils from New Caledonian C. macroptera and C. hystrix from leaves, Waikedre et al. (2010)\textsuperscript{35}, reported the presence of 38 constituents. The obtained essential oils were characterized by high contents of terpinen-4-ol (13.0%), ã-pinene (10.9%), á-terpineol (7.6%), 1,8-cineole (6.4%), citronellol (6.0%) and p-cimene (5.6%), but poor in limonene (4.7%).

However, from the kaffir lime peels (from Masjid Tanah, Melaka in Malaysia), ã-pinene (39.3%), limonene (14.2%), citronellal (11.7%), and terpinen-4-ol (8.9%) were identified as the principal components. Then, ã-pinene (23.5%) and sabine (20.1%) appeared as the major components followed by citronellal (12.6%), limonene (11.8%), and ã-citronellol (3.3%) found in C. hystrix peel and reported by Nor (1999)\textsuperscript{36}.

Other study on essential oils extraction using automated steam distillation process with uncontrolled temperature carried out by Nurhani et al. (2013)\textsuperscript{37} reported that the oil composition was as follows: sabine (31.224%), ã-pinene (32.967%), limonene (20.687%), ã-pinene (3.338%), camphene (0.135%), myrcene (1.735%), á-terpineol (0.938%) and citronellal (7.531%). Other compounds were identified using the same process but in the controlled temperature like terpinolene, linalool, terpinen-4-ol and citronellol. It was also reported that the essential oil isolated from Malaysian variety of kaffir lime peel contained sabine (36.0% - 49.0%), limonene (17.0%–33.0%), citronellal (3.0%–11.0%), and ã-pinene (8.0%–14.0%) as major components\textsuperscript{36}. However, citronellal (66.9%) and á-citronellol (6.6%) were the major components essential oil in kaffir lime peel (from Selangor), obtained using the hydro-distillation method. Other research also reported that the essential oil of fresh fruit-peel is mainly consisted of monoterpene hydrocarbons, with limonene (30.73%) and ã-pinene (18.76%) as the principal components with other minor components such as terpinene-4-ol (10.63%), á-terpineol (8.35%), á-terpine (6.18%), á-terpine (5.09) and terpinolene (4.33%)\textsuperscript{38}. In other study, citronellol was found to be the major component (80.04%) in C. hystrix leaf oil; In contrast, C. hystrix fruit peel essential oil consisted of other components: limonene (40.65%), terpinen-4-ol (13.71%) and á-terpineol (13.20%)\textsuperscript{39}. Recent research of Aumeeruddy-Elalfi et al. (2016)\textsuperscript{40} found that the main compounds of essential oils from C. hystrix leaves as á-pinene (3.02%), limonene (83.89%), á-pinene (0.78%) and á-myrcene (0.89%) with traces of Methyl-eugenol (0.21%). From the study of Juraithip et al. (2010)\textsuperscript{41}, we can draw another conclusion, that C. hystrix peel and leaf showed similar patterns of essential oils chemical compositions. The major constituents of C. hystrix peel and leaf were citronellal (about 23.85–23.41%) and trace components were elemol (6.59-4.17%), á-cadinene (5.96-4.74), geranylacetate (5.12-4.45%), á-terpineol (5.15-5.40%), L-linalool (4.22-4.36%), á-pinene (1.82%) and á-humulene (1.09-0.94%). The chemical structure of these essential oils compounds are given in Fig. 5.

**Phenolic and Flavonoid compounds**

C. hystrix also function to scavenge radical activities, due to their phenolic compounds which has beneficial implications in human health, phenolic compounds (PC) are widely distributed in fruits and vegetables\textsuperscript{42}. In terms of chemical structure, phenolic compounds have at least one aromatic ring to which one or more hydroxyl
Fig. 1. Tree (a), Fruit (b), Seeds (c), Flowers (d) and Leaves (e) of *Citrus hystrix*.

Fig. 2. Analysis of RAPD and SCAR data (Left) and cpDNA markers (Right); Evidence of *C. hystrix* cluster distinction from the “Citron” cluster. 

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Fig. 3. Chloroplast matK gene sequences analysis of genus *Citrus*; Maximum likelihood tree (Left) and neighbor-joining tree (Right) showing the three clusters: Citron, Pummelo and Mandarin. *Penjor et al.* (2013)
groups are bonded to aromatic or aliphatic structures\textsuperscript{43}.

Phenolic compounds range from simple phenolic molecules to highly polymerized compounds. However, these phenolic compounds were obtained mainly in ethanolic extracts\textsuperscript{44}.

Besides their antioxidant activities, flavonoids have been demonstrated to have a wide range of biochemical and pharmacological effects including anti-inflammatory, anti-viral, anti-allergenic, anti-carcinogenic, anti-ageing activity, anti-oxidant and anti-allergic effects\textsuperscript{45,46}. Flavonoids represent the widely distributed group of plant phenolics, including the anthocyanin pigments, flavonols, flavones, flavanols, and isoflavones. The flavanols tend to polymerize to condensed tannins\textsuperscript{47}.

The group of non-flavonoids is mainly represented by benzoic and cinnamic acid known as phenolic acids\textsuperscript{44}. Flavonoids are the most common and widely distributed group of plant phenolic compounds that are characterized by a benzopyrene structure, which is ubiquitous in fruits and vegetables; and can be analyzed using colorimetric method by reaction with sodium nitrite and the development of coloured flavonoid–aluminium complex formation using aluminium chloride. The presence of polyphenolic compounds like gallic acid, hesperidin, and naringin in citrus fruits have been suggested to be responsible for the anti-diabetic activity\textsuperscript{48,49}. Interestingly, the peels of \textit{C. hystrix} have been reported to contain a variety of phenolic compounds, mainly flavanone, flavone and flavonol\textsuperscript{50}. In \textit{C. hystrix}, hesperidin is reported as component, which is responsible for radical scavenging activity\textsuperscript{51,52}.

Using supercritical carbon dioxide extraction, vanillic acid, p-coumaric acid, sinapic acid, m-coumaric acid, benzoic acid and cinnamic acid were isolated from the plant leaves\textsuperscript{53}. Three known coumarins, bergamottin, oxypeucedanin and 5-\{(6’\textprime,7’\textprime-dihydroxy-3’, 7’-dimethyl-2-octenyl)oxy\} psoralen were exhibited inhibitory activities against both lipopolysaccharide (LPS) and interferon-\(\alpha\) (IFN-\(\alpha\))-induced nitric oxide (NO) generation in RAW 264.7 cells\textsuperscript{54}. The chemical structures of these main phenolic compounds found in \textit{C. hystrix} are showed in Fig. 6\textsuperscript{55}

**Other extracts**

Two glyceroglycolipids were isolated by Murakami \textit{et al.} (1995)\textsuperscript{56}, from the leaves of \textit{C. hystrix}, and identified as 1,2-di-O-a-linolenoyl-3-O-beta-galactopyranosyl-sn-glycerol (DLGG) and a mixture of two compounds, 1-O-a-linolenoyl-2-O-palmitoyl-3-O-beta-galactopyranosyl-sn-glycerol and its counterpart (LPGG). These compounds inhibit the tumor-promoting activity of 12-O-Tetradecanoylphorbol 13-Acetate in Mouse skin. However, in addition of two coumarins (hystrixarin and hopeyhopin, an benzenoid derivatives (hystroxene-I), and an quinolinone alkaloaid (hystrolinone), as shown in Fig. 6, were isolated from the crude acetone extract of root of \textit{C. hystrix}\textsuperscript{57}.

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**Fig. 4.** Neighbor-joining method based on \textit{DArT} microarrays study of 23 Citrus species\textsuperscript{24}
Biological activities

According to the diversity of chemical compounds extract from *C. hystrix*, several works have undertaken for the assessment of some biological activities both *in vitro* or *in vivo* systems.

Antimicrobial: antibacterial and antifungal activities

Waikedre *et al.* (2010)\(^3\) have tested the leaves essential oil against three Gram positive bacteria (*Staphylococcus aureus*, *Staphylococcus epidermidis* and *Bacillus subtilis*), two Gram negative bacteria (*Klebsiella pneumonia* and *Escherichia coli*), and five fungal strains (*Aspergillus fumigates*, *Candida albicans*, *Cryptococcus neoformans*, *Saccharomyces cerevisiae* and *Trichophyton mentagrophytes*). The tested essential oil was inactive against

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**Fig. 5.** Structures of essential oils obtained from leaves and peels of *C. hystrix*
Table 1. Summary of main essential oils compounds from *C. hystrix* DC. leaves and peels as reported by previous studies

<table>
<thead>
<tr>
<th>Location and years of study</th>
<th>Main components</th>
<th>%</th>
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<th>references</th>
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<tr>
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<td>β-Pinene</td>
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<td>Malaysia, 2013</td>
<td>Peels</td>
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Bacteria but showed moderate activity against *Cryptococcus neoformans* and *Saccharomyces cerevisiae* with MIC of (50 mg/ml). This value is about tenfold lower than the used antifungal agent standard (ketoconazole: 5 mg/ml). The GC-MS of the tested essential oil was characterized by high contents of terpinen-4-ol (13.0%), α-terpineol (7.6%), 1,8-cineole (6.4%), and citronellol (6.0%). Testing the antibacterial activities of the two essential oils of makrut leaf and makrut fruit peel against 411 isolates of groups A, B, C, F, G *Streptococci, Streptococcus pneumoniae,*
Haemophilus influenzae, Staphylococcus aureus (Methicillin-Resistant and -Sensitive S. aureus) and Acinetobacter baumannii, obtained from patients with respiratory tract infections, Vimol et al. (2012) report that both essential oils were effective against all tested pathogens with minimal inhibitory concentration (MIC) ranges of 0.06–68 mg/ml and 0.03–17.40 mg/ml, respectively for leaves and fruit essential oils from C. hystrix. The GC-MS analysis of the used essential oils revealed that citronellal was found to be the major component (80.04%) in the leaf essential oil and had the lowest

Fig. 6. Structure of phenolic compounds, coumarins and a quinolinone alkaloid from C. hystrix DC
MIC. In contrast, fruits peel essential oil consisted of mixture of components (limonene 40.65%, terpinene-4-ol 13.71%, á-terpineol (13.20%), and the most active fraction was á-terpineol, followed by terpinene-4-ol, and limonene\cite{59}. The antimicrobial activities of volatile oils and extracts of eight Thailand species, \textit{C. hystrix} essential oil were investigated against eight bacteria (3 Gram positive bacteria, \textit{Bacillus subtilis} (ATCC 6051), \textit{Staphylococcus epidermidis} (ATCC 12228), \textit{S. aureus} (ATCC25923); 5 Gram negative bacteria: \textit{Escherichia coli} (ATCC 25922), \textit{Enterococcus faecalis} (ATCC 1406), \textit{Proteus mirabilis} (ATCC 14153), \textit{Pseudomonas aeruginosa} (ATCC 27853); \textit{Mycobacterium: Mycobacterium phlei} (ATCC 11758) and three fungi (\textit{Candida albicans} (ATCC 10231), \textit{C. parasilosis} (ATCC 90018) and \textit{C. tropicalis} (ATCC 13803)\cite{41}.

The volatile oil of \textit{C. hystrix} leaf, did not show any inhibitory activity against tested organisms, but interestingly, growth of \textit{Mycobacterium phlei} was inhibited by the volatiles of \textit{C. hystrix} peel with MIC of 3.5 mg/ml, with the similar patterns of essential oils in both \textit{C. hystrix} peels and leaves. This bioactive fraction composed of citronellal (about 23%) and similar trace components as L-linalool (4.22%), á-pinene (1.82%) and limonene (1.13%)\cite{41}.

It have been also reported hydrodistillation and ethyl acetate extract of \textit{C. hystrix} peels showed broad spectrum of inhibition against three Gram-positive bacteria (\textit{Staphylococcus aureus}, \textit{Bacillus cereus} and \textit{Listeria monocytogenes}), one yeast (\textit{Saccharomyces cerevisiae} var. sake) and one mold (\textit{Aspergillus fumigatus} TISTR 3180)\cite{58}. The tested ethyl acetate extracted essential oils of \textit{C. hystrix} peel had stronger antibacterial activity than the volatile obtained from hydrodistillation.

It exhibited minimum inhibitory concentration (MIC) values of 0.28 and 0.56 mg/ml against the tested yeast and \textit{B. cereus}, respectively while the minimum bactericidal concentration (MBC) values against both microbes were 0.56 mg/ml. The MIC values of the ethyl acetate extracted essential oils against \textit{L. monocytogenes}, \textit{S. aureus} and the mold were 1.13 mg/ml while the MBC values against \textit{L. monocytogenes} as well as the mold \textit{A. fumigatus} TISTR 3180 and \textit{S. aureus} were 2.25 and 1.13 mg/ml, respectively. The GC-MS analyses revealed that the major components of the ethyl acetate extracted essential oil were limonene (31.64 %), citronellal (25.96 %) and á-pinene (6.83 %) whereas á-pinene (30.48 %), sabine (22.75 %) and citronellal (15.66 %) appeared to be major compounds of the essential oil obtained by hydrodistillation\cite{58}.

\textbf{Anti-inflammatory and Antioxidant activities}

On studying anti-Inflammatory response with a model based on lipopolysaccharide-activated RAW 264.7 Murine Macrophages, Tuntipopipat \textit{et al.} (2009)\cite{59}, found that among 13 plants, the extract (with 70% ethanol) of the freeze-dried fresh leaf \textit{C. hystrix}, with extract from seven other plant, inhibited NO and TNF-a production in a dose-dependent manner without exerting cytotoxicity. Kaffer lime extract should be used in a concentration of 29.2±2.1 μg/mL and 35.4±1.5 μg/mL to reach the IC$_{50}$, respectively for the inhibition of NO Production and for TNF-a Secretion by LPS-Activated RAW 264.7 Cells.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Fig.7.png}
\caption{Structure of two glyceroglycolipids from \textit{C. hystrix} with anti-tumor promoting activity.}
\end{figure}
For antioxidant activity evaluation, based on DPPH radical scavenging capacity method, the methanolic extracts from leaf and peel of *C. hystrix*, have promising a potent antioxidant activity with IC₅₀ of 24.6 and 66.3 microg/ml respectively for leaves and peel. The antioxidant activity of fresh juice of *C. hystrix* was evaluated by employing different *in vitro* assays covering applied for the contents of total phenolics, tannins, and total flavonoids ranged that tanged respectively 836.90 mg gallic acid equivalent (GAE)/L, 507.61 mg gallic acid equivalent (GAE)/L and 224.88 mg rutin equivalent/L. Antioxidant potential based on FRAP assay show a value of 30504.40 mmol of ferrous equivalents/L juice, DPPH• and ABTS•⁺ scavaging are respectively about 10903.28 mmol of trolox equivalents/L juice, and 33830.69 mg of EDTA equivalents/L juice. Both superoxide radical and hydroxyl radical scavenging activity are in the range of 19.89 % and 42.91 %, respectively. Also, metal chelating activity reaches 7.73 mg of EDTA equivalents/L juice. These results indicated that fresh juice of *C. hystrix* could be used as a source of antioxidant agents.

**Hepatoprotective activity**

Abirami *et al.* (2015) have evaluated the hepatoprotective effects of *C. hystrix* methanolic leaf extracts on paracetamol induced toxicity in a

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**Fig. 8.** Chemical structure of citrusosides A-D; 6'-hydroxy-7'-methoxybergamottin; 6', 7'-dihydroxybergamottin and isoimperatorin and other molecules responsible of cholinesterase inhibition activity.
Swiss albino mice model. Leaf extracts were administrated at the dose of 200 mg/kg body weight for 7 days and toxicity was induced by paracetamol (2 g/kg) on day 5. Liver function markers (ALT, AST, ALP), total bilirubin and total protein in blood serums and hepatic antioxidants (SOD, CAT, GSH and GPx) in liver homogenate were estimated after that animals were sacrificed on the 7th day.

However, the recent study conducted by Abirami et al. (2015) shows that methanolic extracts of *C. hystrix* leaf possess hepatoprotective action against murine paracetamol induced hepatotoxicity; The level of enzyme markers (alanine transaminase, aspartate transaminase and alkaline phosphatase) in experimental rats were significantly restored of by the interventions *C. hystrix* leaves extract to the comparable level of normal control.

Pretreatment with *C. hystrix* extracts brought back the oxidative stress markers (superoxide dismutase, catalase and glutathione peroxidase) in the range of normal control rats. In the same study, the histopathological examination have confirmed that pretreatment with methanolic extracts of *C. hystrix* leaf in paracetamol intoxicated rats showed recovery of the hepatocytes from necrosis indicating that sample extracts preserved the structural integrity of the hepatocellular membrane and liver cell architecture damaged by paracetamol action.

**Anti-Cancer Effect**

The early study to assess the anticancer properties of methanolic extract of *C. hytrix* fresh leaves by Murakami et al. (1995) reported the plant anti-tumor properties on mouse skin in a two-stage induced by dimethylbenz[a]anthracene (DMBA) and 12-O-tetradecanoylphorbol 13-acetate (TPA). The results showed that the *IC* 50 values of two compounds were strikingly lower than those of representative used cancer preventive agents such as α-linolenic acid, beta-carotene, or (-) epigallocatechin gallate. Also, one compound exhibited anti-tumor-promoting activity was identified as 1,2-di-O-α-linolenoyl-3-O-beta-galactopyranosyl-sn-glycerol (DLGG). The second compound was identified as a mixture of 1-O-α-linolenoyl-2-O-palmitoyl-3-O-beta-galactopyranosyl-sn-glycerol and its counterpart (LPGG) as shown in Fig 7.

However, its well known that several medicinal plant exhibit anti-proliferative activity, Manosroi et al. (2006), have investigated the anti-proliferative activity of essential oil extracted from 17 Thai medicinal plants on human mouth epidermal carcinoma (KB) and murine leukemia (P388) cell lines using MTT assay. The IC50 value of both *C. hytrix* fruit and leaf, was, respectively, 0.0997 - 1.1479 mg/ml in KB cell line and 0.0746- 0.3977 in P388 cell line. Then, these two *C. hytrix* essential oils have an anti-proliferative activity on cervical cancer (KB cell) and mouse leukemia (P388 cell).

Five fractions of crude extract (hexane, ethanol, ethyl acetate, butanol and methanol) from the leaves of *C. hystrix* were investigated in vitro for their potential cytotoxic activity on 4 leukemic cell lines (HL60, K562, Molt4, U937), and normal human peripheral blood mononuclear cells (PBMCs) using the MTT assay.

The cytotoxicity bioassays showed that the ethyl acetate fraction exhibited the highest cytotoxicity, with *IC* 50 values of 19.0±0.6, 35.3±1.4, 21.8±0.4, and 19.8±1.0 ìg/ml, in response to the 4 leukemic cell lines (HL60, K562, Molt4, and U937), respectively.

These were higher than those of fractions from hexane, ethanol, and butanol. However, none of the five fractions had cytotoxic effects on PBMCs; Also, the methanol fraction did not exhibit any cytotoxic activity.

The cytotoxicity effects and apoptosis induced by three different kaffir lime leaves extract (ethanol, ethyl acetate, and hexane) to cervical cancer cell line (HeLa cells) were studied by Nastiti et al. (2013). They used cytotoxicity assay via
MTT assay, and apoptosis test with double staining method (ethidium bromide-acrydine orange). The three kaffir lime crude extract exhibited dose dependently HeLa cells proliferation inhibition. The IC<sub>50</sub> of ethanolic and ethyl acetate extract was 82,034 and 57,845 μg/mL, respectively, these two extract were able to induce apoptosis of HeLa cells by increasing the number of apoptotic cells. On the other hand hexane extract was not cytotoxic with IC<sub>50</sub> of 203,992 μg/mL. In addition, the results showed that ethyl acetate extract of kaffir lime was the most potential to induce apoptosis in HeLa cells.

The cytotoxic effect of kaffir lime leaf extracts on cervical cancer and neuroblastoma cell lines based on the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay was carried out by Woro et al. (2014).<sup>66</sup> They showed that both ethyl acetate and chloroform extracts have an IC<sub>50</sub> for HeLa cells, UKF-NB3, IMR-5 and SK-N-AS parental cells of 40.7-17.6; 28.4-18.9; 14.1-6.4 and 25.2-9.4 (μg/mL) respectively. Then, kaffir lime extract reduces the viability of cervical and neuroblastoma cell lines and may have potential as anti-cancer compounds.

**Cholinesterase inhibition activity**

Increasing communication between nerve cells that use acetyl-choline as a chemical messenger produce a therapeutic effect in patients with Alzheimer’s disease, glaucoma, myasthenia gravis, and for the recovery of neuromuscular block in surgery, then, acetylcholine breakdown in the brain can be prevented by the inhibition of acetyl cholinesterase activity, which subsequently increases the concentration of acetylcholine<sup>67</sup>. The juices of C. hystrix possess strong anti-cholinesterase activity of 79.74% against 86.89% of the used reference compound (Eserine)<sup>69</sup>. From the hexanes and dichloromethane extracts of the peels of C. hystrix fruits, Youkwan et al. (2010)<sup>68</sup> have isolated 4 new citrusosides A-D, six furanocoumarins, a sesquiterpene (eudesmane-4b,11-diol), 5 monoterpenes, and 1-O-isopropyl-beta-D-glucopyranoside. The Butyryl-cholinesterase inhibitory activity of the isolated fractions was investigated and 62,72-dihydroxybergamottin and isoimperatorin showed IC<sub>50</sub> values of 15.4+/−0.3 and 23+/−0.2 μM, respectively. These molecules are represented in Fig. 8.

In the study undertaken by Wantida et al. (2010)<sup>69</sup>, essential oils of C. hystrix were tested for their acetyl-cholinesterase (AChE) and butyryl-cholinesterase (BChE) inhibitory activities. The tested essential oil exhibited inhibitory activity on BChE higher than on AChE.

Among sixteen compounds isolated by Chonticha et al. (2016)<sup>70</sup>, from the ethyl acetate extract of the fruit peels of C. hystrix, only one flavonol compound, (3-O-beta-D-glucopyranosyl-3,5,7,4′-tetrahydroxy-6,8,3′-trimethoxyflavonol nucleus in the prenylfuranocoumarin–HMGA conjugate, Fig. 9) showed very potent butyryl-cholinesterase inhibitory activity with IC<sub>50</sub> value of 10.12 ± 0.22 μM, against galanthamine, a positive control compound with IC50 value of 11.2 ± 0.09 μM.

**Insecticidal and larvicidal activities**

The study of insecticidal properties of essential oil from Citrus hystrix DC fresh leaves against tobacco armyworm Spodoptera litura fabricius, using topical application bioassay on uniform weighted second instar larvae, demonstrated considerable repellant activity against the armyworm larvae after 24 and 48 h of treatment with LD50 values of 29.25 and 26.75 μg/mL, respectively. Also, the growth and development study in the antifeedant test showed that weight gained of larvae treated with C. hystrix essential oil were lower as compared to control treatment.

GCMS analyses of the tested essential oil revealed the presence of 29 compounds with dominance of beta-citronellal as major compound (66.85%) of total essential oil followed by beta-citronellol (6.59%), linalool (3.90%) and citronellol (1.76%). In another study, Mya et al. (2015)<sup>71</sup>, used ethanol extract of Citrus hystrix leaves to assess their larvicidal effects against Aedes aegypti which is the primary vector of dengue<sup>72</sup>; Result suggests that high concentrations of Citrus hystrix leaves ethanol extract can be used for the eradication of A. aegypti; then, concentrations of 2.4-2.1-1.8-1.5 and 1.2% of the tested leaves ethanol extract caused 99.5-85.5-62.5-26.5 and 2% mortality of
Aedes larvae in 24 hrs, respectively.

Ansori et al. (2015)\textsuperscript{73}, tested (methanol) and non-polar (n-hexane) extract fractions of C. hystrix leaves, with concentrations of 500 ppm, 1375 ppm, 2250 ppm, 3125 ppm, and 4000 ppm against the 3rd instar larvae of A. aegypti; The number of mosquito larvae mortality was calculated after 24 hours of treatment. The results reported that non-polar extract fraction is more toxic and is an effective biolarvicide with LC\textsubscript{90} = 2,885 ppm compared with polar extract fraction which has an LC\textsubscript{90} = 3,180 ppm.

Using an excito-repellency test system, Nararak et al. (2016)\textsuperscript{74}, studied the effect of essential oils of the leaf and peel of kaffir lime at four different concentrations (0.5, 1.0, 2.5, and 5.0\% v/v) for their repellency, excitation, and knockdown properties against laboratory strains of A. aegypti (L.) and Anopheles minimus Theobald at the 3–5 day aged old mosquito starved 24 h before testing.

For repellency against A. aegypti, leaf volatile oil produced the greatest response for both contact (56.1\% escape) and non-contact trials with 63.3\% escape at 2.5\%, while peel volatile oil produced the strongest response with 46.5\% escape at 2.5\%.

Against Anopheles minimus Theobald, essential oil from C. hystrix leaf had strong combined irritant and repellent activity responses at 1–5\% concentrations (90.0–96.4\% escape) and the strongest spatial repellent activity at 1\% and 2\% (85.9\% and 87.2\% escape), respectively. The peel essential oil exhibited good excitation with repellency at concentrations of 2.5\% (89.8\% escape) and 5\% (96.28\% escape), while concentrations 1–5\% showed more moderate repellent activity.

However, knockdown responses above 50\% were only observed in A. aegypti exposed to 5\% leaf essential oil. Then, the tested Kaffir lime essential oils were more active against Anopheles minimus Theobald than A. aegypti mosquitoes\textsuperscript{74}. Leaf significantly appear more active then peel essential oils at each concentration against Anopheles minimus in contact and non-contact trials, except at the highest (5\%) concentration.

**Others bioactivities: Antifertility, Tyrosinase inhibitory activity and Cardioprotective effect**

Pawinee et al.\textsuperscript{(1995)}\textsuperscript{75} investigated the effect of oral administration of both alcohol and chloroform extract of C. hystrix DC fruit peel for antifertility activity in pregnant adult female rats (Wistar) by oral administration at different periods of gestation.

They showed an enhancement of the uterotrophic effect of estradiol when both extract were simultaneously given; additionally, the extract stimulated uterine contractions. These two effects may be responsible for the interruption of pregnancy associated with the extract. Then, alcohol and chloroform extract of C. hystrix were found to effectively inhibit implantation, produce abortion and slightly hasten labor time when it was given from day 2 to 5, day 8 to 12 and day 15 until labor, respectively.

Administration of the chloroform extract in a dose of 1 g/kg produces a 62.2 f 14.5\% inhibition of implantation. However, administration of the chloroform extract at a dose of 1 g/kg twice a day from day 8 to 12 interrupted pregnancy by 91.9\%+5.5\% while the same amount of the alcohol extract produced the effect by 86.3\+9.6\%.

According to the anti-implantation effect, they founded that chloroform extract also possesses a higher abortifacient activity than the alcohol extract\textsuperscript{75}.

Tyrosinase is responsible for the formation of melanin in the human body; however, surplus expression of tyrosinase is a major problem which can lead to several skin hyperpigmentation disorders such as, seborrheic keratoses, melasma, diabetic dermopathy, tinea versicolor, melasmas disorders such as, seborrheic keratoses, melasma, diabetic dermopathy, tinea versicolor, melasmas and malignant melanomas\textsuperscript{76}. Abirami et al. (2014)\textsuperscript{60}, reported that C. hystrix juice exhibited excellent tyrosinase inhibitory activity of 80.79\%, against 90.87\% of the used reference compound (Kojic acid).

A recent study conducted by Aumeeruddy-Elalfi et al. (2016)\textsuperscript{77} showed the potency of 19 essential oils from exotic and endemic medicinal plants from Mauritius. The results tend to show that essential oils extracted from these medicinal plants can exhibit anti-tyrosinase activities and may be potential candidates for the cosmetic, food and pharmaceutical industries. Results showed that C. hystrix essential oils exhibit an IC\textsubscript{50} of 2.08 ± 0.253ig/ml. Putri et al. (2013)\textsuperscript{78}, analyzed the effects of C. hystrix peel ethanolic extract on blood serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST)
activity, and observed by light microscope the cardio-hepato-histopathology of a doxorubicin-induced cardiac and hepatic toxicity animal model (female Sprague Dawley rats). In the animal groups receiving 500 mg/kg to 1000 mg/kg C. hystrix peel ethanolic extract, cardiohistopathology profile of doxorubicin induced cardiotoxicity and hepatotoxicity rats was repaired, but neither hepatohistopathology profile was did repaired nor serum activity of aminotransferase (ALT) and aspartate aminotransferase (AST) was reduced; Thus, Putri et al. (2013)78, conclude that the ethanolic extract of C. hystrix, can be developed as cardioprotector agent.

**Safety issues**

Regarding the chemical structures and the biological activities of biomolecules from C. hystrix, and also, the several domestic uses of this plant and their extracts, it appears that a bio-safety issue for this plant may be highlighted.

No information was founded on toxicity profile of C. hystrix biomolecules in the second edition of a Guide for Health Care Professionals of Essential Oil Safety79.

From, the released draft tentative report of the Cosmetic Ingredient Review Expert Panel (May and October, 2016), entitled “Safety Assessment of Citrus Flower- and Leaf-Derived Ingredients as Used in Cosmetics”. Thus, it appear that C. hystrix leaf extract produced by extracting dried leaves with 80% ethanolic solution is reported as safe and to be non-irritating and non-sensitizing.

**CONCLUSION**

*C. hystrix* has been used as food and medicine with long history mainly in the Asian region. It is also a flavor food with health value. The present study reported phylogenetic taxonomy based on bootstrap analysis of RAPD, SCAR data, cpDNA markers and the chloroplast *matK* gene sequences analysis showed that C. hystrix belong to Pummelo cluster which is genetically distinct from the Citron cluster and the Mandarin cluster. The chemical structures of bioactives molecules explain its traditional uses and his potential to be used in cosmeceutical and pharmaceuticals.

*C. hystrix* essential oils are the mostly studied biomolecules and they have potential beneficial therapeutic actions in the management of bacterial and fungal infections. The chemical composition revealed that essential oil of leaves and fruit peel of *C. hystrix* have generally a different profile; then, among reported studies, the *C. hystrix* leaves are characterized by citronellal, α-citronellol and terpinen-4-ol as major components. However, citronellyl acetate, α-pinene, limonene, alpha-terpineol, 1,8-cineole, citronellol, p-cimene, and limonene were identified as minor components. Whereas, kaffir lime peels content respectively limonene, α-pinene, sabiniine and citronellal as major components with other minor components like terpine-4-ol, a-terpineol, g-terpinene, a-terpinene and terpinolene. Other essential oils compounds of *C. hystrix* were detected in little amount in both leaves and peels such as elemol, delta-cadinene, geranylacetate and L-linalool. The reported studies show that some of tested essential oils were inactive against bacteria. mainly those content terpinene-4-ol as major compounds. However, essential oils with citronellal as major component were more effective against bacterial strains.

In the future, more deep analysis and profiling of the volatile oils are needed to allow further elaboration of a chemotype of *C. hystrix* based on essential oils profile. Interestingly, phenol compounds of the peels of *C. hystrix* contain a variety of flavanone, flavone and flavonol; Vanillic acid, p -coumaric acid, sinapic acid, m –coumaric acid, benzoic acid and cinnamic acid were isolated from *C. hystrix* leaves. In addition, flavonoids such as cyanidin, myricetin, peonidin, quercetin, luteolin, hesperetin, apigenin and isorhamnetin, as flavanone compounds, didymin and hespiridine were isolated from both leaves and fruit juice and as flavone compounds, rutin and diosmin were isolated just from the leaves. Glyceroglycolipids, 1,2-di-O-a-linolenoyl-3-O-beta-galactopyranosyl-sn-glycerol (DLGG) and a mixture of 1-O-a-linolenoyl-2-O-palmitoyl-3-O-beta-galactopyranosyl-sn-glycerol and its counterpart (LPGG) were identified in the *C. hystrix* leaves. Benzenoid derivatives, (hystroxene-I), quinolinone alkaloid (hystrolinone) were isolated from the crude acetone extract of root of *Citrus hystrix*. 
With diversity of contents in the total phenolics, both ethanolic and methanolic extracts have promising a potent anti-inflammatory, antioxidant activity and hepatoprotective effects. Fresh leaves methanolic extract content 1,2-di-O-a-linolenoyl-3-O-beta-galactopyranosyl-sn-glycerol (DLGG) which appear an anti-tumor-promoting agent. Also, some essential oils extracted from fruit and leaves C. hystrix have an anti-proliferative activity on cervical cancer (KB cell) and mouse leukemia (P388 cell). Also, Ethyl acetate fraction of leaves of kaffir lime exhibited the highest cytoxicity activity on leukemic cell lines inducing, highest apoptosis in HeLa cells and reducing cytotoxicity activity on leukemic cell lines (P388 cell) as reported. Ethanol and chloroform extract of C. hystrix showed excellent tyrosinase inhibitory activity, cardioprotective and hepatoprotective effects on rat model. Finally, we think that the safety issues in terms of toxicity profile related to the daily use of these biomolecules should be further investigated. In addition, further studies on non-conventional extraction processes involving less solvent and energy use are also needed.

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