Journal of Applied Pharmaceutical Science Vol. 7 (05), pp. 204-218, May, 2017 Available online at http://www.japsonline.com DOI: 10.7324/JAPS.2017.70534 ISSN 2231-3354 CC BY-NC-SR

Phytochemistry and Biological Activities of the Genus *Ocotea* (Lauraceae): A Review on Recent Research Results (2000-2016)

Wan Mohd Nuzul Hakimi Wan Salleh^{1*}, Farediah Ahmad²

¹Department of Chemistry, Faculty of Science and Mathematics, Universiti Pendidikan Sultan Idris (UPSI), 35900 Tanjong Malim, Perak, Malaysia. ²Department of Chemistry, Faculty of Science, Universiti Teknologi Malaysia (UTM), 81310 Johor Bahru, Johor, Malaysia.

ARTICLE INFO

Article history: Received on: 29/12/2016 Accepted on: 11/02/2017 Available online: 30/05/2017

Key words: Phytochemistry; pharmacology; *Ocotea*; Lauraceae.

ABSTRACT

Ocotea (family: Lauracea), which comprises nearly 350 species, are distributed throughout tropical America, Africa, and Asia. Up to now, the reported constituents from the genus *Ocotea* involve neolignans, alkaloids, sesquiterpenes, flavonoids, lignans, butanolides, benzopyrans, steroids, essential oils and several other types of compounds (alkylphenols, arylpropene, coumarin, ester, saponin). Studies have shown that *Ocotea* and its active principles possess a wide range of pharmacological activities, such as anti-inflammatory, cytotoxicity, antimicrobial, larvicidal, and antiproliferative activities. The outcome of these studies will further support the therapeutic potential of the genus *Ocotea*, and provide convincing evidences to its future clinical applications in modern medicine. Thus, increasing amount of data supports application and exploitation for new drug development.

INTRODUCTION

The plant genus Ocotea, one of the largest members of the Lauraceae family, comprises approximately 350 species that are distributed throughout tropical and subtropical climates. Most species are found in America from Mexico to Argentina, seven species are found in Africa, one species is found in the Canary Islands, and about 34 recognized species are found in Madagascar (Rohwer, 2000; van der Werff, 2013). It can be recognized by the simple, alternate, stiff and aromatic elliptic to obovate leaves and fruits often borne in a cup. This family has a considerable economic importance worldwide because it is used as a source of timber for construction and furniture (Nectandra, Ocotea, Persea spp.), as a crop (Persea americana), and to obtain flavours for food industry, perfumery, and medicines (Cinnamomum zeylanicum, C. cassia) (Chaverri et al., 2011). Several plants of this genus have been used for the treatment of various diseases. Among them, the stem wood of O. bullata has been used to treat headache and male urinary tract infections

* Corresponding Author

Email: wmnhakimi @ fsmt.upsi.edu.my

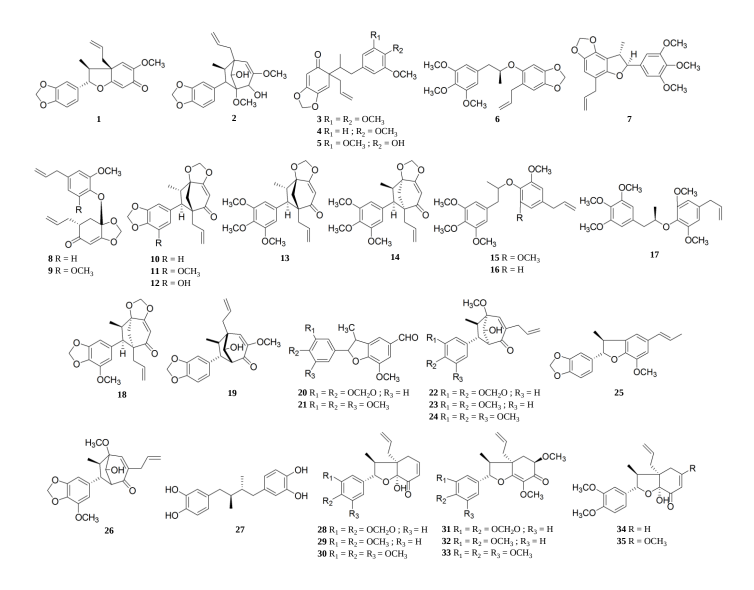
(Rakotondraibe et al., 2015). O. puchury-major is popular in local medicine, as possible sedative, gastroenteric, and antirheumatic properties. It is reported mainly for its leaves and bark, as well as cosmetic applications involving the essential oil of the leaves (Christophel et al., 1996). O. quixos is used as disinfectant, local anaesthetic and anti-diarrheic infusion (Ballabeni et al., 2007). Meanwhile, O. lancifolia is used as antiparasitic, and O. caparrapi is used to treat insect bites, snake bites, bronchitis, and cancerous tumours (Fournet et al., 2007). The woody calyces of O. bofo collected from mature fruits are traditionally used to aromatize infusions by ethnic groups. It possesses a strong anise like aroma and thus may represents a potential aniseed, fennel, or tarragon substitute or adulterant (Guerrini et al., 2006). A number of plants in the genus Ocotea are the sources of secondary metabolites, including neolignans, alkaloids, flavonoids, sesquiterpenes, butanolides, benzopyrans, steroids, alkylphenols, lignans, arylpropenes, coumarins, esters, and saponins; many of which exhibited interesting antiproliferative, antifungal, antiherpetic, antiinflammatory, and antimicrobial activities (Camargo et al., 2013; Castro et al., 2011; Cuca et al., 2009; Destryana et al., 2014; Garcez et al., 2011; Garett et al., 2012; Yamaguchi et al., 2013).

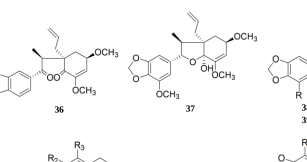
The extensive reading and investigation were actualized by systematically searching the scientific databases (PubMed, Scopus, SciFinder, and the Web of Science) for topics related to factors like the essential oils composition, chemical constituents, and pharmacological effects of the genus *Ocotea*. A bibliographic search, carried out from the year 2000 to 2016 of the genus *Ocotea* revealed that about 43 species were investigated at chemical or biological level.

Due to the ethnobotanic importance of this genus, further studies on *Ocotea* species are urgently needed. Thus, the aim of this review is to provide an overview on chemical and pharmacological studies on the essential oil, extracts, and isolated compounds from the genus *Ocotea* from year 2000 to 2016. Also included are the biological activities of compounds isolated in recent years. This should be helpful for professionals in ethnopharmacology and natural product chemistry and promote the application of plants of this genus.

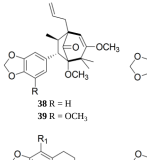
PHYTOCHEMISTRY STUDIES

The chemical constituents of *Ocotea* compounds (**Figure 1**) includeneolignans, alkaloids, sesquiterpenes, flavonoids lignans, butanolides, benzopyrans, steroids, alkylphenols, arylpropene, coumarins, ester, and saponin. Their structures are shown below, and their names and the corresponding plant sources are listed in the **Table 1.** In addition, the chemical compositions of the *Ocotea* essential oils are also discussed and summarized in **Table 2**.





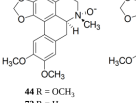
R₆



`R₃

Ĥ

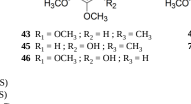


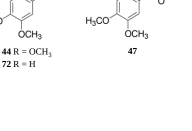


42 $R_1 = R_2 = OCH_2O$; $R_3 = H$; $R_4 = R_5 = OCH_3$; $R_6 = CH_3$; (6aS) **48** $R_1 = R_2 = OCH_2O$; $R_3 = H$; $R_4 = R_5 = OCH_2O$; $R_6 = CH_3$; (6aS) **53** $R_1 = R_2 = OCH_3$; $R_3 = H$; $R_4 = R_5 = OCH_2O$; $R_6 = CH_3$; (6a-7) **54** $R_1 = R_2 = OCH_3$; $R_3 = H$; $R_4 = R_5 = OCH_2O$; $R_6 = CH_3$; (6aS) $\textbf{55} \ \ R_1 = R_2 = OCH_3 \ ; \ R_3 = H \ ; \ R4 = R_5 = OCH_2O \ ; \ R_6 = COCH_3 \ ; \ (6aS)$ **56** $R_1 = R_2 = OCH_2O$; $R_3 = OCH_3$; $R_4 = R_5 = OCH_2O$; $R_6 = H$; (6aS) **57** $R_1 = R_2 = OCH_2O$; $R_3 = OCH_3$; $R_4 = R_5 = OCH_3$; $R_6 = CH_3$; (6a-7)

R₁

R₄





OCH

0 H₃CO

=0

OCH₃O

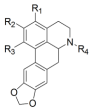
0

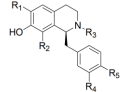
OCH

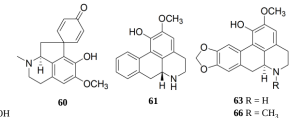
40 R = H

41 $R = OCH_3$

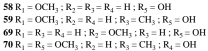
OCH₃

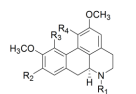




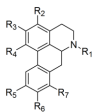


49 $R_1 = R_2 = OCH_3$; $R_3 = OH$; $R_4 = H$ 50 $R_1 = R_2 = OCH_3$; $R_3 = OH$; $R_4 = COOCH_2CH_3$ **51** $R_1 = R_2 = OCH_3$; $R_3 = OH$; $R_4 = COH$ **52** $R_1 = R_2 = OCH_3$; $R_3 = OH$; $R_4 = COOCH_3$

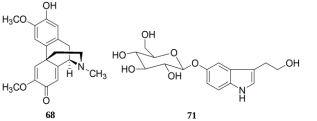


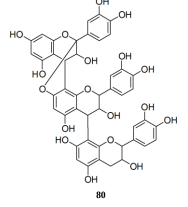


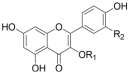
 $R_1 = R_3 = H$; $R_2 = OH$; $R_4 = OCH_3$ $R_1 = R_3 = H$; $R_2 = R_4 = OH$ $R_1 = CH_3$; $R_2 = H$; $R_3 = R_4 = OH$ $R_1 = CH_3$; $R_2 = R_4 = OH$; $R_3 = H$



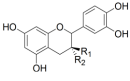
 $R_1 = CH_3$; $R_2 = R_5 = R_6 = OCH_3$; $R_3 = R_4 = OCH_2O$; $R_7 = OH$ $\mathbf{R}_1=\mathbf{R}_7=\mathbf{H}$; $\mathbf{R}_2=\mathbf{R}_5=\mathbf{R}_6=\mathbf{OCH}_3$; $\mathbf{R}_3=\mathbf{R}_4=\mathbf{OCH}_2\mathbf{O}$ $R_1 = R_2 = R_7 = H$; $R_3 = R_4 = OCH2O$; $R_5 = R_6 = OCH_3$ $R_1 = CH_3$; $R_2 = R_3 = R_4 = R_5 = R_6 = OCH_3$; $R_7 = H$ $R_1 = R_2 = R_7 = H$; $R_3 = OH$; $R_4 = OCH_3$; $R_5 = R_6 = OCH_2O$ $R_1 = CH_3$; $R_2 = R_7 = H$; $R_3 = OH$; $R_4 = R_5 = R_6 = OCH_3$ $R_1 = CH_3$; $R_2 = R_7 = H$; $R_3 = R_4 = R_5 = OCH_3$; $R_6 = OH$







81 $R_1 = Glucose$; $R_2 = OH$ 82 $R_1 = Xylose$; $R_2 = OH$ 83 R_1 = Glucuronic acid ; R_2 = OH 84 R_1 = Rhamnose ; R_2 = OH 85 $R_1 = Rhamnose$; $R_2 = H$ **88** $R_1 = H$; $R_2 = OH$ **89** $R_1 = R_2 = H$



86 $R_1 = OH$; $R_2 = H$ **87** $R_1 = H$; $R_2 = OH$

H₃CO

H₃CO

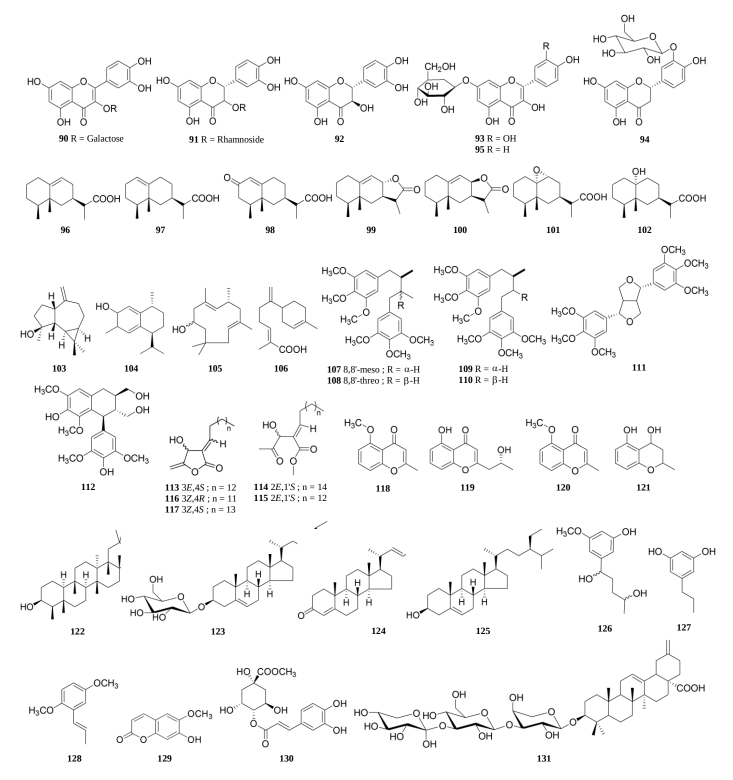


Fig. 1: Chemical structures of the compounds isolated from the genus Ocotea.

Table 1: Chemical constituents iso	ted from the genus Occ	otea (2000-2016).
------------------------------------	------------------------	-------------------

No.	Compound class and name	Source	References
	Neolignans		
1	Burchellin	O. cymbarum	Narciso et al., 2014
		O. elegans	Oliveira et al., 2006
2	Canelin	O. elegans	Oliveira et al., 2006
3	Cymosalignan A	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
4	Cymosalignan B	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
5	Cymosalignan C	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
6	$3',4'$ -Methylenedioxy-3,4,5-trimethoxy- $\Delta^{8'}$ -8.0.6'-neolignan	O. cymosa O. cymosa	Rakotondraibe <i>et al.</i> , 2015
7	Ococymosin		Rakotondraibe <i>et al.</i> , 2015
8	Didymochlaenone B	O. cymosa O. cymosa	
		~	Rakotondraibe <i>et al.</i> , 2015
9	Didymochlaenone C	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
10	Sibyllenone	O. bullata	Zschocke <i>et al.</i> , 2000
11	Demethoxysibyllenone	O. cymosa	Rakotondraibe et al., 2015
12	5-O-Demethylsibyllenone	O. cymosa	Rakotondraibe et al., 2015
13	$(7R, 8S, 1'S, 3'S) - \Delta^{8'} - 3, 4, 5$ -Trimethoxy-3',4'-methylenedioxy-1',2',3',6'-tetrahydro-	O. cymosa	Rakotondraibe et al., 2015
	6'-oxo-7.1'-8.3'-neolignan		
14	$(7R, 8R, 1'R, 3'R)$ - $\Delta^{8'}$ - $3, 4, 5$ -Trimethoxy- $3', 4'$ -methylenedioxy- $1', 2', 3', 6'$ -tetrahydro-	O. cymosa	Rakotondraibe et al., 2015
	6'-oxo-7.1'-8.3'-neolignan	-	
15	3,4,5,3',5'-Pentamethoxy-1'-allyl-8.O.4'-neolignan	O. cymosa	Rakotondraibe et al., 2015
16	3,4,5,3'-Tetramethoxy-1'-allyl-8.O.4'-neolignan	O. cymosa	Rakotondraibe et al., 2015
17	Virolongin B	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
18	Ocobullenone	O. cymosa	Rakotondraibe <i>et al.</i> , 2015
10		O. bullata	Zschocke <i>et al.</i> , 2000
10	2'-Epiguianin	O. macrophylla	Suárez <i>et al.</i> , 2011
19	2 -Epigutatili		
		O. macrophylla	Coy-Barerra <i>et al.</i> , 2009
20	Ocophyllal A	O. macrophylla	Coy-Barerra et al., 2009
		O. macrophylla	Suárez et al., 2011
21	Ocophyllal B	O. macrophylla	Coy-Barerra et al., 2009
		O. macrophylla	Suárez et al., 2011
22	Ocophyllol A	O. macrophylla	Coy-Barerra et al., 2009
	1 V	O. macrophylla	Suárez et al., 2011
23	Ocophyllol B	O. macrophylla	Coy-Barerra et al., 2009
	FJ	O. macrophylla	Suárez <i>et al.</i> , 2011
24	Ocophyllol C	O. macrophylla	Coy-Barerra <i>et al.</i> , 2009
24 25	(+)-Licarin B		Coy-Barerra <i>et al.</i> , 2009
	rel (75,8 <i>R</i> ,1'5,2' <i>R</i> ,3'S)- $\Delta^{8'}$ -2'-hydroxy-5,1',3'-trimethoxy-3,4-methylenedioxy-7,3',	O. macrophylla	
26		O. heterochroma	Cuca et al., 2009
	8,1'-neolignan		
27	meso-Dehydroguaiaretic acid	O. heterochroma	Cuca <i>et al.</i> , 2009
28	Ferrearin C	O. catharinensis	Funasaki <i>et al.</i> , 2009
29	Ferrearin E	O. catharinensis	Funasaki et al., 2009
30	Ferrearin G	O. catharinensis	Funasaki et al., 2009
31	Armenin B	O. catharinensis	Funasaki et al., 2009
32	5'-Methoxyporosin	O. catharinensis	Funasaki et al., 2009
33	$(75,85,1'R,3'R)$ -3,4,5,3',5'-Pentamethoxy-4'-oxo- $\Delta^{1,3,5,5',8'}$ -8.1',7.0.6'-neolignan	O. catharinensis	Funasaki <i>et al.</i> , 2009
34	<i>rel-</i> (75,85,1' <i>R</i> ,2' <i>S</i>)-2'-Hydroxy-3,4-dimethoxy-3'-oxo-Δ ^{1,3,5,4',8'} -8.1',7.0.2'-	O. catharinensis	Funasaki <i>et al.</i> , 2009
54	neolignan	C. cumun mensis	i unusuki (<i>i ui.</i> , 2007
35	<i>rel</i> - $(7R,8S,1'R,2'S)-2'$ -Hydroxy-3,4,5'-trimethoxy-3'-oxo- $\Delta^{1,3,5,4',8'}-8.1',7.0.2'$ -	O. catharinensis	Funasaki et al., 2009
33		5. cumur mensis	1 ullasaki ei ul., 2009
20	neolignan	O antheni	Europelri et al. 2000
36	<i>rel</i> - $(8S, 1'R, 5'R)$ -3,4,3',5'-Tetramethoxy-7,2'-dioxo- $\Delta^{1,3,5,3',8'}$ -8.1'-neolignan	O. catharinensis	Funasaki <i>et al.</i> , 2009
37	<i>rel</i> $(7R,8S,1'R,2'S)$ -2'-Hydroxy-3,4-methylenedioxy-5,3',5'-trimethoxy- $\Delta^{1,3,5,3',8'}$ -	O. catharinensis	Funasaki et al., 2009
	8.1',7.O.2'-neolignan		
38	rel- (7S,8R,1'R,3'R)-4'-Hydroxy-3,4-methylenedioxy-3',5'-dimethoxy-2',4'-dioxo-	O. catharinensis	Funasaki et al., 2009
	$\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan		
39	(7S,8R,1'R,3'R)-4'-Hydroxy-3,4-methylenedioxy-3',5',5-trimethoxy-2',4'-dioxo-	O. catharinensis	Funasaki et al., 2009
	$\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan		,,
40	<i>rel-</i> (75,8 <i>R</i> ,1′ <i>R</i> ,3′ <i>R</i>)-4′-Hydroxy-3,4-methylenedioxy-3′,5′-dimethoxy-2′,4′-dioxo-	O. catharinensis	Funasaki et al., 2009
-10	$\Delta^{1,3,5,5,8,6}$ -8.1',7.3'-neolignan dimer	5. cumur mensis	1 unusaki t <i>i ut.</i> , 2007
41	(7 <i>S</i> ,8 <i>R</i> ,1' <i>R</i> ,3' <i>R</i>)-4'-Hydroxy-3,4-methylenedioxy-3',5',5-trimethoxy-2',4'-dioxo-	0 aathanin main	Europeki et al. 2000
41		O. catharinensis	Funasaki et al., 2009
	$\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan dimer		
	Alkaloids		
42	(+)-Dicentrine	O. puberula	Montrucchio et al., 2012

	Alkaloids		
42	(+)-Dicentrine	O. puberula	Montrucchio et al., 2012
		O. macrophylla	Coy-Barrera and Cuca-Suárez, 2009
		O. acutifolia	Garcez et al., 2011; Guterres et al., 2013
43	(+)-Ocoteine	O. acutifolia	Garcez et al., 2011; Guterres et al., 2013
44	(+)-6S-Ocoteine N-oxide	O. acutifolia	Garcez et al., 2011; Guterres et al., 2013
45	(+)-Leucoxine	O. acutifolia	Garcez et al., 2011; Guterres et al., 2013

46	(+)-Norocoxylonine	O. acutifolia	Garcez et al., 2011
47	(+)-Thalicminine	O. acutifolia	Garcez et al., 2011; Guterres et al., 2012
48	(+)-Neolitsine	O. acutifolia	Garcez et al., 2011; Guterres et al., 2012
		O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
19	S-3-methoxy-nordomesticine	O. macrophylla	Pabon and Cuca 2010
50	S-N-ethoxycarbonyl-3-methoxy-nordomesticine	O. macrophylla	Pabon and Cuca 2010
51	S-N-formyl-3-methoxy-nordomesticine	O. macrophylla	Pabon and Cuca 2010
52	S-N-methoxycarbonyl-3-methoxy-nordomesticine	O. macrophylla	Pabon and Cuca 2010
53	Dehydronantenine	O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
54	(+)-Nantenine	O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
55	(+)-N-acetyl-nornantenine	O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
56	(+)-Cassythidine	O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
57	Didehydroocoteine	O. macrophylla	Coy-Barrera and Cuca-Suárez 2009
58	Coclaurine	O. lancifolia	Fournet <i>et al.</i> , 2007
		O. duckei	Silva et al., 2002
59	(-)-N-Methycoclaurine	O. lancifolia	Fournet <i>et al.</i> , 2007
NMS	Crostparine	O. lancifolia	Fournet et al., 2007
50	Glaziovine	O. lancifolia	Fournet <i>et al.</i> , 2007
61	(-)-Caaverine	O. lancifolia	Fournet <i>et al.</i> , 2007
62	(+)-Laurotetanine	O. lancifolia	Fournet <i>et al.</i> , 2007
6 3	(+)-Nordomesticine	O. lancifolia	Fournet <i>et al.</i> , 2007
53 54	(+)-Norisoboldine	O. lancifolia	Fournet <i>et al.</i> , 2007
NMS	(+)-Norantenine	O. lancifolia	Fournet <i>et al.</i> , 2007
65	(+)-Corytuberine	O. lancifolia	Fournet $et al., 2007$
66	(+)-Domesticine	O. lancifolia	Fournet <i>et al.</i> , 2007
67	(+)-Isoboldine	O. lancifolia	Fournet <i>et al.</i> , 2007
68	(S)-Pallidine	O. lancifolia	Fournet <i>et al.</i> , 2007
69	(+)-Norjuziphine	O. lancifolia	Fournet <i>et al.</i> , 2000
70	(+)-Reticuline	O. lancifolia	Fournet <i>et al.</i> , 2007
71	Tryptophol-5- <i>O</i> -β-D-glucopyranoside	O. minarum	Garcez <i>et al.</i> , 2005
NMS	Lequesnamine	O. leucoxylon	Imler <i>et al.</i> , 2003
72	(+)-6S-Dicentrine N-oxide	O. acutifolia	Garcez <i>et al.</i> , 2011
73	(+)-Oxocylonine	O. acutifolia	Garcez <i>et al.</i> , 2011
74	(+)- <i>O</i> -Methylcassyfiline	O. acutifolia	Garcez <i>et al.</i> , 2011
75	(+)-Nordicentrine	O. acutifolia	Garcez <i>et al.</i> , 2011
76	(+)-Thalicsimidine	O. acutifolia	Garcez <i>et al.</i> , 2011
77	(+)-Isodomesticine	O. acutifolia	Garcez <i>et al.</i> , 2011
78	(+)-Predicentrine	O. acutifolia	Garcez <i>et al.</i> , 2011
79	(+)- <i>N</i> -Methyllaurotethanine	O. acutifolia	Garcez et al., 2011
79	(+)-N-Methyllaurotethanine Flavonoids	O. acutifolia	Garcez et al., 2011
	Flavonoids	O. acutifolia O. notata	Garcez et al., 2011 Garett et al., 2012
30	<i>Flavonoids</i> A-type proanthocyanidin trimer	O. notata	Garett et al., 2012
30	Flavonoids		
30 31	<i>Flavonoids</i> A-type proanthocyanidin trimer Isoquercitrin	O. notata O. notata O. corymbosa	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010
30 31	<i>Flavonoids</i> A-type proanthocyanidin trimer	O. notata O. notata O. corymbosa O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012
80 81 82	<i>Flavonoids</i> A-type proanthocyanidin trimer Isoquercitrin Reynoutrin	O. notata O. notata O. corymbosa	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010
80 81 82 83	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012
80 31 32 33 34	<i>Flavonoids</i> A-type proanthocyanidin trimer Isoquercitrin Reynoutrin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010
80 31 32 33 34 35	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012
80 81 82 83 84 85 86	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012
80 81 82 83 84 85 86 87	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012
80 81 82 83 84 85 86 87 88	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin	O. notata O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012
30 31 32 33 34 35 36 37 38 39	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol	O. notata O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012
30 31 32 33 34 35 36 37 38 39 90	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol Quercetin-3-O-β-D-galactoside	O. notata O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010
30 31 32 33 34 35 36 37 38 39 90 90	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol Quercetin-3- O - β -D-galactoside rel - ($2R$, $3R$)-dihydroquercetin-3- O - α -L-rhamnoside (astilbin)	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010
80 81 82 83 84 85 86 87 88 88 89 90 91 92	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol Quercetin-3- O - β -D-galactoside rel - $(2R,3R)$ -dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010
30 31 32 33 34 35 36 37 38 88 39 90 91 92 93	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol Quercetin-3-O- β -D-galactoside rel- (2R,3R)-dihydroquercetin-3-O- α -L-rhamnoside (astilbin) Taxifolin Quercetin-7-O- β -D-glucopyranoside	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010 Garecz <i>et al.</i> , 2005 Garecz <i>et al.</i> , 2005
30 31 32 33 34 35 36 37 38 37 38 39 90 90 91 92 29 3 94	Flavonoids A-type proanthocyanidin trimer Isoquercitrin Reynoutrin Miquelianin Quercitrin Afzelin Catechin Epicatechin Quercetin Kaempferol Quercetin-3- O - β -D-galactoside rel - $(2R,3R)$ -dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010
30 31 32 33 34 35 36 37 38 37 38 39 90 90 91 92 29 3 94	FlavonoidsA-type proanthocyanidin trimer IsoquercitrinReynoutrinMiquelianin QuercitrinAfzelinCatechinEpicatechin Quercetin Kaempferol Quercetin-3- O - β -D-galactoside $rel-(2R,3R)$ -dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin Quercetin-7- O - β -D-glucopyranoside Eriodictyol-3'- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranoside	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum O. minarum	Garett et al., 2012 Garett et al., 2012 Batista et al., 2010 Garett et al., 2010 Garett et al., 2012 Batista et al., 2012 Garett et al., 2011 Batista et al., 2010 Batista et al., 2010 Garcez et al., 2005 Garcez et al., 2005 Garcez et al., 2005
79 80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 995 96	FlavonoidsA-type proanthocyanidin trimer IsoquercitrinReynoutrinMiquelianin QuercitrinAfzelinCatechinEpicatechin QuercetinQuercetin Kaempferol Quercetin-3- O - β -D-galactoside $rel-(2R,3R)$ -dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin Quercetin-7- O - β -D-glucopyranoside Eriodictyol-3'- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranosideSesquiterpenes	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum O. minarum	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005
80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96	FlavonoidsA-type proanthocyanidin trimer IsoquercitrinReynoutrinMiquelianin QuercitrinAfzelinCatechinEpicatechin QuercetinQuercetin Kaempferol Quercetin-3- O - β -D-galactoside rel- (2R,3R)-dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin Quercetin-7- O - β -D-glucopyranoside Eriodictyol-3'- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranosideSesquiterpenes rel-4 β ,5 β ,7 β -eremophil-9-en-12-oic acid	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum O. minarum O. minarum	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005
80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96 97	FlavonoidsA-type proanthocyanidin trimer IsoquercitrinReynoutrinMiquelianin QuercitrinAfzelinCatechin Epicatechin Quercetin-3- O - β -D-galactoside rel- (2R,3R)-dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin Quercetin-7- O - β -D-glucopyranoside Eriodictyol-3'- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranosideSesquiterpenesrel-4 β ,5 β ,7 β -eremophil-9-en-12-oic acid rel-4 β ,5 β ,7 β -eremophil-1 (10)-en-12-oic acid	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum O. minarum O. minarum O. minarum	Garett et al., 2012 Garett et al., 2012 Batista et al., 2010 Garett et al., 2012 Batista et al., 2012 Batista et al., 2012 Garett et al., 2011 Batista et al., 2010 Batista et al., 2010 Garez et al., 2005 Garez et al., 2013
80 81 82 83 84 85 86 87 88 89 90 91 92 93 94 95 96	FlavonoidsA-type proanthocyanidin trimer IsoquercitrinReynoutrinMiquelianin QuercitrinAfzelinCatechinEpicatechin QuercetinQuercetin Kaempferol Quercetin-3- O - β -D-galactoside rel- (2R,3R)-dihydroquercetin-3- O - α -L-rhamnoside (astilbin) Taxifolin Quercetin-7- O - β -D-glucopyranoside Eriodictyol-3'- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranoside Naringenin-7- O - β -D-glucopyranosideSesquiterpenes rel-4 β ,5 β ,7 β -eremophil-9-en-12-oic acid	O. notata O. notata O. corymbosa O. notata O. corymbosa O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. notata O. acutifolia O. corymbosa O. elegans O. minarum O. minarum O. minarum O. minarum	Garett <i>et al.</i> , 2012 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2010 Garett <i>et al.</i> , 2010 Garett <i>et al.</i> , 2012 Batista <i>et al.</i> , 2012 Garett <i>et al.</i> , 2011 Batista <i>et al.</i> , 2010 Batista <i>et al.</i> , 2010 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005 Garcez <i>et al.</i> , 2005

101	<i>rel</i> -4β,5β,7β-eremophil-9α,10α-epoxy-12-oic acid	O. lancifolia	Camargo et al., 2013
102	$4\beta,5\beta,7\beta$ -eremophil-11-en- 10 α -ol	O. lancifolia	Camargo et al., 2013
103	Spathulenol	O. lancifolia	Camargo et al., 2013
104	rel- (1R, 4S)-7-hydroxycalamenene	O. elegans	Batista et al., 2010
105	rel- (8R)-Humulan-1,4-dien-8-ol	O. catharinensis	Funasaki et al., 2009
106	Lanceolic acid	O. minarum	Garcez et al., 2005
	Lignans		
107	meso-3,4,5,3',4',5'-Hexamethoxy-8.8'-lignan	O. macrophylla	Coy-Barerra et al., 2011
108	threo-3,4,5,3',4',5'-Hexamethoxy-8.8'-lignan	O. macrophylla	Coy-Barerra et al., 2011
109	erythro-Diarylbutane	O. macrophylla	Suárez et al., 2011
110	threo-Diarylbutane	O. macrophylla	Suárez et al., 2011
111	Yangambin	O. heterochroma	Cuca et al., 2009
		O. duckei	Neto et al., 2007; 2008
112	Lyonyresinol	O. minarum	Garcez et al., 2005
	Butanolides		
113	Macrocarpolide A	O. macrocarpa	Liu et al., 2015
114	Macrocarpolides B	O. macrocarpa	Liu et al., 2015
115	Macrocarpolides C	O. macrocarpa	Liu et al., 2015
116	Linderanolide B	O. macrocarpa	Liu et al., 2015
117	Isolinderanolide	O. macrocarpa	Liu et al., 2015
	Benzopyrans		
118	2-Methyl-5-methoxy-benzopyran-4-one	O. corymbosa	Teles et al., 2005
119	(2'S)-2- (propan-2'-ol)-5-hydroxy-benzopyran-4-one	O. corymbosa	Teles <i>et al.</i> , 2005
120	(2R)-2,3-dihydro-2-methyl-5-methoxy-benzopyran-4-one	O. corymbosa	Teles <i>et al.</i> , 2005
121	2,3-Dihydro-2-methyl-benzopyran-4,5-diol	O. corymbosa	Teles <i>et al.</i> , 2005
	Steroids		
122	β-Friedelanol	O. heterochroma	Cuca et al., 2009
122	$3-O-\beta$ -D-glucopyranosyl stigmasterol	O. minarum	Garcez <i>et al.</i> , 2005
123	Stigmasta-4,22-dien-3-one	O. minarum	Garcez <i>et al.</i> , 2005
125	β-Sitosterol	O. minarum	Garcez <i>et al.</i> , 2005
	Alkylphenols		
126	3- (1,4-Dihydroxypentyl)-5-methoxyphenol	O. minarum	Garcez et al., 2005
120	5-Propylresorcinol	O. minarum O. minarum	Garcez <i>et al.</i> , 2005
	Arylpropene		
128	trans-Asarone	O. minarum	Garcez et al., 2000
120		0. minur um	Jailez ei ui., 2000
	Coumarin	<u> </u>	
129	Scopoletin	O. minarum	Garcez et al., 2005
	Ester		
130	Ester 4-O-E-caffeoylquinic acid methyl ester	O. corymbosa	Batista et al., 2010
130		O. corymbosa	Batista <i>et al.</i> , 2010

 $\label{eq:NMS-no-molecular} NMS- no molecular structure provided.$

Species	Locality	Parts/Major components	References
D. odorifera	Brazil Brazil	Leaves: Safrole (92.0%) Leaves: Camphor (43.0%), safrole (42.0%)	Oltramari <i>et al.</i> , 2004 Mossi <i>et al.</i> , 2014
). quixos	Italy	Leaves: 1,8-Cineole (8.2%), sabinene (4.6%), α-pinene (4.3%)	Enrico et al., 2014
-	USA	Leaves: <i>trans</i> -Caryophyllene (28.2%), methyl cinnamate (19.5%), β -selinene (10.4%), α -humulene (10.1%)	Destryana et al., 2014
	Ecuador	Leaves: trans-Cinnamaldehyde (27.8%), methyl cinnamate (21.6%)	Tognolini et al., 2006
	Ecuador	Leaves: β -Caryophyllene (15.1%), cinnamyl acetate (11.4%), sabinene (7.6%), geranial (5.6%)	Sacchetti et al., 2006
	Ecuador	Flower calices: <i>trans</i> -Cinnamaldehyde (27.9%), methylcinnamate (21.6%), 1,8-cineole (8.0%)	Bruni et al., 2003
). nigrescens	Brazil	Leaves: β-Caryophyllene (37.9%), α-pinene (6.7%), β-pinene (6.9%), α-copaene (6.2%)	Yamaguchi et al., 2013
O. splendens	Brazil	Leaves: β -Caryophyllene (51.0%), caryophyllene oxide (9.9%), α -humulene (6.2%)	Yamaguchi et al., 2013
). puchury-major	Brazil	Leaves: Safrol (39.4%), eucaliptol (28%), sabinene (8.5%), α-terpineol (7.9%)	Leporatti et al., 2014
). macrophylla	Brazil	Leaves: Germacrene A (22.7 %), β -caryophyllene (22.9 %), α -pinene (8.7%), β -pinene (6.9%)	Garrett <i>et al.</i> , 2010
	Colombia	Leaves: Spathulenol (15.9%), γ -muurolene (15.4%), bicyclogermacrene (14.5%)	Prieton et al., 2010
D. gomezii	Costa Rica	Leaves: Spannenou (13.5%), γ -inducible (13.4%), bicyclogennactene (14.5%) Leaves: Pentan-2-ol (12.5%), epi- α -cadinol (9.8%), δ -cadinene (7.7%), 1,8-cineole (6.0%)	Chaverri <i>et al.</i> , 2011
		Bark: δ-Cadinene (14.5%), 1,10-diepi-cubenol (7.7%), α-muurolene (6.9%)	Chaverri et al., 2011
		Wood: epi- α -Muurolol (15.0%), epi- α -cadinol (10.0%), δ -cadinene (7.7%)	Chaverri et al., 2011
O. morae	Costa Rica	Leaves: β -Pinene (17.5%), α -pinene (10.4%), bicyclogermacrene (8.8%), germacrene	Chaverri <i>et al.</i> , 2011
7. morue	costa Mica	D (7.5%), 1,8-cineole (7.3%), β -caryophyllene (7.1%)	Chaven11 et al., 2011
		Bark: 1,8-Cineole (12.8%), β-caryophyllene (6.1%)	Chaverri et al., 2011
)	D	Wood: (<i>E</i>)-Nerolidol (11.4%), 1,8-cineole (7.1%), epi- α -muurolol (6.3%), δ -cadinene (6.2%), α -cadinol (6.0%)	Chaverri <i>et al.</i> , 2011
). puberula	Brazil	Bark: Spathulenol, β -pinene, bicyclogermacrene, germacrene D and α -pinene	Farago <i>et al.</i> , 2010
). longifolia	Colombia	Leaves: α-Terpinolene (80.9%), α-phellandrene (4.7%)	Prieto et al., 2010
). sp.	Colombia	Stem/leaves/flowers: α-Pinene (42.0%), <i>p</i> -cymene (14.6%), β-pinene (12.7%)	Olivero et al., 2010
). duckei	Brazil	Roots: Elemol (24.3%), β -elemene (16.6%), β -eudesmol (13.4%), (%)	Barbosa-Filho et al., 2008
		Stems: β-Eudesmol (27.5%), α-pinene (9.0%), dl-limonene (6.6%), 1-borneol (6.1%) Leaves: <i>trans</i> -Caryophyllene (60.5%), α-humulene (4.6%), δ-selinene (4.4%) Fruits: dl-Limonene (30.1%), β-pinene (12.2%), α-pinene (9.8%), epiglobulol (8.1%)	Barbosa-Filho <i>et al.</i> , 2008 Barbosa-Filho <i>et al.</i> , 2008 Barbosa-Filho <i>et al.</i> , 2008
D. endresiana D. praetermissa	Costa Rica Costa Rica	Leaves: α-Pinene (47.9%), β-pinene (21.0%), α-humulene (14.3%) Leaves: α-Pinene (30.1%), β-pinene (19.0%), (<i>E</i>)-caryophyllene (9.1%), limonene (9.0%)	Agius <i>et al.</i> , 2007 Agius <i>et al.</i> , 2007
O. veraguensis	Costa Rica	(9.0%) Leaves: Bulnesol (29.5%), p -cymene (19.8%), spathulenol (8.5%)	Moriarity <i>et al.</i> , 2007; Takaku <i>et al.</i> , 2007
D. whitei	Costa Rica	Leaves: Spathulenol (15.3%), β -caryophyllene (15.2%), (<i>E</i> , <i>E</i>)-farnesyl acetate (10.1%)	Moriarity <i>et al.</i> , 2007; Takaku <i>et al.</i> , 2007;
O. meziana	Costa Rica	Leaves: Germacrene D (50.6%), β -caryophyllene (13.2%), δ -cadinene (8.0%)	Moriarity <i>et al.</i> , 2007; Takaku <i>et al.</i> , 2007;
D. sp. nov. ''los lanos''	Costa Rica	Leaves: α-Pinene (27.5%), β -pinene (17.2%)	Wright <i>et al.</i> , 2007
D. sp. nov. ''small eaf''	Costa Rica	Leaves: Germacrene D (60.4%), α-pinene (4.3%)	Wright et al., 2007
O. floribunda	Costa Rica	Leaves: α-Pinene (22.5%), β -pinene (21.3%)	Takaku <i>et al.</i> , 2007; Werk <i>et al.</i> , 2007
O. bracteosa	Brazil	Stem bark: δ-Cadinene (12.4%), ledene (11.1%), globulol (10.1%)	Coutinho et al., 2007
O. austinii	Costa Rica	Leaves: α -Pinene (33.2%), β -pinene (13.0%) and δ -cadinene (5.7%) Twig wood: α -Pinene (14.9%), β -pinene (8.2%), β -eudesmol (9.1%), α -eudesmol	Chaverri <i>et al.</i> , 2007 Chaverri <i>et al.</i> , 2007
O. tonduzii	Costa Rica	(8.8%) Leaves: α-Pinene (41.4%), β-pinene (25.1%), α-humulene (6.9%), β-caryophyllene (5.8%)	Takaku <i>et al.</i> , 2007; Bans <i>et al.</i> , 2007
). holdridgeana	Costa Rica	Leaves: α-Pinene (29.9%), germacrene D (19.9%), <i>trans</i> -2-hexenal (8.4%)	Takaku <i>et al.</i> , 2007
). sinuate	Costa Rica	Leaves: trans-2-hexenal (17.0%), camphene (16.2%), germacrene D (15.4%), α -pinene (15.0%)	Takaku <i>et al.</i> , 2007 Takaku <i>et al.</i> , 2007
D. valeriana	Costa Rica USA	Leaves: Germacrene D (69.7%), α-cadinol (4.5%) Leaves: 1,4-Cineole (19.6%), (Z)-anethole (13.4%), β-pinene (6.8%), α-gurjunene (6.5%)	Takaku <i>et al.</i> , 2007 Scora and Scora, 2001
O. bofo O. foetens	Ecuador Portugal	Floral calyces: Stragole (48.7%), <i>R</i> -phellandrene (19.6%), sabinene (10.4%) Aerial parts: <i>p</i> -Coumarate (69.6%), γ -muurolene (5.2%), β -caryophyllene (4.9%)	Guerrini <i>et al.</i> , 2006 Pino <i>et al.</i> , 2004
O. comoriensis O. opifera	Africa Bolivia	Bark: Camphene (18.1%), α-pinene (13.7%), bornyl acetate (13.8%) Stem bark: Asaricin (84.6%)	Menut <i>et al.</i> , 2002 Lorenzo <i>et al.</i> , 2001
0. botrantha	USA	Leaves: Germacrene D (35.2%), β -caryophyllene (13.4%), δ -elemene (11.2%)	Scora and Scora, 2001

Table 2: Chemical components identified in the essential oils of the genus Ocotea (2000-2016).

Neolignans

Forty one neolignans were isolated from eight Ocotea species. Rakotondraibe et al. (2015)successively isolated ten new neolignans including the 6'-oxo-8.1'-lignans, cymosalignan A-C (3-5), 8.O.6'-neolignan (6), ococymosin (7), didymochlaenone C (9), and the bicyclo[3.2.1]octanoids (11-14) were isolated along with the known compounds, didymochlaenone B (8), 3,4,5,3',5'pentamethoxy-1'-allyl-8.O.4'-neolignan (15), 3.4.5.3'tetramethoxy-1'-allyl-8.O.4'-neolignan (16), virolongin B (17), ocobullenone (18), and sibyllenone (10) from the stems or bark of the Madagascan plant O. cymosa. Compound (10) and (18) were also identified from the stem bark of O. bullata (Zschocke et al., 2000). Phytochemical exploration of the leaves of O. macrophylla afforded two new di-nor-benzofuran neolignan aldehydes, ocophyllals (20-21), and three new macrophyllin-type bicyclo[3.2.1]octanoid neolignans, ocophyllols A-C (22-24). The known compounds of 2'-epi-guianin (19) and licarin B (25) were also isolated (Coy-Barrera et al., 2009; Suarez et al., 2011). Funasaki et al., (2009) reported the isolation of fourteen neolignans (28-41) from the leaves of O. catharinensis. They managed to isolate seven new compounds; (7S, 8S, 1'R, 3'R)-3,4,5,3',5'-pentamethoxy-4'-oxo-Δ^{1,3,5,5',8'}-8.1',7.O.6'-neolignan (33), rel- (7R,8S,1'R,2'S)-2'-hydroxy-3,4,5'-trimethoxy-3'-oxo- $\Delta^{1,3,5,4',8'}$ -8.1',7.O.2'-neolignan (**35**), *rel*- (8*S*,1'*R*,5'*R*)-3,4,3',5'tetramethoxy-7,2'-dioxo- $\Delta^{1,3,5,3',8'}$ -8.1'-neolignan (36), rel-(7R,8S,1'R,2'S)-2'-hydroxy-3,4-methylenedioxy-5,3',5'trimethoxy- $\Delta^{1,3,5,3',8'}$ -8.1',7.O.2'-neolignan (37), rel-(7S,8R,1'R,3'R)-4'-hydroxy-3,4-methylenedioxy-3',5'-dimethoxy-2'.4'-dioxo- $\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan (**38**), rel- (75,8R,1'R,3'R)-4'-hydroxy-3,4-methylenedioxy-3',5'-dimethoxy-2',4'-dioxo- $\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan dimer (40), and (7S,8R,1'R,3'R)-4'hydroxy-3,4-methylenedioxy-3',5',5-trimethoxy-2',4'-dioxo- $\Delta^{1,3,5,5',8'}$ -8.1',7.3'-neolignan dimer (41). In addition, burchellin (1) was isolated from the leaves of O. cymbarum (Narciso et al., 2014), while canelin (2) from the stem of O. elegans (Oliveira et al., 2006). Besides, a new bicyclo[3.2.1]octanoid neolignan (26) was isolated from the fruit of O. heterochroma, together with a known compound, meso-dehydroguaiaretic acid (27) (Cuca et al., 2009).

Alkaloids

Most of alkaloids isolated from several Ocotea species were aporphine and benzylisoquinoline alkaloids. Forty one alkaloids were successfully identified from six Ocotea species. Garcez et al. (2005) isolated a new indole alkaloid; tryptophol-5-O- β -D-glucopyranoside (71) from the fruit of O. minarum. Six years later, they managed to isolate two new aporphinoid alkaloids; (+)-6S-ocoteine N-oxide (44) and norocoxylonine (46) from the leaves and trunk bark of O. acutifolia, along with thirteen aporphine analogues (+)-dicentrine (42), (+)-ocoteine (43), (+)leucoxine (45), (+)-thalicminine (47), (+)-neolitsine (48), (S)pallidine (68), (+)-reticuline (70), tryptophol-5-*O*-β-Dglucopyranoside (71), lequesnamine (NMS), (+)-6S-dicentrine Noxide (72), (+)-oxocylonine (73), (+)-O-methylcassyfiline (74),

(+)-nordicentrine (75), and (+)-thalicsimidine (76) and one morphinan alkaloid, N-methyllaurotethanine (79) (Garrett et al., 2010). Compound (42) was also isolated from the fruit of O. puberula (Montrucchio et al., 2012). Coy-Barrera et al., (2009) reported the isolation of seven aporphine alkaloids from the leaves of O. macrophylla. They were identified as dicentrine (42), neolitsine (48), dehvdronantenine (53), nantenine (54), N-acetylnornantenine (55), cassythidine (56), and didehydroocoteine (57). A year later, Pabon and Cuca (2010) managed to isolate four aporphine alkaloids from the wood of O. macrophylla, which were (S)-3-methoxy-nordomesticine (49). (S)-N-ethoxycarbonyl-3methoxy-nordomesticine (50), (S)-N-formyl-3-methoxynordomesticine (51), and (S)-N-methoxycarbonyl-3-methoxynordomesticine (52). Fournet and co-workers (2007)had successfully isolated thirteen known isoquinoline alkaloids from the stem bark of O. lancifolia. It comprises three benzyltetrahydroisoquinolines, coclaurine (58), N-methycoclaurine (59), norjuziphine (69), reticuline (70); two pro-aporphines: crostparine (NMS), glaziovine (60); eight aporphines, laurotetanine (62), nordomesticine (63), norisoboldine (64), domesticine (66), isoboldine (67), norantenine (NMS), caaverine (61), corytuberine (65), and the morphidanedienone alkaloid (S)-pallidine (68). Compound (58) was also isolated from the stem bark of O. duckei (Silva et al., 2002). Besides, Imler et al., (2003)had successfully isolated a new oxoaporphine alkaloid, lequesnamine from the wood of O. leucoxylon.

Flavonoids

Sixteen flavonoids were isolated from five Ocotea species. Garett et al., (2012) have successfully isolated an A-type proanthocyanidin trimer (80), isoquercitrin (81), reynoutrin (82), miquelianin (83), quercitrin (84), afzelin (85), and four minor compounds; catechin (86), epicatechin (87), quercetin (88), and kaempferol (89) from the leaves of O. notata. They also managed to isolate compound (89) from the leaves of O. acutifolia (Garcez et al., 2011). Compounds (81-82) were also isolated from the leaves of O. corymbosa, together with quercetin-3-O-β-Dgalactoside (90) (Batista et al., 2010). In addition, Batista et al., (2010) also isolated a flavonoid, rel- (2R,3R)-dihydroquercetin-3-O- α -L-rhamnoside (91) from the leaves of O. elegans. Garcez et al., (2005) obtained four flavonoids which were taxifolin (92), quercetin-7-O-β-D-glucopyranoside (93), eriodictyol-3'-O-β-Dglucopyranoside (94), and naringenin-7-O- β -D-glucopyranoside (95) from the fruit of O. minarum.

Sesquiterpenes

Eleven sesquiterpenes were isolated from four species of *Ocotea*. Camargo *et al.*, (2013) isolated six new eremophilane sesquiterpenes, *rel*-4 β ,5 β ,7 β -eremophil-9-en-12-oic acid (**96**), *rel*-4 β ,5 β ,7 β -eremophil-1 (10)-en-12-oic acid (**97**), *rel*-4 β ,5 β ,7 β -eremophil-1 (10)-en-2-oxo-12-oic acid (**98**), *rel*-4 β ,5 β ,7 β -eremophil-9-en-12,8 α -olide (**99**), *rel*-4 β ,5 β ,7 β -eremophil-9-en-12, 8 β -olide (**100**), and *rel*-4 β ,5 β ,7 β -eremophil-9 α ,10 α -epoxy-12-oic acid (**101**), from the leaves of *O. lancifolia*, together with two

known sesquiterpenes, 4β , 5β , 7β -eremophil-11-en-10 α -ol (102) reported fors the first time in the genus *Ocotea*, and the aromadendrene sesquiterpene, spathulenol (103). Besides, the sesquiterpene *rel*- (1*R*,4*S*)-7-hydroxycalamenene (104) isolated from the leaves of *O. elegans*, has already been isolated from *O. corymbosa*. However, there was no report regarding this compound on other Lauraceae genera (Batista *et al.*, 2010). Additionally, a new sesquiterpene of humulane type, *rel*- (8*R*)humulan-1,4-dien-8-ol (105), besides known spathulenol (103) was isolated form the leaves of *O. catharinensis* (Suarez *et al.*, 2011), while lanceolic acid (106) was identified from the trunk bark of *O. minarum* (Garcez *et al.*, 2005).

Lignans

Six lignans were isolated from four species of *Ocotea*. Two known lignans were isolated from the leaves of *O. macrophylla* which were *meso*-3,4,5,3',4',5'-hexamethoxy-8.8'lignan (**107**) and *threo*-3,4,5,3',4',5'-hexamethoxy-8.8'-lignan (**108**) (Coy-Barrera *et al.*, 2011). Meanwhile, Suarez *et al.*, (2011)also managed to isolate*erythro*-diarylbutane (**109**) and *threo*-diarylbutane (**110**) from the same species. Yangambin (**111**), a furofuran lignan was obtained from the leaves of *O. duckei* (Neto *et al.*, 2007, 2008) and fruits of *O. heterochroma* (Cuca *et al.*, 2009). Besides, lyonyresinol (**112**) was isolated from the heartwood of *O. minarum* (Garcez *et al.*, 2005).

Butanolides

Five butanolides were isolated from the root of O. *macrocarpa*. Macrocarpolide A (113), B (114) and C (115) were isolated for the first time from genus *Ocotea*. Compounds (114-115) belong to the class of secobutanolides (Liu *et al.*, 2015).

Benzopyrans

Four benzopyrans derivatives were isolated from the leaves of *O. corymbosa*. (2'S)-2- (propan-2'-ol)-5-hydroxy-benzopyran-4-one (**119**) and 2,3-dihydro-2-methyl-benzopyran-4,5-diol (**121**) were firstly isolated from the genus *Ocotea* (Teles *et al.*, 2005).

Steroids

Three steroids, $3-O-\beta$ -D-glucopyranosyl stigmasterol (123), stigmasta-4,22-dien-3-one (124), and β -sitosterol (125) were isolated from the heartwood and trunkbark of *O. minarum* (Garcez *et al.*, 2005). β -Friedelanol (122) was obtained from the fruit of *O. heretochroma* (Cuca *et al.*, 2009).

Alkylphenols

Two alkylphenols, 3- (1,4-dihydroxypentyl)-5methoxyphenol (**126**) and 5-propylresorcinol (**127**) were isolated from the heartwood of *O. minarum* (Garcez *et al.*, 2005)

Miscellaneous compounds

The arylpropene, *trans*-asarone (128), and a coumarin, scopoletin (129) were successfully isolated from the heartwood

and fruits of *O. minarum* (Garcez *et al.*, 2005). An ester, 4-*O*-*E*-caffeoylquinic acid methyl ester (**130**) was obtained from the leaves of *O. corymbosa* (Batista *et al.*, 2010), while a saponin, guaianin (**131**) was found from the leaves of *O. elegans* (Oliveira *et al.*, 2006).

Essential oils

Since 2000 to 2016, thirty one Ocotea species have been reported on their essential oils composition as shown in Table 2. Sesquiterpenoids were found as the major group components in most of the Ocotea essential oils, which are O. quixos, O. nirescens, O. splendens, O. macrophylla, O. gomezii, O. puberula, O. duckei, O. veraguensis, O. whitei, O. meziana, and O. bracteosa (Barbosa-Filho et al., 2008; Chaverri et al., 2007; Coutinho et al., 2007; Destryana et al., 2014; Farago et al., 2010; Garrett et al., 2010; Moriarity et al., 2007; Prieto et al., 2010; Sacchetti et al., 2006; Takaku et al., 2007; Wright et al., 2007; Yamaguchi et al., 2013). In addition, monoterpenoids were also detected from the essential oils of O. marae, O. longifolia, O. endresiana, O. praetermissa, O. floribunda, O. austinii, O. tonduzii, and O. holdridgeana (Agius et al., 2007; Bansal et al., 2007; Chaverri et al., 2011; Chaverri and Ciccio, 2007; Moriarity et al., 2007; Prieto et al., 2010; Takaku et al., 2007; Werka et al., 2007). Meanwhile, phenylpropanoids were also reported as the major group components isolated in O. odorifera, O. puchurymajor, and O. opifera (Leporatti et al., 2014; Lorenzo et al., 2001; Mossi et al., 2014; Oltramari et al., 2004).

BIOLOGICAL ACTIVITIES

The traditional medicinal applications of the *Ocotea* species have inspired many pharmacological investigations. The pharmacological activities were compiled in this section. Nineteen different biological activities have been reported from the extracts, essential oils, and their isolated compounds.

Cytotoxicity activity

The cytotoxicity of O. gomezii and O. morae essential oils were tested on cell lines (CCF-STTG1, Hep3B, HepG2, H-460, AGS, N-87, SW-620, MCF-7, and VERO), and found that astrocytoma cells were the most resistant. The leaves and bark oils of O. gomezii gave the best activity against SW620 (colon) with LD₅₀ of 122 and 94 µg/mL, respectively. While, the wood oil showed cytotoxicity activity against HepG2 (liver) with LD₅₀ of 94 µg/mL. In addition, the leaf, bark, and wood oils of O. morae showed activity against AGS (stomach) (LD₅₀ of 183 µg/mL), SW620 (colon) (LD₅₀ of 166 µg/mL), and HepG2 (liver) (LD₅₀ of 183 µg/mL), respectively (Chaverri et al., 2011). O. praetermissa and O. endresiana leaf oils were notably cytotoxic on MCF-7 cells (with >97% killing at 100 µg/mL) (Agius et al., 2007). O. whitei oil showed activity against M-14 melanoma cells (with 65% killing at 100 µg/mL). In addition, O. veraguensis oil was active against the estrogen receptor (ER) negative cell line, MDA-MB 231 (with 93% killing at 100 µg/mL) (Moriarity et al., 2007). O.

meziana, Ocotea sp. nov. 'los llanos', and Ocotea sp. nov. 'small leaf' have been reported to exhibit in vitro cytotoxic activity on MCF-7 human mammary adenocarcinoma cells (with 100% killing at 100 µg/mL) (Wright et al., 2007). O. floribunda oil has showed notable cytotoxic activity ($\geq 50\%$ killing on at least one cell line) with the best activity against HepG2 cell (78.8% killing at 100 µg/mL) (Werka et al., 2007). The (+)-ocoteine (43) showed the most potent effect against Hep-2 cells and (+)-neolitsine (48) being the most cytotoxic alkaloid to MCF-7 and B16-F10 cells. Thalicminine (47) on the other hand, was the most active compound against 786-0 cells, although it proved only marginally inhibitory to Hep-2 cells and nontoxic to MCF-7 and B16-F10 cells. While, (+)-6S-ocoteine N-oxide (44) was found to be only weakly active against Hep-2 and B16-F10 cells (IC50 of 32.7 and 30.7 µg/mL, respectively) and to be devoid of cytotoxicity against MCF-7 and 786-0 cells, as demonstrated by an IC₅₀ value higher than 50 µg/mL in both cases. These results indicated that, compared with (+)-ocoteine (43), N-oxide functionality in (44) reduces its cytotoxic activity (Garcez et al., 2011). Yangambin (111) isolated from O. duckei presented cytotoxicity to murine macrophages, measured by the Trypan blue dye exclusion test and MTT reduction assay, resulting in high IC₅₀ values of 187.0 µg/mL and 246.7 µg/mL, respectively (Neto et al., 2008).

Antibacterial and antifungal activities

O. praetermissa and O. endresiana leaf oils were slightly antibacterial against Bacillus cereus with MIC values of 78 and 156 µg/mL, respectively (Agius et al., 2007). O. bofo oil showed fair inhibiting properties against Escherichia coli, Staphylococcus aureus, and Bacillus subtilis with MIC values of 0.16 mg/mL, while a good inhibition against yeast, Yarrowia lypolytica with MIC values of 0.10 mg/mL (Guerrini et al., 2006). The O. quixosoil also showed a dose-dependent antifungal activity against Candida albicans and Saccharomyces cerevisiae (MIC of 0.024 mg/ml), while the antifungal activity against the dermatophyte and phytopathogen strains by the essential oil was relatively good. The growth inhibition percentage against the dermatophyte Trichophyton mentagrophytes was 60% at the highest concentration tested. However, O. quixosoil performed better against the phytopathogen Pythium ultimum, with an inhibitory action of 85% at 500 mg/mL (Bruni et al., 2003). The in vitro antifungal activity of essential oil from O. odorifera had weak activity against C. albicans and C. tropicalis strains involved in oral cavity infections.

A slight antifungal activity was observed with 2.5 mg/mL and MIC in 68% strains (Castro *et al.*, 2011). The essential oil of *O. longifolia* showed significant fumigant activity against *Sitophilus zeamais* (CL₅₀ of 280.5 μ L/L) (Prieto *et al.*, 2010). The alkaloid fraction of *O. macrophylla* was active against *Fusarium oxysporum* at 250 μ g/ μ L. The inhibitory activity against the growth of the fungi was moderate at 5 μ g/ μ L for (*S*)-3-methoxynordomesticine (**49**) while the other alkaloids were ineffective, suggesting that the presence of electron withdrawing substituents on nitrogen atom decreased the antifungal activity. In addition,

alkaloid (**49**) also showed antimicrobial activity against *Staphylococcus aureus* and *Enterococcus faecalis* with values of 30 AU (Pabon and Cuca, 2010).

Anti-inflammatory activity

The essential oils of O. nigrescens and O. splendens have showed a low anti-aggregant factor with 10.8% and 11.74%, respectively, as compared with the positive standard acetylsalicylic acid which strongly inhibits platelet aggregation (100%). The low antiplatelet activity against ADP shown by both essential oils might be due to the absence of phenylpropanoids in their composition (Yamaguchi et al., 2013). O. quixos essential oil significantly reduced LPS-induced NO release from J774 macrophages at non-toxic concentrations, inhibited LPS-induced COX-2 expression and increased forskolin-induced cAMP production. The essential oil in carrageenan-induced rat paw edema showed anti-inflammatory effect without damaging gastric mucosa (Ballabeni et al., 2010). The diastereomeric lignans, the meso-isomer (107) and threo-isomer (108) were found to be the potent COX-2/5-LOX dual inhibitor and PAF-antagonist. Compound (107) displayed a lower IC_{50} value than (108), in contrast to their COX-inhibition. The IC_{50} value of (107) for 5-LOX and PAFwere 46.4 and 38.9 µM, respectively, while COX-2 for (108) was 15.6 µM (Coy-Barrera et al., 2011). 2'-epi-Guianin (19) showed the most potent inhibition of platelet activating factor (PAF)-induced aggregation of rabbit platelets among other neolignans isolated from O. macrophyla, with IC₅₀ value 1.6 µM (Coy-Barrera et al., 2009). Sibyllenone (10) was the only compound from O. bullata which showed good inhibitory activity towards 5-lipoxygenase with IC50 value of 18.6 µM (Zschocke et al., 2000).

Antileishmanial activity

The ethanolic leaves extract of *O. macrophylla* showed the efficacy assay against both Leishmania parasites, *L. panamensis* and *L. braziliensis* with EC₅₀ values of 98.0 and 85.7 µg/mL,respectively. In addition, *erythro*-diarylbutane (**109**) and ocophyllal B (**21**) the isolated compounds from *O. macrophylla* have shown the best activity against *L. panamensis* (IC₅₀ value 26.6 µg/mL) and *L. braziliensis* (IC₅₀ value 36.3 µg/mL), respectively (Suarez *et al.*, 2011). The crude ethanolic extract, lignoid fraction, and the purified compound, yangambin (**111**) obtained from *O. duckei* presented antileishmanial activity with IC₅₀ values of 135.7, 26.5, and 49.0 µg/mL, respectively on *Leishmania chagasi*. Meanwhile, for *Leishmania amazonensis*, the IC₅₀ values were 143.7, 48.2, and 64.9 µg/mL for the crude ethanolic extract, the lignoid fraction, and the purified compound yangambin (**111**), respectively (Neto *et al.*, 2007).

Cruzain inhibitory activity

O. praetermissa and *O. endresiana* leaf oils inhibited cruzain with IC₅₀ values of 87.5 and 18.8 μ g/mL, respectively (Agius *et al.*, 2007). The leaf essential oils of ten species of *Ocotea* from Costa Rica were examined on the enzyme inhibitory

activities against cruzain using a fluorometric assay. The *O. meziana* leaf oil was the most active (IC₅₀ value 14.9 µg/mL) followed by *O. whitei* (15.8 µg/mL), *Ocotea* sp. nov. 'los llanos' (17.1 µg/mL), *Ocotea* sp. nov. 'small leaf' (19.2 µg/mL), and *O. holdridgeana* (76.9 µg/mL). The leaf oils of *O. floribunda*, *O. tonduzii*, and *O. valeriana* were somewhat active (IC₅₀ value 100-200 µg/mL), but *O. sinuata* and *O. veraguensis* essential oils were inactive (IC₅₀>500 µg/mL). The cruzain inhibitory activities of the essential oils can be attributed to active sesquiterpenoid components as well as synergistic effects between two or more components (Setzer *et al.*, 2006).

Mutagenic and recombinogenic activities

The somatic mutation and recombination test (SMART) in wing cells of Drosophila melanogaster was used to test the mutagenic and recombinogenic activities of alkaloids isolated from O. acutifolia. Third-stage larvae derived from the standard cross with wing cell markers mwh and/or flr³ were treated chronically. The frequencies of mutant spots observed in marked heterozygous descendants revealed significant dose-dependent genotoxicity for alkaloids; thalicminine (47), (+)-dicentrine (42), (+)-ocoteine (43), and (+)-6S-ocoteine N-oxide (44). These alkaloids also induced mitotic recombination. The presence of a methoxyl group at C-3 (as in compound 43) lowers its genotoxic effect relative to that of unsubstituted analogue (42), and the introduction of N-oxide functionality (43 vs. 44) further reduces its genotoxicity. The very planar conformation of oxoaporphine alkaloid (47) may account for its higher genotoxicity vs. its lessplanar analogues (43) and (44) (Guterres et al., 2013).

Antiherpetic activity

The flavonoid fraction from *O. notata* leaves extract showed antiherpes activity against both herpes simplex viruses;type 1 (HSV-1) and 2 (HSV-2) with EC₅₀ values of 35.8 and 23.5 μ g/mL, respectively. Moreover, this fraction was more active against HSV-2 than HSV-1. The mechanisms of antiviral action of the flavonoid fraction against the herpesvirus types 1 and 2 were evaluated and the inhibition of different steps of the virus replication cycle was observed. The percentage inhibition obtained for HSV-2 was higher than 90% in all performed experiments. Differently, for HSV-1, the flavonoid fraction had no effect in pretreatment of cells and showed 60% of inhibition in virucidal assay (Garett *et al.*, 2012).

Antioxidant activity

The antioxidant activity (TBARS method) of the essential oil from *Ocotea* sp., showed the lowest mean effective concentration with EC₅₀ value of 31.1 µg/mL (Olivero *et al.*, 2010). *O. bofo* oil revealed a remarkable inhibitory scavenging effect on DPPH with inhibition of 64.4%, while in β -carotene bleaching test gave 75.8% inhibition. In the photochemiluminescence (PCL) assay, the oil gave 3.14 mmol of Trolox/L (Guerrini *et al.*, 2006). The *O. quixos* oil exerted a relatively good capacity to act as a non-specific donor of hydrogen

atoms or electrons whenchecked by the DPPH assay, quenching 52% of the radical. On the other hand, it showed weak effects in inhibiting theoxidation of linoleic acid when assayed by the β -carotene bleaching test (Bruni *et al.*, 2003).

Cardiovascular activity

The pharmacological activity of *O. duckei* essential oil showing significant cardiovascular effects. The leaves oil induced significant hypotension, followed by intense bradycardia. The fruits also induced a marked hypotension, which was followed by bradycardia. The stem bark and roots were both able to induce hypotension and bradycardia. Among all of the essential oils tested, the hypotensive effect was more potent on diastolic arterial blood pressure compared with the effect induced on systolic pressure (Barbosa-Filho *et al.*, 2008).

Antithrombotic activity

O. quixos essentially possesses potent and safe antithrombotic activity attributable to its antiplatelet and vasorelaxant effects. The best inhibitory potency against platelet aggregation in guinea pig PRP induced by arachidonic acid (IC₅₀ of 47 μ g/mL), while platelet aggregation in human PRP induced by thromboxane A₂ receptor agonist U46619 (IC₅₀ of 115 μ g/mL). The antithrombotic activity could be related to its ability to block both platelet aggregation and clot retraction and to inhibit vasoconstriction (Ballabeni *et al.*, 2007).

Antiproliferative activity

Macrocarpolides A-C (**113-115**), linderanolide B (**116**), and isolinderanolide (**117**) showed good antiproliferative activities against the A2780 ovarian cell line, with IC₅₀ values of 2.57, 1.98, 1.67, 2.43, and 1.65 μ M, respectively. The similar IC₅₀ values for the five compounds suggested that they have a similar mechanism of action, possibly as Michael acceptors (Liu *et al.*, 2015). A benzopyran (**119**) was found to induce cell proliferation: 70% on HeLa (human cervix tumour) cells and 25% on CHO (Chinese hamster ovary) cells (Teles *et al.*, 2005).

Antinociceptive activity

Dicentrine (42) produced dose-related inhibition of acetic acid-induced pain without causing changes in the motor performance of mice. Furthermore, the chloroform fraction from *O. puberula* fruit produced marked antinociception in different models of chemical pain, and this effect appears to be, at least in part, due to the presence of dicentrine (42). The mechanisms by the extract and the alkaloid produced antinociception still remains unclear, but the adenosinergic or opioid system seems unlikely to be involved in this action (Montrucchio *et al.*, 2012).

Larvicidal activity

Burchellin (1) isolated from *O. cymbarum* interfered with the development cycle of *Aedes aegypti*, where its strongest toxic effect was 100% mortality in larvae (L3) at concentrations \geq 30 ppm (Narciso *et al.*, 2014). The new 8.O.6'-neolignan (6), dihydrobenzofuranoid (7), and bicyclo[3.2.1]octanoid (11) havein vitro activities against Aedes aegypti, with \geq 80% mortality at 4 mg/mL (Rakotondraibe *et al.*, 2015).

Lethality activity

O. praetermissa and *O. endresiana* leaf oils were active in the brine shrimp lethality test against *Artemia salina* with LC_{50} values of 31.6 and 6.9 µg/mL, respectively (Agius *et al.*, 2007). The *O. floribunda* oil showed notable brine shrimp toxicity against *Artemia salina* with LC_{50} value of 3.7 µg/mL. On the other, the oil also showed antibacterial activity on *Staphylococcus aureus* with MIC value of 78 µg/mL (Werka *et al.*, 2007).

Antiprotozoal activity

(-)-Caaverine (**61**) has shown the most interesting antiprotozoal activity against *Leishmaniabraziliensis*, *Leishmania amazonensis*, and *Leishmania donovani* with IC₁₀₀ of 10 µg/mL, respectively. Meanwhile, (-)-caaverine (**61**) and (+)-domesticine (**66**) showed the best activity against *Trypanosoma cruzi* parasite with IC₅₀ of 155 and 105 µg/mL, respectively (Fournet *et al.*, 2007).

Toxicity activity

The essential oil of *O. notata* was evaluated by means of the brine shrimp lethality test and showed a high toxicity with an LC_{50} value of 2.37 µg/mL (Garrett *et al.*, 2010). The *Ocotea* sp. essential oil has showed high cytotoxicity (LC_{50} value 7.84 µg/mL) against *Artemia franciscana* on the brine shrimp assay (Olivero *et al.*, 2010).

Molluscicidal activity

The stem bark oil of *O. bracteosa* showed significant molluscicidal activity against *Biomphalaria glabrata*, with an LC_{90} value of 8.3 µg/mL, which falls below the threshold of 100 µg/mL, set for potential molluscicidal activity by the World Health Organization (Coutinho *et al.*, 2007).

Antiplasmodial activity

Ococymosin (7) was the most active antiparasitic component than other neolignans, with an IC₅₀ value of 0.45 μ M against the Dd2 strain of *Plasmodium falciparum* (Rakotondraibe *et al.*, 2015).

Antilmalarial activity

The antimalarial activity of the *O. comoriensis* oil is interesting, with an IC₅₀ value of 10 μ g/mL (Menut *et al.*, 2002).

CONCLUSION

The genus *Ocotea* is widespread all over the world, and many species of this genus have been used as traditional herbal medicines. The chemical investigation of *Ocotea* species has revealed that many components from this genus exhibit significant bioactivities. Nevertheless, there are still several *Ocotea* species that have received no or only little attention, and phytochemical and biological studies now should focus on these plants.

Financial support and sponsorship: NIL.

Conflict of Interests: There are no conflicts of interest.

REFERENCES

Agius BR, Setzer MC, Stokes SL, Walker TM, Haber WA, Setzer WN. Composition and bioactivity of essential oils of Lauraceae from Monteverde, Costa Rica. Int J Essent Oil Ther, 2007; 1: 167-171.

Ballabeni V, Tognolini M, Giorgio C, Bertoni S, Bruni R, Barocelli E.*Ocotea quixos* Lam. essential oil: *In vitro* and *in vivo* investigation on its anti-inflammatory properties. Fitoterapia,2010; 81:289-295.

Ballabeni V, Tognolini M, Bertoni S, Bruni R, Guerrini A, Rueda GM, Barocelli E. Antiplatelet and antithrombotic activities of essential oil from wild *Ocotea quixos* (Lam.) Kosterm. (Lauraceae) calices from Amazonian Ecuador. Pharmacol Res, 2007; 55: 23-30.

Bansal A, Moriarity DM, Takaku S, Setzer WN. Chemical composition and cytotoxic activity of the leaf essential oil of *Ocotea tonduzii* from Monteverde, Costa Rica. Nat Prod Commun, 2007; 2: 781-784.

Barbosa-Filho JM, Cunha RM, Dias CS, Athayde-Filho PF, Silva MS, Leitao da-Cunha EV, Machado MIL, Craveiro AA, Medeiros IA. GC-MS analysis and cardiovascular activity of the essential oil of *Ocotea duckei*. Rev Bras Farmacogn, 2008; 18: 37-41.

Batista ANL, Batista JMJ, Lopez SN, Furlan M, Cavalheiro AJ, Silva DHS, Bolzani VS, Nunomura SM, Yoshida M. Aromatic compounds from three Brazilian Lauraceae species. Quim Nova, 2010; 33: 321-323.

Bruni R, Medici A, Andreotti E, Fantin C, Muzzoli M, Dehesa M, Romagnoli C, Sacchetti G. Chemical composition and biological activities of Ishpingo essential oil, a traditional Ecuadorian spice from Ocotea quixos (Lam.) Kosterm. (Lauraceae) flower calices. Food Chem, 2003; 85: 415-421.

Camargo MJ, Miranda MLD, Kagamida CM, Rodrigues ED, Garcez FR, Garcez WS. Sesquiterpenes of *Ocotea lancifolia* (Lauraceae). Quim Nova, 2013; 36: 1008-1013.

Castro RD, Lima EO. Antifungal activity of Brazilian sassafras (*Ocotea odorifera* Vell.) and rosemary (*Rosmarinus officinalis* L.) essential oils against genus Candida. Rev Bras Plant Med, 2011; 13: 203-208.

Coutinho DF, Dias CS, Barbosa-Filho JM, Agra MF, Martins RM, Silva TMS, da-Cunha EVL, Silva MS. Composition and molluscicidal activity of the essential oil from the stem bark of *Ocotea bracteosa* (Meisn.) Mez. J Essent Oil Res, 2007; 19: 482-484.

Chaverri C, Diaz C, Ciccio JF. Chemical analysis of essential oils from *Ocotea gomezii* W.C. Burger and *Ocotea morae* Gomez-Laur. (Lauraceae) collected at "reserva biologica Alberto M. Brenes" in Costa Rica and their cytotoxic activity on tumor cell lines. J Braz Chem Soc, 2011; 22: 741-745.

Chaverri C, Ciccio JF. Essential oils from *Ocotea austinii* C. K. Allen (Lauraceae) from Costa Rica. J Essent Oil Res, 2007; 19: 439-443.

Christophel DC, Kerrigan R, Rowett AI. The use of cuticular features in the taxonomy of the Lauraceae. Ann Miss Bot Garden, 1996; 83: 419-432.

Coy-Barrera ED, Cuca-Suarez LE. Aporphine alkaloids from leaves of *Ocotea macrophylla* (Kunth) (Lauraceae) from Colombia. Biochem Syst Ecol, 2009; 37: 522-524.

Coy-Barrera ED, Cuca-Suarez LE, Sefkow M. PAFantagonistic bicyclo[3.2.1]octanoid neolignans from leaves of *Ocotea macrophylla* Kunth. (Lauraceae). Phytochemistry, 2009; 70: 1309-1314.

Coy-Barrera ED, Cuca-Suarez LE. In vitro anti-inflammatory effects of naturally-occurring compounds from two Lauraceae plants. Anais Acad Bras Cien, 2011; 83: 1397-1402.

Coy-Barrera ED, Cuca-Suarez LE, Sefkow M. COX, LOX and platelet aggregation inhibitory properties of Lauraceae neolignans. Bioorg Med Chem Lett, 2009; 9:6922-6925.

Cuca LE, Leon P, Coy ED. A bicyclo[3.2.1]octanoid neolignan and toxicity of the ethanol extract from the fruit of *Ocotea heterochroma*. Chem Nat Comp, 2009; 45: 179-181.

Destryana RA, Young DG, Woolley CL, Huang TC, Wu Hyi, Shih WL.) Antioxidant and anti-inflammation activities of *Ocotea*, Copaiba and Blue Cypress essential oils in vitro and in vivo. J Am Oil Chem Soc, 2014; 91: 1531-1542.

Enrico R, Matteo M, Silvia M, Gianni S, Renato B. Comparative phytotoxicity of 25 essential oils on pre- and post-emergence development of *Solanum lycopersicum* L.: A multivariate approach. Ind Crops Prod,2014; 60:280-290.

Farago PV, Padilha-de-Paula JF, Nakashima T, Doll-Boscardin PM, Budel JM, Maia BHLNS. Chemical composition and antibacterial activity of the essential oil from bark of *Ocotea puberula* (Rich.) Ness. Latin Am J Pharm, 2010; 29: 1242-1245.

Fournet A, Ferreira ME, Rojas de Arias A, Guy I, Guinaudeau H, Heinzen H. Phytochemical and antiprotozoal activity of *Ocotea lancifolia*. Fitoterapia, 2007; 78: 382-384.

Funasaki M, Lordello ALL, Viana AM, Santa-Catarina C, Floh EIS, Yoshida M, Kato MJ. Neolignans and sesquiterpenes from leaves and embryogenic cultures of *Ocotea catharinensis* (Lauraceae). J Braz Chem Soc, 2009; 20: 853-859.

Garcez WS, Garcez FR, da Silva LMGE, Shimabukuro AA. Indole alkaloid and other constituents from *Ocotea minarum*. J Braz Chem Soc, 2005; 16: 1382-1386.

Garcez FR, da Silva AFG, Garcez WS, Linck G, Matos MFC, Santos ECS, Queiroz LMM. Cytotoxic aporphine alkaloids from *Ocotea acutifolia*. Planta Med, 2011; 77: 383-387.

Garrett R, Cruz RAS, Rocha L, Santos MG, da Silva AJR. Chemical composition and toxicity of *Ocotea notata* (Nees) Mez essential oil. J Essent Oil Bear Pl, 2010; 13: 455-459.

Garett R, Romanos MTV, Borges RM, Santos MG, Rocha L, da Silva AJR. Antiherpetic activity of a flavonoid fraction from *Ocotea notata* leaves. Rev Bras Farmacogn, 2012; 22: 306-313.

Guerrini A, Sacchetti G, Muzzoli M, Moreno RG, Medici A, Besco E, Bruni R. Composition of the volatile fraction of *Ocotea bofo* Kunth (Lauraceae) calyces by GC-MS and NMR fingerprinting and its antimicrobial and antioxidant activity. J Agric Food Chem, 2006; 54: 7778-7788.

Guterres Zda R, da Silva AF, Garcez WS, Garcez FR, Fernandes CA, Garcez FR. Mutagenicity and recombinagenicity of *Ocotea acutifolia* (Lauraceae) aporphinoid alkaloids. Mut Res, 2013; 757: 91-96.

Imler MJ, Menachery MD, Rajaraman R, Cava MP, Freyer AJ, Shi SD. 2003. Lequesnamine: A new oxoaporphine alkaloid from *Ocotea leucoxylon*. Abstracts of Papers, 225th ACS National Meeting, New Orleans, LA, United States.

Leporatti ML, Pintore G, Foddai M, Chessa M, Piana A, Petretto GL, Masia MD, Mangano G, Nicoletti M. Chemical, biological, morphoanatomical and antimicrobial study of *Ocotea puchury-major* Mart. Nat Prod Res, 2014; 28: 294-300.

Liu Y, Cheng E, Rakotondraibe LH, Brodie PJ, Applequist W, Randrianaivo R, Rakotondrafara A, Ratsimbason M, Rasamison VE, Kingston DG. Antiproliferative compounds from *Ocotea macrocarpa* from the Madagascar dry forest. Tetrahedron Lett, 2015; 56: 3630-3632.

Lorenzo D, Loayza I, Leigue L, Frizzo C, Dellacassa E, Moyna P. Asaricin, the main component of *Ocotea opifera* Mart. essential oil. Nat Prod Lett, 2001; 15: 163-170.

Menut C, Bessiere JM, Hassani MS, Buchbauer G, Schopper B. Chemical and biological studies of *Ocotea comoriensis* bark essential oil. Flav Fragr J, 2002; 17: 459-461.

Montrucchio DP, Miquel OG, Zanin SMW, Araujo da Silva G, Cardozo AM, Santos ARS. Antinociceptive effects of a chloroform extract and the alkaloid dicentrine isolated from fruits of *Ocotea puberula*. Planta Med, 2012; 78: 1543-1548. Moriarity DM, Bansal A, Cole RA, Takaku S, Haber WA, Setzer WN. Selective cytotoxic activities of leaf essential oils from Monteverde, Costa Rica. Nat Prod Commun, 2007; 2: 1263-1268.

Mossi AJ, Zanella CA, Kubiak G, Lerin LA, Cansian RL, Frandoloso FS, Prá VD, Mazutti MA, Costa JAV, Treichel H. Essential oil of *Ocotea odorifera*: An alternative against Sitophilus zeamais. Renew Agric Food Syst, 2014; 29: 161-166.

Narciso JO, Soares RO, Reis dos Santos Mallet J, Guimarães AÉ, de Oliveira Chaves MC, Barbosa-Filho JM, Maleck M. Burchellin: study of bioactivity against *Aedes aegypti*. Parasites Vect, 2014; 7: 172.

Neto RLM, Sousa LMA, Dias CS, Barbosa Filho JM, Oliveira MR. Yangambin cytotoxicity: A pharmacologically active lignan obtained from *Ocotea duckei* Vattimo (Lauraceae). Z Naturforsch C: J Biosci, 2008; 63: 681-686.

Neto RLM, Barbosa-Filho JM, Sousa LMA, Athayde Filho PF, Dias CS, Oliveira MR. Crude ethanolic extract, lignoid fraction and yangambin from *Ocotea duckei* (Lauraceae) show antileishmanial activity. Z Naturforsch C: J Biosci, 2007; 62: 348-352.

Oliveira R, Heringer A, Figueiredo M, Futuro D, Kaplan M. Isolation of neolignans from *Ocotea elegans* by CCC. J Liq Chromatogr RT, 2006; 29: 229-234.

Olivero VJ, Gonzalez-Cervera T, Guette-Fernandez J, Jaramillo-Colorado B, Stashenko E. Chemical composition and antioxidant activity of essential oils isolated from Colombian plants. Rev Bras Farmacogn, 2010; 20: 568-574.

Oltramari AC, Wood KV, Bonham C, Verpoorte R, Caro MSB, Viana AM, Pedrotti EL, Maraschin RP, Maraschin M. Safrole analysis by GC-MS of prototrophic (*Ocotea odorifera* (Vell.) *Rohwer*) cell cultures. Plant Cell Tiss Org Cult, 2004; 78: 231-235.

Pabon LC, Cuca LE. Aporphine alkaloids from *Ocotea macrophylla* (Lauraceae). Quim Nova, 2010; 33: 875-879.

Pino JA, Fernandes P, Marbot R, Sa Fontinha S. Chemical composition of the leaf oil of *Ocotea foetens* (Alt.) Benth. et Hook. from Madeira. J Essent Oil Res, 2004; 16: 131-132.

Prieto JA, Pabon LC, Patino OJ, Delgado WA, Cuca LE. Chemical constituents and insecticidal and antifungal activities of the essential oils of leaves of two Colombian species of the genus *Ocotea* (Lauraceae). Rev Colomb Quim, 2010; 39: 199-209.

Rakotondraibe LH, Graupner PR, Xiong Q, Olson M, Wiley JD, Krai P, Brodie PJ, Callmander MW, Rakotobe E, Ratovoson F, Rasamison VE, Cassera MB, Hahn DR, Kingston DG, Fotso S. Neolignans and other metabolites from *Ocotea cymosa* from the Madagascar Rain Forest and their biological activities. J Nat Prod, 2015; 78: 431-440.

Rohwer JG. Toward a phylogenetic classification of the Lauraceae: evidence from matK sequences. Syst Bot, 2000; 60-71.

Sacchetti G, Guerrini A, Noriega P, Bianchi A, Bruni R. Essential oil of wild *Ocotea quixos* (Lam.) Kosterm. (Lauraceae) leaves from Amazonian Ecuador. Flav Frag J, 2006; 21: 674-676.

Scora RW, Scora PE. Essential leaf oil of *Persea subgenus* Eriodaphne and closely related *Perseoid* genera. J Essent Oil Res, 2001; 13: 37-42.

Setzer WN, Takaku S, Stokes SL, Penton AF. Inhibition of cruzain by *Ocotea* leaf essential oils from Monteverde, Costa Rica. Pharmacologyonline, 2006; 3: 785-793.

Silva IG, Barbosa-Filho JM, Silva MS, Lacerda CDG, Cunha EVL. Coclaurine from *Ocotea duckei*. Biochem Syst Ecol, 2002; 30: 881-883.

Suarez SJ, Coy-Barrera ED; Cuca LE, Delgado G. Leishmanicidal and cytotoxic activities of extracts and naturally-occurring compounds from two Lauraceae species. Nat Prod Commun, 2011; 6: 231-234.

Takaku S, Haber WA, Setzer WN. Leaf essential oil composition of 10 species of *Ocotea* (Lauraceae) from Monteverde, Costa Rica. Biochem Syst Ecol, 2007; 35: 525-532.

Teles HL, Silva GH, Castro-Gamboa I, Bolzani VS, Pereira JO, Costa-Neto CM, Haddad R, Eberlin MN, Young MCM, Araujo AR.

Benzopyrans from *Curvularia* sp., an endophytic fungus associated with *Ocotea corymbosa* (Lauraceae). Phytochemistry, 2005; 66: 2363-2367.

Tognolini M, Barocelli E, Ballabeni V, Bruni R, Bianchi A, Chiavarini M, Impicciatore M. Comparative screening of plant essential oils: Phenylpropanoid moiety as basic core for antiplatelet activity. Life Sci,2006; 78:1419-1432.

van der Werff H. A revision of the genus *Ocotea* Aubl. (Lauraceae) in Madagascar and the Comoro Islands. Adansonia, 2013; 35: 235-279.

Werka JS, Boehme AK, Setzer WN. Biological activities of essential oils from Monteverde, Costa Rica. Nat Prod Commun, 2007; 2: 1215-1219.

Wright BS, Bansal A, Moriarity DM, Takaku S, Setzer WN. Cytotoxic leaf essential oils from neotropical Lauraceae: synergistic effects of essential oil components. Nat Prod Commun, 2007; 2: 1241-1244.

Yamaguchi KKL, Alcantara JM, Lima ES, Veiga-Junior VF. Chemical composition and platelet aggregation activity of essential oils of two species of the genus *Ocotea* (Lauraceae). J Essent Oil Bear Pl, 2013; 16: 518-523.

Zschocke S, van Staden J, Paulus K, Bauer R, Horn MM, Munro OQ, Brown NJ, Drewes SE. Stereostructure and anti-inflammatory activity of three diastereomers of ocobullenone from *Ocotea bullata*. Phytochemistry, 2000; 54: 591-595.

How to cite this article:

Salleh WMN, Ahmad F. Phytochemistry and Biological Activities of the Genus *Ocotea* (Lauraceae): A Review on Recent Research Results (2000-2016). J App Pharm Sci, 2017; 7 (05): 204-218.