RSC Advances

REVIEW

rsc.li/rsc-advances

Cite this: RSC Adv., 2017, 7, 14114

Received 6th December 2016 Accepted 22nd February 2017 DOI: 10.1039/c6ra27830b

An overview of chemical constituents from Alpinia species in the last six decades†

Xiao-Ni Ma,^{ab} Chun-Lan Xie,^{ab} Zi Miao,^a Quan Yang^b and Xian-Wen Yang^{*a}

Alpinia species is one of the most important genera of the Zingiberaceae family. In Asia, they have been widely used as food and traditional medicines for centuries. This review focuses on their chemical constituents and their relevant biological activities with 252 references covering from 1955 to 2015. In total, 544 compounds were isolated from 35 Alpinia species. The major ones are terpenoids (207) and diarylheptanoids (143). The crude extracts and identified compounds exhibited a broad spectrum of bioactivities including antiemetic, antiulcer, antibacterial, anti-inflammatory, anti-amnesic, anticancer, etc.

> Up to 2015, this genus contributed about 252 papers. However, only seven review articles were published, five of which were on chemical constituents and biological activities of single plant. And the rest two were on two major components of Alpinia species. The first review came out in 2010 regarding distributions, physiological activities and 13 C NMR spectroscopic data of 307 naturally occurring diarylheptanoids, which were mainly isolated from Alpinia species.⁴¹ In 2011, the pharmacological and phytochemical studies of A. galanga (L.) Willd were summarized with 30 references. Although it was claimed to concern new phytoconstituents that have appeared in recent years for A. galangal, it actually collected all reported compounds including volatile oil.⁴² In 2012, structural characterization and biological effects of constituents from the seeds of A. katsumadai was described. Sixty compounds were reported together with their structures and bioactivities with 18 references.⁴³ In 2013, chemical constituents in fruits of A. oxyphylla and their pharmacological activities were summarized. Eighty-five compounds were obtained from this species between 2001 and 2012, with the major component of sesquiterpenes (61.2%). It possessed a variety of pharmacological activities, including neuroprotection, learning and memoryimproving function, anticancer, anti-aging, anti-inflammation, and anti-anaphylaxis.⁴⁴ In 2015, a comprehensive review on the ethnomedical uses, chemical constituents, and the pharmacological profile of A. calcarata Roscoe was published with particular attention given to the pharmacological effects of the essential oil.⁴⁵ In the same year, the phytochemistry of A. purpurata with pharmacological properties of antioxidant, antibacterial, larvicidal, cytotoxic, and vasodilator activities were reported together with another ornamental ginger, Hedychium coronarium. As a matter of fact, little research was performed on

1. Introduction

The genus Alpinia is an important member of the Zingiberaceae family. It includes ca. 230 species.¹ Most of them are distributed in tropical and subtropical Asia, including India, Malaysia, China, and Japan. A few are found in Australia and the Pacific Islands.¹⁻³ Plants of this genus have been extensively used for different purposes for centuries. For example, A. vittata, A. purpurata (Vieill.) K. Schum., A. calcarata Rosc., and A. zerumbet are cultivated as ornamental plants;^{3,4} A. *blepharocalyx* K. Schum. is a natural dye;⁵ A. galanga (L.) Willd is an important ingredient for curries and has been broadly utilized as a flavoring in the preparation of meats and soups in Southeast Asia⁶⁻⁸ and in the preparation of beverages in Europe;⁹ and A. officinarum Hance, listed as medicinal and edible food by the Chinese Ministry of Health, are used in medicinal diets,¹⁰⁻¹⁵ wines,¹⁶ sauces, and flavorings.¹⁷⁻¹⁹ Moreover, A. galanga (L.) Willd is also applied to preserve food and fruits.^{8,20} Most important of all, Alpinia plants are also broadly used as traditional medicines in India, China, and Japan to treat many diseases such as indigestion, gastralgia, vomiting, enterozoa $etc.²¹⁻²³$ Thus, a growing investigation on the chemical constituents and bioactivities of this genus has been carried out since 1955.²⁴ Consequently, *Alpinia* species were proved to have various biological activities including antiulcer, 25 antiemetic, $26-28$ antibacterial,²⁹⁻³¹ antitumor,³²⁻³⁴ hypoglycemic,³⁵ cardioprotection,³⁶ antifungi,³⁷ neuroprotection,^{38,39} and antianxiety activities.⁴⁰ **EXAMELIAR SEAL SET AND SERVIEW AND SPECIES IN OVERVIEW Of chemical constituents from Alpin species in the last six decades?

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A. purpurata.⁴⁶ In addition, the isolation, synthesis, and characterization of dihydro-5,6-dehydrokavain, the major constituent of A. zerumbet were also reviewed.⁴⁷ However, so far there has been no comprehensive review for chemical constituents of this species. Herein, we describe all isolated compounds and their 14114 | RSC Adv., 2017, 7, 14114-14144 This journal is © The Royal Society of Chemistry 2017

[&]quot;State Key Laboratory Breeding Base of Marine Genetic Resources, Key Laboratory of Marine Genetic Resources, Fujian Key Laboratory of Marine Genetic Resources, Third Institute of Oceanography, State Oceanic Administration, 184 Daxue Road, Xiamen 361005, PR China. E-mail: yangxianwen@tio,org,cn

b Department of Traditional Chinese Medicine, Guangdong Pharmaceutical University, Guangzhou 510006, China

[†] Electronic supplementary information (ESI) available: The name, source, plant part, and reference for each compound. A comparison of Alpinia species names from the references and the accepted name in The Plant List. See DOI: 10.1039/c6ra27830b

relevant bioactivities of Alpinia species reported in the last six decades from 1955 to 2015.

2. Terpenoids

2.1. Monoterpenoids

A total number of 17 monoterpenoids were obtained from Alpinia species (Fig. 1). Rubraine (1) , isorubraine (2) , and sumadain C (3) were three new monoterpene–chalcone conjugates obtained from A. katsumadai.⁴⁸ They were tested for cytotoxic activities against three tumor cell lines of HepG2, MCF-7, and MAD-MB-435. Sumadain C (3) exhibited very weak effect with IC_{50} value of around 40.0 μ M.⁴⁸ A. katsumadai Hayata yielded a new monoterpene–kavalactone conjugate, katsumadain (4) and a new (E)-1-(1-terpinen-4-olyl)-3 methoxystilbene (5).⁴⁹ While A. densibracteata T. L. Wu and Senjen yielded two diastereoisomers of cinnamate esters, 2a-cinnamoyl cineole (6) and 2 β -cinnamoyl cineole (7).⁵⁰ From rhizomes of A. tonkinensis Gagnep., 2a-(p-hydroxycinnamoyl) cineole (8) was isolated.50,51 Two endoperoxides, (1S,4R,6R)-1,4-epidioxy-p-menth-2-ene (9) and (1R,4S,6R)-1,4-epidioxy-p-menth-2-ene (10), were isolated from aerial parts of A. densibracteata T. L. Wu and Senjen.⁵⁰ Whilst (3R,4R,6S)-3,6-dihydroxy-1-menthene (11) and 1-terpinen-4-ol (12) were obtained from A. sichuanensis Z. Y. Zhu (a synonym of A. jianganfeng T. L. Wu) and A. katsumadai Hayata, respectively.^{49,52} Fruit of A. oxyphylla Miq. was the source of (1R,2R)-p-menth-3-ene-1,2-diol (13).⁵³ And aerial parts of A. densibracteata T. L. Wu and Senjen yielded 3,4-dihydroxy-p-menth-1-ene (14).⁵⁰ Compounds 15-17 were three hydroxyl-1,8-cineole glucopyranosides, which were mainly isolated from rhizomes of A. galanga (L.) Willd.^{54,55} Review Works are equilibrius of Alphais species reported in the last sk. according (align article from a cryptople of distribution (a) operation and the set of the proportion and the set of the set of the set of the set o

2.2. Sesquiterpenoids

To date, 132 sesquiterpenoids were reported from Alpinia species (Fig. 2). They were divided into acyclic sesquiterpenoids (18 and 19), eremophilanes (20–40), eudesmanes (41–84), cadinanes (85– 100), guaianes (101–117), caryophyllanes (118–120), bisabolanes (121–137), humulanes (138–140), drimane (141), elemane (142), carabrane (143), oplopanane (144), and others (145–149).

Seeds of A. katsumadai Hayata produced an acyclic sesquiterpenoid, trans,trans-farnesol (18), which exerted weak neuraminidase inhibitory activity in vitro $[IC_{50} = 81.4 \mu M].^{56}$ Nerolidol (19), another acyclic sesquiterpene, was obtained from rethizoms of A. japonica.⁵⁷

Investigations on fruits of A. oxyphylla Miq. afforded 16 eremophilanes (20–35). Epinootkatol (29) and nootkatone (30) displayed insecticidal activities against larvae and adults of Drosophila melanogaster with IC₅₀ values of 11.5 μ M and 96 μ g per adult, respectively.⁵⁸ While 9ß-hydroxynootkatone (31), (11S)-12-chloronootkaton-11-ol (32), and (11R)-12-chloronootkaton-11-ol (33) displayed anti-acetylcholinesterase (AChE) activities by TLC-bioautographic assays.^{59,60} 12-Nornootkaton-6-en-11-one (35) was a novel nor-eremophilane. It showed potent anti-AChE bioactivity at 10 nM using the same TLC-bioautographic assay.⁵⁹ The rest of five eremophilanes $(36-40)$ were isolated from three different species. Eremophilen-10 β -ol (36) and eremophilen-11-ol (37) were obtained from A. intermedia Gagnep. and A. japonica (Thunb.) Miq., respectively,^{61,62} whilst

nootkatene (38), valencene (39), and dehydro-nootkatone (40) were all identified from A. oxyphylla Miq.^{59,63-65}

Among 44 eudesmane sesquiterpenoids, oxyphyllones A and B $(41$ and $42)$ were isolated from A. oxyphylla. They were the first two examples of 4,5-secoeudesmanes in the Zingiberaceae family.⁶⁶ Oxyphyllone A displayed moderate anti-AChE activity.⁵⁹ Also obtained from A. oxyphylla Miq. were compounds 43-63.^{67,68} A. *intermedia* Gagnep. was the source of intermedeol (64) and β selinene (65).⁶¹ Investigations of A. japonica (Thunb.) Miq. led to the identification of 66-75.^{21,57,69,70} Two novel trinoreudesmanes, oxyphyllanenes A (76) and B (77) were obtained from A. oxyphylla, together with four known ones $(78-81)$.^{71,72} Investigation on A. oxyphylla Miq. provided three nor-eudesmane sesquiterpenoids, oxyphyllanene C (82), (5R,7S,10S)-5-hydroxy-13-noreudesma-3 en-2,11-dione (83), and 4-methoxy-oxyphyllenone A (84) .^{67,71,73}

A new 1,10-seco-15-norcadinane sesquiterpene nominated oxyphenol A (85) was isolated from A. oxyphylla.⁶⁵ Fruits of A. oxyphylla Miq. also provided one tricyclic sesquiterpene, mustakone (86), nine nor-cadinanes, $87-94$ and 2β -hydroxy- δ -cadinol (95) .^{53,59,68,74} A. *oxymitra* K. Schum. was the source of $(-)$ - $(1R,4S)$ -8-hydroxy-13calamenenoic acid (96) .⁷⁵ Alpiniaterpene A (97) was provided by A. officinarum Hance,⁷⁶ while 4(15)-cadinene-6,10-diol (98) by A. tonkinensis Gagnep.⁵¹ Two new compounds (99 and 100) were isolated from fruits of A. oxyphylla Miq. And 100 exhibited moderate hypoglycemic activity with inhibitory rate of 11.5%, compared to 41.9% of the positive control acarbose (41.9%) at 90 μ M.⁷⁷

Rhizomes of A. japonica (Thunb.) Miq. produced alpinenone (101) , an inhibitor of AChE.^{59,60} Hanamyol (102) , containing a cyclic ether linkage, was also isolated from A. japonica (Thunb.) Miq.⁷⁸ Rhizomes of A. intermedia Gagnep. provided hanalpinol peroxide (103), isohanalpinol (104), and aokumanol (105).⁶¹ While A. intermedia Gagnep. and A. japonica (Thunb.) Miq. produced hanalpinol (106), hanalpinone (107), and isohanalpinone (108).61,79 From A. japonica (Thunb.) Miq. and A. intermedia Gagnep., furopelargones A (109) and B (110) were obtained.^{61,78,80} Later on, 110 was also found from A. formossana.⁸¹ Compounds 111-114 were four secoguaiane-type sesquiterpenes with an α , β -unsaturated butenolide. A. intermedia Gagnep. produced epialpinolide (111), whilst A. japonica (Thunb.) Miq. yielded alpinolide peroxide (112), 6-hydroxy-alpinolide (113), and alpinolide (114). $61,78,79$ A 1,10-secoguaiane sesquiterpene, (+)-mandassidion (115), and two 1,10-seco-15-norguaiane sesquiterpenes, mandassions A (116) and B (117) were obtained from fruits of A. oxyphylla Miq.⁶⁵

Caryophyllene oxide (118), caryophyllenol-I (119), and caryophyllenol-II (120) were caryophyllanes from A. galanal. In addition, caryophyllene oxide was also distributed in rhizomes of A. conchigera Griff.^{24,82}

Investigation of the aerial parts of A. densibracteata T. L. Wu and Senjen led to the isolation of two bisabolane endoperoxides (121 and 122), three bisabolane hydroperoxides (123–125), and one 3,4-dihydroxy-bisabola-1,10-diene (126).⁵⁰ Compounds 127– 137 were reported from rhizomes of A. japonica (Thunb.) Miq.⁸³

A. oxyphylla Miq. was the source of $3(12)$, $7(13)$, $9(E)$ humulatriene-2,6-diol (138).⁸⁴ While A. formossana and A. japonica produced humulene epoxideII (139).^{57,81} (9E)-Humulene-2,3;6,7diepoxide (140) was reported from the fruits of A. oxyphylla Miq. However, its relative configuration remained undetermined. It

exhibited moderate anti-AChE activity in bioautographic assay at 10 nM.59,84 Interestingly, the structure and molecular formula for 140 (CAS Registry Number: 21956-93-4) provided by Scifinder were not correct. It should be $C_{15}H_{24}O_2$ instead of $C_{14}H_{21}O_2$.

Rhizomes of A. calcarata Rosc. affored a drimane-type sesquiterpene (γ -bicyclohomofarnesal, 141),⁸⁵ and an elemane one (shyobunone, 142).⁸³ Pubescone (143) was isolated from A. oxyphylla Miq. and showed weak anti-AChE activity at the concentration of 100 μ M.⁵⁹ (-)-Oplopanone (144) and oxyphyllone F (145) were obtained from fruits of A. oxyphylla Miq.⁸⁴ (Z)-4-(2,6-Dimethylhepta-1,5-dien-1-yl)-1-methyl-cyclobut-1-ene (146) was a novel nor-sesquiterpene incorporating cyclobutene ring from A. oxyphylla Miq.⁷⁴ Seeds of A. galanga (L.) Willd produced caryolane-1,9b-diol (147), which suppressed the proliferation of four cancer cell lines of HeLa, A549, HepG2, and SMMC-7721 with IC_{50} values ranged from 252 to 378 µM.⁸⁶ A. japonica (Thunb.) Miq. yielded

alpiniol (148).⁸⁷ Compound 2-ethyl-6-isopropyl-7-hydroxymethyl naphthalene (149) was a noval naphthalene from A. oxyphylla.⁷⁷ It showed bioactive activity with the inhibitory rates of 10.3%, compare to 41.9% of the positive control acarbose at 0.9 mM.⁷⁷

Noteworthily, compounds 22–31, 34, 48–56, 58–63, 79–82, 87– 89, 117, and 129–136 exerted NO production inhibitory activities at different levels.^{58-60,65,67,71,73,83,88-90} While (10R)-13-noreudesma-4,6-dien-3,11-dione (46), (5S,8R,10R)-2-oxoeudesma-3,7(11)-dien-12,8-olide (47), (5R,7S,10S)-5-hydroxy-13-noreudesma-3-en-2,11 dione (83), and (4S)-10-nor-calamenen-10-one (90) showed potent auxo-action of NO production at 10 μ M induced by lipopolysaccharide (LPS) in microglia.⁷¹

2.3. Diterpenoids

Labdane diterpenes is undoubtedly predominant in Zingiberaceae family, notably in Alpinia genus. Almost all diterpenes are

labdanes (150–205). Only one grayanane diterpene was found (206) (Fig. 3).

 (E) -Labda-8(17),12-diene-15,16-dial (150) is widely distributed in Alpinia. It exhibited a number of bioactivities, such as

antibacterial, 91 α -glucosidase inhibition, 92 NO production inhibition,⁸⁸ antifungal,⁹³ antiglycation,⁹⁴ HIV-1 integrase, and neuraminidase inhibitory activities.⁹⁵ A. katsumadai Hayata, A. galanga (L.) Willd, and A. nigra yielded (E) -8 β ,17-epoxylabd-12-

ene-15,16-dial (151). It exhibited extensive antibacterial activities, especially against Candida guilliermondii and Candida tropicalis.^{49,91,93,96} Moreover, 151 also showed a-glucosidase

inhibitory activity with IC₅₀ value between 5 μ M and 10 μ M.⁷⁰ The α -glucosidase inhibitory activity of 151 was even much higher than the positive control, acarbose (IC₅₀ = 400 μ M),

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 $(152),$ ^{81,85,96} while (E) -labda-8(17),13-dien-15-al (153) was only obtained from A. pahangensis Ridley.⁹⁶ Flowers of A. chinensis Rosc. provided compounds 154-161.81,85,97 A. tonkinensis

Gagnep. and A. speciosa K. Schum. (the accepted name is A. zerumbet (Pers.) B. L. Burtt & R. M. Sm.) were the sources of (E) -15-nor-16-oxo-8(17),12-labdadiene (162).^{51,98} Both A. zerumbet

(Pers.) Burtt and P. M. Smith and A. pahangensis Ridley gave birth to zerumin B (163).^{96,99} (11E)-15,16-Epoxylabda-8(17),11,13-trien-16-ol (164) and (E)-15-hydroxylabda-8(17),11,13-trien-16,15-olide

(165) were found in the flowers of A. *chinensis* Rosc.⁹⁷ It is noteworthy that 164 was actually a mixture of two epimers. Rhizomes of A. calcarata Rosc. produced calcaratarins A–D (166–169) and labda-8(17),11,13-trien-15(16)-olide (170).⁸⁵ Rhizomes of A. malaccensis yielded coronarin A (171), coronarin E (172), and hedyforrestin B (173) .¹⁰⁰ Coronarin E (172) was also isolated from A.

zerumbet (Pers.) Burtt and P. M. Smith, and A. chinensis Rosc.97,99,100 Three antibacterial constituents, zerumin A (174), pahangensin B (175), and sceptrumlabdalactone B (176), were isolated from A. pahangensis Ridley.⁹⁶ Interestingly, zerumin A (174) was also obtained from A. calcarata Rosc. and A. zerumbet (Pers.) Burtt and P. M. Smith.^{85,99} Compound 175 was also found in A. japonica (Thunb.) Miq., with NO production inhibition (IC $_{50}$ $= 34.3 \mu M$) in LPS-induced RAW264.7 macrophages.¹⁰¹ Galanolactone (177) was isolated from A. katsumadai Hayata and A. galanga. It was reported to have moderate antifungal activity to Candida guilliermondii PW44 and Candida tropicalis PW30 with both MIC values of 25 $\mu\mathrm{g\,mL}^{-1}$. $^\mathrm{93}$ Isocoronarin D (178) was found in A. galanga (L.) Willd and A. calcarata Rosc., which weakly suppressed the proliferation of four cancer cells lines of HeLa, A549, HepG2, and SMMC-7721 in a concentration-dependent way with IC_{50} values ranging from 69.1 to 87.0 μ g mL⁻¹.^{64,67} Seeds of A. galanga yielded galaganin (179), which showed moderate cytotoxicity towards DU145, MCF-7, H522, and k562 cells with IC_{50} values of 8.2, 13.8, 17.8, and 16.1 μ M, respectively.¹⁰² Rhizomes of A. pinnanensis T. L. Wu et Senjen produced labda-8(17),13(14)-di-en-15,16-olide (180) and ottensinin (181).⁹⁶ A. japonica provided compounds 182–187, of which 182 and 183 were norlabdanes.¹⁰¹ Compounds 182, 185, and 186 exhibited

2,3,22,23-Tetrahydroxyl-2,6,10,15,19,23-hexamethyl-6,10,14,18-tetracosatetraene 207

Fig. 4 Triterpenoids from Alpinia species.

significant NO production inhibitory effects in LPS-induced RAW264.7 macrophages, with respective IC_{50} values of 25.9, 14.6, and 25.6 μ M, compare to 39.6 μ M of the positive control, Nmonomethyl-L-arginine (L-NMMA).¹⁰¹ Ethanol extract of A. oxyphylla Miq. provided 188, which showed moderate hypoglycemic effect with inhibitory rates of 10.0% at 60 μ M.⁷⁷ Ottensinin showed moderate antibacterial activity on the Gram-positive bacteria of *Bacillus cereus* with MIC value of 0.25 µg μ L⁻¹.⁹⁶ Alpindenosides A–D (189–192) were four labdane glycosides from A. densespicata Hayata. They didn't show cytotoxic activities against four human tumor cell lines of Hela, KB, Doay, and WiDr at 20 µM. Instead, they all exhibited moderate NO inhibitory activities with IC_{50} ranging from 30 to 49 μ M.¹⁰³ Leaves of A. flabellate provided rel-labda-12-en-15(16)-olid-7-one-8R-spiro-1'-[2S-(2,4,5-trimethoxyphenyl)-3-cyclohexene] (193), a unique labdane diterpene coupled with a phenylbutenoid.¹⁰⁴ Noralpindenosides A (194) and B (195) were two norditerpene glycosides from A. densespicata Hayata, both of which showed moderate inhibitory effects on NO production with IC_{50} values of 34.2 and 49.3 μ M, respectively.¹⁰³ (*E,E*)-15-Hydroxylabda-8(17),11,13-trien-16-al (196) and its diastereoisomer (197) from A. chinensis Rosc. may arise by direct oxygenation of (E,E) -15hydroxylabda-8(17),11,13-trien-16-al.⁹⁷ From flowers of A. chinensis Rosc., coronarin B (198) containing a seven-membered endoperoxide hemiacetal was isolated.⁹⁷ It should be noted that although the structure and its NMR and MS spectroscopic data referred to coronarin B (CAS number: 119188-38-4) in the reference, the author gave a wrong name for this compound as coronarin C (CAS number: 119188-35-1) which was previously isolated from *Hedychium coronarium*.¹⁰⁵ Galanals A (199) and B (200) were obtained from A. galanga (L.) Willd. Both compounds showed signicant antifungal activities against

Candida guilliermondii PW44 with MIC values of 12.5 µg mL⁻¹. Furthermore, galanal A exhibited potent cytotoxic activity against KB cells (IC₅₀ = 3.25 µg mL⁻¹).^{7,93} Compound 201 was a novel metabolite conjugated of labdane diterpene with chalcone from aerial parts of A. katsumadai Hayata.⁴⁹ A. pahangensis Ridley provided pahangensins A (202) and C (203) as antibacterial constituents.^{96,106} A. pahangensis Ridley produced calcaratarins D (204) and E (205), both of which were cytotoxic against human KB cells *in vitro* with IC₅₀ value of 0.21 and 0.15 μ g mL⁻¹, respectively.¹⁰⁷ From seeds of A. katsumadai Hayata, a grayanane diterpenoid was isolated and characterized as rhodomollein I (206).¹⁰⁸

2.4. Triterpenoids

Up to now, only one triterpene was found from this genus (Fig. 4). It was named as 2,3,22,23-tetrahydroxyl-2,6,10,15,19,23-hexamethyl-6,10,14,18-tetracosatetraene (207), an acyclic triterpenoid, isolated

from the seeds of A. katsumadai L.¹⁰⁹ It showed weak cholesterol acyltransferase inhibitory activity with IC_{50} value of 47.9 μ M.¹⁰⁹

3. Diarylheptanoids

A total of 143 diarylheptanoids (208–350, Fig. 5) were isolated from Alpinia species, including 66 acyclic diarylheptanoids (208–273), 11 cyclic diarylheptanoids (274–284), 50 diarylheptanoid and flavonoid conjugates (285-334), 10 dimeric diarylheptanoids (335–344), and six others (345–350).

Compounds 208–210 were isolated from rhizomes of A. officinarum Hance. They were moderate or weak NO production inhibitors.¹¹⁰ From fruits of A. $oxyphylla$, oxyphyllacinol (211) and yakuchinones A–B (212–213) were isolated, of which 211 was a NO production inhibitor, while 212 and 213 exhibited anti-tumor activities to human promyelocytic leukemia (HL-60) cells in a concentration-related manner.^{32,67} In addition, 212

also possessed insecticidal,³⁶ anti-adipocyte differentiation,¹¹¹ NO production inhibitory,⁴⁶ and cardiotonic activities.¹¹² Compounds 213-216 were also yielded by fruits of A. oxyphylla.^{113,114} Seeds of A.

blepharocalyx K. Schum. gave birth to 217-225.¹¹⁵⁻¹¹⁷ Among these compounds, 1,7-bis(4-hydroxyphenyl)-3-hydroxy-1,3-heptadien-5 one (223) signicantly inhibited platelet aggregation induced by

collagen with IC₅₀ value of 14.7 μ g mL⁻¹.¹¹⁷ (3*S*,6*E*)-Methoxy-1,7-bis(4-hydroxyphenyl)-6-hepten-5-one (224) and (3S,5S)- 3,5-dihydroxy-1,7-bis(4-hydroxyphenyl)heptane (225) showed

significant antiproliferative activities against murine colon 26-L5 carcinoma and human HT-1080 fibrosarcoma with IC_{50} values of 5.2 and 12.8 μ M, respectively.^{115,116} Both A. pinnanensis

T. L. Wu et Senjen and A. katsumadai Hayata provided (3S,5S) trans-3,5-dihydroxy-1,7-diphenyl-1-heptene (226).^{118,119} It did not showed antimycobacterial activity (MIC ≥ 64 mg L^{-1}). Instead,

it exhibited weak neuraminidase inhibitory activity $[IC_{50} =$ 29.75 \pm 8.15 μM) *in vitro*.^{56,120} (*E,E*)-5-Hydroxy-1,7-diphenyl-4,6heptadien-3-one (227), (S)-1,7-diphenyl-6(E)-hepten-3-ol (228),

Fig. 5 Diarylheptanoids from Alpinia species (continued).

and alnustone (229) were isolated from A. katsumadai Hayata with significantly neuraminidase inhibitory in vitro with IC_{50} values between 1.0 and 6.1 μ M.⁵⁶ In addition, 229 also possessed antiemetic, 121 antimycobacterial activities, 120 and significantly inhibited proliferation of Bel 7402 and L0-2 cells.¹²² Investigation of A. katsumadai Hayata also led the isolation of

Fig. 5 Diarylheptanoids from Alpinia species (continued).

compounds 230-238.49,119,121,123-125 49,119,121,123–¹²⁵ 1,7-Bis(4-hydroxyphenyl)- 1,4,6-heptatrien-3-one (239) and bisdemethoxycurcumin (240) were obtained from rhizomes of A. galanga (L.) Willd, both of which significantly inhibited the proliferation of melanoma cells and indistinctively inhibited cellular tyrosinase.¹²⁶ A planar structure of 1,7-diphenyl-5-hydroxy-6-hepten-3-one (241) was reported from A. nutans Rosc.,¹²⁷ A. rafflesiana Wall.ex.Bak.,¹²⁸ and A. officinarum Hance.¹²⁹ While its enantiomers, 5S (241a) and $5R$ (241b) counterparts, were identified from A. mutical Roxb.¹³⁰ and A. katsumadai Hayata,¹¹⁹ respectively. It was shown that a large amount of diarylheptanoids (242–276) were obtained from the rhizomes of A. officinarum Hance.^{27,131-136} 7-(3,4-Dihydroxyphenyl)-1-(4-hydroxy-3-methoxyphenyl)-4-en-3-heptanone (257) displayed moderate cytotoxicity against human tumor cell lines of HepG2, MCF-7, and SF-268. While (4E,6E)-5-hydroxy-1- (4-hydroxy-3-methoxyphenyl)-7-phenylhepta-4,6-dien-3-one (258)

showed weak cytotoxicity against two cancer cell lines of MCF-7 and T98G with IC₅₀ values of 22.68 and 4.44 μ M, respectively.¹³⁵ Meanwhile, 258–267 were proved to be inhibitors of Helicobactor pylori (Hp-Sydney and Hp-F44).¹²⁹ AO-5 (263) showed antiinflammatory activity induced by 12-O-tetradecanoylphorbol-13acetate (TPA), platelet-activating factor (PAF), and NO.110,136,137 Moreover, it exhibited very weak cytotoxic activity against human glioblastoma T98G cells (IC₅₀ = 27 μ M).¹³⁸ The acetone extract of the rhizomes of A. officinarum Hance showed 5x-reductase inhibitory effect, which was superior to the drug used in the treatment of androgen-dependent disorders. Therefore, a bioactivity-guided isolation was performed and resulted in the isolation of 263–266 which exerted 5 α -reductase inhibitory effect with IC₅₀ values ranging from 220 to 390 μ M, indicating potent usage in treating androgen-dependent diseases.¹³⁹ Besides, AO-1 (266) also showed anti-helicobacter pylori, hypolipidemic activities, and NO

production inhibitory activity.^{110,140,141} AO-2 (267) was identified as an inhibitor of prostaglandin (PG) biosynthesis and exerted antioxidant activity.142,143 It is interesting to note that dihydroyashabushiketol (264), AO-1 (266), and AO-2 (267) were firstly reported

as planar structures, and later, their absolute configurations were established as 264a, 266a, and 267a, respectively.^{136,144}

7-(4"-Hydroxy-3"-methoxyphenyl)-1-phenyl-3,5-heptadione (268) also exhibited prostaglandin biosynthesis inhibitory effect

Fig. 7 Flavonoids from Alpinia species

with IC₅₀ values of 50 μ M.¹⁴³ AO-4 (269) was found to have marked inhibitory effect on TPA-induced inflammation and antioxidant activity.142,144 6-Hydroxy-1,7-diphenyl-4-en-3-heptanone (270) was

a PAF inhibitor.¹³⁷ AO-3 (271) and (5S)-5-methoxy-1,7-diphenyl-3 heptanone (272) displayed potent inhibitory effects on TPAinduced inflammation in mice with 50% of inhibition at a dose

of 0.8–2.7 µmol per ear.¹⁴⁴ $(3R,5R)$ -1- $(4$ -Hydroxyphenyl)-7-phenyl-3,5heptanediol (273) showed significantly antiemetic effect induced by CuSO₄ with 37.7% inhibition at a dose of 50 mg kg⁻¹.^{27,145}

Investigation on seeds of A. blepharocalyx K. Schum. led to the isolation of ten cyclic diarylheptanoids $(274-283).^{115,146-148}$ Rhizomes of A. officinarum Hance provided $3,6$ -furan-7- $(4'' -$

Fig. 8 Phenolics from Alpinia species

hydroxy-3"-methoxyphenyl)-1-phenylheptane (284).¹³¹ From the seeds of A. katsumadai, 285-292 were obtained,¹⁴⁹ three of which (285–287) displayed weak antiproliferative activities against four cancer cell lines of NCI-H460, HeLa, SMMC-7721, and HCT-116 with IC_{50} values of 15.39-42.24 mM.^{115,149} A. blepharocalyx K. Schum. was the source of 293-305.^{115,148,150,151} However,

the stereochemistry at $C-9''$ of six stereoisomerics (294/295, 296/ 297, 298/299, 300/301, 302/303, 304/305) remained unsolved. Calyxin J (298), epicalyxin J (299), calyxin K (300), and epicalyxin K (301) showed marked anti-proliferative activity against human HT-1080 fibrosarcoma cells with ED_{50} values from 0.3-8.2 μ M.^{115,152} Compounds 302-305 were proved to inhibit NO production in endotoxin activated murine macrophage J774.1 with 90-94% inhibitory rate at a concentration of 100 µg mL⁻¹.¹⁵¹ Seeds of A. katsumadai Hayata provided 306-318. Calyxins Q (306) and R (307) exerted potent antiproliferative activities against four cancer cell lines of NCI-H460, HeLa, SMMC-7721, and HCT-116 at the level of IC_{50} values of 15.3-42.2 μ M.¹⁴⁹ Calyxin B (319) and epicalyxin B (320) were obtained from A. blepharocalyx K. Schum. and A. pinnanensis as NO production inhibitiors.115,151 In addition, 319 showed potent antiproliferative activity against human HT-1080 fibrosarcoma cells with an ED_{50} value of 0.69 μ M.¹⁴⁸ Both A. *pinnanensis* T. L. Wu et Senjen and A. katsumadai Hayata yielded alpinnanin B (321).118,124 From A. katsumadai Hayata and A. blepharocalyx K. Schum., epicalyxin H (322) and calyxin H (323) were isolated.^{118,124,153} Epicalyxin H was identified as NO production inhibitor.115,153 Seeds of A. blepharocalyx yielded 324– 330.^{115,152,154} It's worth mentioning that all three structures of calyxin L (325), epicalyxin F (327), and calyxins F (328) in the Scifinder were wrong. Out of a serious of diarylheptanoids bearing a chalcone or a flavanone moiety, epicalyxins $I(326)$, F (327), and calyxin F (328) were shown to possess strong antiproliferative activities toward colon 26-L5 carcinoma and HT-1080 fibrosarcoma with IC_{50} values ranging from 0.5 to 10.1 µM.^{115,150} Meanwhile, 326 and 327 were cytotoxic against human fibrosarcoma cells with IC_{50} values ranging from 0.9 to 12.1 μ M.¹⁵² 6-Hydroxycalyxin F (329) and calyxin A (330) demonstrated NO production inhibitory activities with IC_{50} values of

49 and 62 μ M, respectively.^{115,150} Rhizomes of A. pinnanensis T. L. Wu et Senjen provided deoxycalyxin A (331), alpinnanins A (332), and C (333).¹¹⁸ In addition, 331 was also found in A. ble $pharccalyx$ K. Schum.¹¹⁵ While officinin A (334) was obtained from rhizomes of A. officinarum Hance.¹⁵⁵

Five dimeric diarylheptanoids (335–339) were obtained from rhizomes of A. officinarum Hance.135,136,138,156,157 Only alpinin C (338) displayed selective cytotoxic against MCF-7 ($IC_{50} = 62.3$) μ M) and T98G cells (IC₅₀ = 57.3 μ M).¹³⁵ Seeds of A. *blepharocalyx* K. Schum. provided 340–344 possessing two diarylheptanoid units and a chalcone moiety.^{115,146,153} Both blepharocalyxins A (340) and B (341) showed concentration-dependent inhibition in the range of $1-100 \mu g \text{ mL}^{-1}$ against NO production in endotoxin-activated murine macrophages J774.1.¹⁵⁸ Blepharocalyxins C–E (342–344) were tested for antiproliferative activities against two tested cancer cells, blepharocalyxin D (343) exhibited the strongest effect against highly liver-metastatic murine colon 26-L5 carcinoma cells ($ED_{50} = 3.6 \mu M$), whereas blepharocalyxin E (344) showed the strongest activity against human HT-1080 fibrosarcoma cells (ED₅₀ = 9.02 µM).^{115,146,159} It is worth mentioning that the stereochemistry at C-I-5 position for 343 in Scifinder was S , which was not correct and should be revised as R. Moreover, the two diarylheptanoid moieties in 344 were wrongly connected through C-I-6 and C-II-5 by Scifinder. Instead, it should be joined through C-I-6 and C-II-7. Two unusual diarylheptanoid derivatives, neocalyxin A (345) and its epimer neocalyxin B (346), were found from the seeds of A. blepharocalyx K. Schum., with the stereochemistry at $C-9$ ⁿ undetermined.115,152

Rhizomes of A. officinarum Hance produced officinaruminane B (347) , a diarylheptanoid coupled with a monoterpene unit.¹³¹ Investigation on seeds of A. katsumadai Hayata identified two novel anti-emetic diaryllheptanoids, katsumadains A (348) and B

(349).¹⁶⁰ Besides, 348 also exerted promising neuraminidase inhibitory effect against human influenza virus A/PR/8/34 (IC $_{50}$ = 1.05 mM).⁵⁶ 4-Phenethyl-1,7-diphenyl-1-heptene-3,5-dione (350) was isolated from rhizomes of A. officinarum Hance. It exhibited weak antibacterial activity against Hp-Sydney and Hp-F44 with the MIC values of 23.6-31.4 and 78.5 μ M, respectively.¹²⁹

Fig. 11 The percentage of each type of compounds from Alpinia species.

Fig. 12 The number of published papers for each investigated Alpinia species on chemical constituents and their bioactivities over last six decades since 1955.

4. Lignans

Twenty-four lignans (351–374) were reported from the genus of Alpinia (Fig. 6). Separation for leaves of A. flabellata Ridley resulted in the isolation of 351–353, three phenylbutanoid dimers bearing a novel tetracyclic moiety.^{161,162} cis-1-(2,4,5-Trimethoxy-E-styryl)-2-(2,4,5-trimethoxy-Z-styryl)cyclobutane (351) and trans-1-(2,4,5-trimethoxy-E-styryl)-2-(2,4,5-trimethoxy-Zstyryl)cyclobutane (352) showed weak antibacterial against Staphylococcus aureus with MIC values of 5.0 and 2.5 mM, respectively.¹⁶¹ Furthermore, 351 signicantly decreased the ovalbumin permeability in intestinal cells.¹⁶¹

Rhizomes of A. officinarum Hance yielded 354–358 containing a rare $\beta-\gamma$ linkage. All five compounds exhibited weak antioxidant activities against the autoxidation of methyl linoleate in bulk phase.¹⁶³ Extracts of seeds of A. katsumadai Hayata afforded antiemetic katsumadin (359) with antiemetic activity on CuSO₄-induced emesis in young quail.¹²¹ Galanganol B (360) was isolated from rhizomes of A. galanga (L.) Willd.¹⁶⁴ Investigation on the whole plant of A. conchigera afforded eight rare 8– 9' linked neolignans 361-368.¹⁶⁵ Although conchigeranals D (364) and E (365) shared the same planar structure, their relative configurations were not be determined. Galanganal (366), galanganols A (367), and B (368) were also found from rhizomes of A. galanga (L.) Willd.¹⁶⁶ Compounds 361-367 exhibited significant cytotoxic activity against cancer Hela cells with IC_{50} values ranging from 1.5 to 5.29 μ g mL⁻¹.¹⁶⁵ Interestingly, 366 and 368 also inhibited NO production in mouse peritoneal macrophages.¹⁶⁶ Galanganol C (369) was obtained from rhizomes of A. galanga (L.) Willd as a NO production inhibitor.¹⁶⁶ The whole plant of A. conchigera yielded three unusual sesquineolignans, conchignans A–C (370–372) bearing a tetrahydropyrane ring.¹⁶⁷ 7-Methoxycoumarin (373) is a coumarin known from A. calcarata Rosc.⁸⁵ Open Access Article. Published on 02 March 2017. Downloaded on 1/3/2025 10:07:37 PM. This article is licensed under a [Creative Commons Attribution 3.0 Unported Licence.](http://creativecommons.org/licenses/by/3.0/) **[View Article Online](https://doi.org/10.1039/c6ra27830b)**

Citrusin B (374) and 2,3-dihydro-2- $(4-\beta)$ -p-glucopyranosyl-3methoxyphenyl)-3-hydroxymethyl-7-hydroxy-5-

benzofranpropanol (375) were the only two lignan glycosides isolated from leaves of A. speciosa.¹⁶⁸

5. Flavonoids

To date, 71 flavanoides (Fig. 7) were isolated from the Alpinia species, including seven flavones (376-382), 14 flavonols (383-396), four flavanones (397-400), seven flavanonols (401-407), two dihydrochalcones (408 and 409), 13 chalcones (410–422), four flavanols $(423-426)$, and 18 flavonoid glycosides $(427-444)$, two flavonoid oligomers (445 and 446).

Tectochrysin (376) and chrysin (377) were isolated from A. $oxyphylla$ Miq. and exhibited moderate anti-inflammatory activities against LPS-induced NO production in RAW264.7 macrophage cells.¹⁶⁹ Both A. bracteata and A. officinarum Hance produced apigenin (378), which displayed moderate activity on scavenging DPPH free radicals (EC₅₀ = 90 \pm 1.5 μ M).¹⁷⁰ A. galanga (L.) Willd was the source of 379–381 and A. tonkinensis Gagnep. produced 5-hydroxy-3',4',7-trimethoxy flavanone (382).^{102,171} Kaempferol-3,4'-dimethylether (383) was afforded by A. sichuanensis Z. Y. Zhu.⁵² Galangin (384) and kaempferide (385) were the major flavonols distributed in several plants of Alpinia, both of which exhibited inhibitory against penicillinase and potent antioxidant activities.^{113,172} In addition, galangin effectively inhibited the TPA-induced invasion and migration of HepG2 cells at concentrations of 2.5–5 μ M.¹⁷³ In 2001, a review summarized anti-genotoxic activity of galangin and demonstrated that galangin was a promising candidate for cancer chemoprevention.¹⁷⁴ Investigation on the whole plant of A. sichuanensis Z. Y. Zhu provided kaempferol $(386)^{52}$ From A. speciosa, A. galanga (L.) Willd, A. katsumadai Hayata, and A. tonkinensis Gagnep., 3-methoxykaempferol (387) was isolated.^{175–178} While A. *flabellata* Ridley, A. $oxyphylla$, and A.

tonkinensis Gagnep. yielded 3,5-dihydroxy-7,4′-dimethoxyflavone (388).^{113,161,171} Izalpinin (389) from different parts of A. oxyphylla Miq. was a NO production inhibitor and exhibited potent antioxidant activity.^{113,176} From rhizomes of A. officina rum , 3-methylethergalangin (390) was identified as an inhibitor of pancreatic lipase with an IC_{50} value of 1.3 mg mL⁻¹.¹⁷⁹ Compounds 391–395 were mainly obtained from A. tonkinensis Gagnep.¹⁷¹ 5-Hydroxy-3,7,4'-trimethoxyflavone (396) was yielded by leaves of A. *flabellata* Ridley.¹⁸⁰ Pinocembrin (397) and alpinetin (398) were distributed in several Alpinia species and both showed antiemetic activities.^{121,181} In addition, 397 also demonstrated several bioactivities, such as cytotoxicity (on human T4 lymphoblastoid cancer cells),¹⁸² anti-inflammation,¹⁶⁹ and antiplatelet aggregation etc.¹⁸³ While, 398 was a PAF receptor binding inhibitor.¹⁸⁴ 7,4'-Dihydroxy-5-methoxy flavanone (399), pinostrobin (400) were reported from several species.^{116,118,128,182} Pinobanksin (401), (2R,3S)-pinobaksin-3cinnamate (402), and 3-O-acetylpinobanksin (403) were mainly obtained from A. galanga (L.) Willd and A. katsumadai Hayata.176,177,185 Compound 402 showed potent neuroprotective effect against PC12 cells.^{177,186} Leaves of A. *flabellata* Ridley provided 404 and 405.¹⁸⁰ Dihydrokaempferol (406) were isolated from A. oxyphylla.¹⁶⁹ Both A. japonica (Thunb.) Miq. and A. galanga (L.) Willd were sources for alpinone (407) .^{176,187} From seeds of A. katsumadai Hayata, a dihydrochalcone uvangoletin (408) was isolated.¹⁰⁸ A. speciosa K. Schum. and A. formosana afforded another dihydrochalcone, dihydroflavokawin B (409).^{81,188} Flavokawin B (410) was isolated from several plants and showed strong cytotoxicity against human T4 lymphoblastoid cancer cells (IC₅₀ = 6.5 μ M) and anti-inflammatory activity.^{182,189} Cardamomin (411) distributed in many Alpinia species^{100,118,123,188,190} and exhibited extensive bioactivities including death receptor 5 $(DR5)$ promotor,¹⁷⁵ antimicrobial,¹⁹¹ antiemetic,¹²¹ anticoagulation,¹⁸³ and anti-inflammation.¹²⁸ Interestingly, it also protected septic mice from acute lung injury by preventing endothelial barrier dysfunction.¹⁹² 2',3',4',6'-Tetrahydroxychalcone (412), which was obtained from A. rafflesiana Wall.ex.Bak., was potently active to DPPH free radical scavenging $\rm (IC_{50} = 55 \ \mu M).^{128}$ Rhizomes of A. *pricei* Hayata yielded $\rm 2',4',6'$ trimethoxychalcone (413) and pinostrobin chalcone $(414).$ ¹⁸⁹ Compounds 415–417 were isolated from the seeds of A. ble $pharocalyx$ K. Schum.,^{116,117} while helichrysetin (415) was also found in A. katsumadai Hayata.¹⁰⁸ Pinocembrin chalcone (418) and 4',6'-dimethylchalconaringenin (419) were provided by A. katsumadai Hayata and A. pinnanensis T. L. Wu et Senjen, respectively.118,181 Compound 418 was also isolated from A. platychilus.¹⁹³ Galanganones A-C (420-422) were three novel chalcones bearing a long-chain alkylphenol from A. galanga.¹⁹⁴ Whilst A. katsumadai Hayata and A. zerumbet (Pers.) B. L. Burttet Smith. provided (+)-catechin (423).^{195,196} Epicatechin (424) and galloepicatechin (425) were yielded by A. oxymitra K. Schum.⁷⁵ (+)-Epicatechin (426) was isolated from A. speciosa K. Schum. and displayed antioxidant activity.¹⁹⁷ Kaempferide-3-O-β-Dglucoside (427) from A. officinarum Hance had an weak inhibitory activity against penicillinase.¹⁷² Study on A. speciosa K. Schum. lead to the isolation of 428-432.¹⁹⁸ Quercetin 3-O-robinobioside (433) and galangoflavonoside (434) were obtained Review Webcomes Cagarep. yielded 3,5-dlinydcox-7,4" dimetabor March 2017. The matrix of the computer of the c

from A. katsumadai Hayata and A. galanga (L.) Swartz., respectively.^{196,199} Compounds 435-437 from A. densespicata Hayata exhibited moderate NO inhibitory activities.¹⁰³ Compounds 438–440 were obtained from the seeds of A. katsumadai Hayata and isorhamnetin-3-O- β -D-galactosyl- $(6 \rightarrow 1)$ - α -L-rhamnoside (441) was isolated from rhizomes of A. tonkinensis Gagnep.^{51,196} Leaves of A. zerumbet (Pers.) B. L. Burttet Smith. contained rutin (442) and kaempferol-3-O-rutinoside (443).¹⁹⁵ The whole plant of A. sichuanensis Z. Y. Zhu yielded hesperidin (444).⁵² Two pairs of enantiomers of flavonoidoligomers (445a and 445b, 446a and 446b) were found from rhizomes of A. platychilus. The compounds mixture of 446a and 446b showed anticoagulant activity on the prolongation of both prothrombin times (PT) and the thrombin times (TT) with a dose-effect relationship at 6.25– 100 mM.¹⁹³

6. Phenolics

A total number of 66 phenolics (447–512) were obtained from Alpinia species (Fig. 8). [Di- $(p$ -hydroxy-cis-styryl]]methane (447) was obtained from A. galanga (L.) Willd.²⁰⁰ Whist alpininone (448) was isolated from A. gagnepainii K. Schum. with antibacterial effect against E. coli, B. subtilis, and S. aureus with the same MIC value of 12.5 μ g mL⁻¹.¹⁹¹ (1E,4Z)-5-Hydroxy-1phenylhexa-1,4-dien-3-one (449) and 2-propenal, 3-[4- (acetyloxy)-3-methoxyphenyl] (450) were provided by A. katsumadai Hayata and A. galanga (L.) Willd, respectively.^{86,108} From A. sichuanensi and A. oxyphylla, dibutyl phthalate (451) was isolated.^{52,201} Two compounds named as (E) -p-coumaryl alcohol (452) and (E) -p-coumaryl alcohol γ -O-methyl ether (453) exhibited potent inhibitory activities against the autoxidation of methyl linoleate in bulk phase.¹⁶³ In addition, compound 453 exerted potent cytotoxic activity against the SNU638 cells with IC₅₀ value of 1.62 µg mL⁻¹.²⁰²

A. galanga (L.) Willd and A. conchigera Griff. produced transp-hydroxycinnamaldehyde (454) and trans-p-hydroxycinnamyl acetate (455).170,203 Compound 454 displayed weak antiallergic effect,²⁰⁴ and NO production inhibitory activities (IC₅₀ = 20 μ M),¹⁶⁶ and 455 exerted no inhibitory activity towards Staphy $lococcus aureus strain VISA (MIC = 203 mM).⁸² trans-p-Coumaryl$ alcohol (456) was a weak NO production inhibitor from A. galanga (L.) Willd (IC₅₀ = 72 µM).¹⁶⁶ trans-p-Coumaryl diacetate (457) from A. galanga showed a number of bioactivities, including anti-allergy,²⁰⁴ efflux pump inhibition,²⁰⁵ NO production inhibition,¹⁶⁶ xanthine oxidase inhibition,²⁰⁶ antileishmania,¹⁶⁴ cytotoxicity,²⁰³ and antibacteria.⁸² trans-p-Acetoxycinnamyl alcohol (458), trans-p-hydroxylcinnamaldehyde acetate (459), and p -coumaric acid (460) were obtained from rhizomes of A. galanga (L.) Willd.^{164,205} In addition, compound 460 was also distributed in A. galanga (L.) Willd,¹⁶⁴ A. sichuanensis Z. Y. Zhu,⁵² A. speciosa,¹⁹⁸ A. blepharocalyx K. Schum.,¹¹⁶ and A. oxyphylla.¹⁶⁹ Both A. formosana and A. speciosa K. Schum. were sources of methyl trans-cinnamate $(461).$ ^{81,188} Seeds of A. blepharocaly yielded methyl p-hydroxycinnamate (462) and methyl p-hydroxycinnamyl ketone (463).¹¹⁶ From rhizomes of A. galanga (L.) Willd, 12 compounds (464–475) were obtained.^{33,82,203,207} Among them, 1S-1'-acetoxychavicol acetate (464) and 1-acetoxyeugenol acetate (465) were the most

abundant phenylpropanoids presented in A. galanga (L.) Swartz., A. officinarum Hance, and A. conchigera Griff. They were reported to have anti-ulcer,²⁴ antileishmanial,¹⁶⁴ and antitumor bioactivities,^{33,202,208} Furthermore, 464 also showed antiallergic,²⁰⁴ efflux pump inhibitory,²⁰⁵ NO production inhibitory,¹⁶⁶ xanthine oxidase inhibitory,²⁰⁶ gastroprotective,²⁰⁹ anti-HIV,²¹⁰ anti-cancer,⁸⁶ antibacterial,^{30,211} plant growth-inhibitory and fungal growthinhibitory activities.²¹² Two compounds, methyleugenol (466) and hydroxychavicol acetate (467), were isolated from A. galanga (L) Willd.^{82,164,166,204,211} It was demonstrated that 467, a chavicol acetate analogue, suppressed T-bet expression in Th cells.²¹¹ Besides, 467 also showed weak antibacterial activity against Staphylococcus aureus strain VISA (MIC = 0.8 mM).⁸² trans-Coniferyl diacetate (468) was proved to be a xanthine oxidase inhibitor.²⁰⁶ Three new phenolics 469, 470, and 471, along with four known ones 472-475 were also yielded by A. galang.^{33,203,207} Chavicol acetate (476) and 1'S-acetoxyeugenol acetate (477) were two known phenolics found from A. conchigera Griff.⁸² Compound 477 possessed antibacterial,⁸² xanthine oxidase inhibitory,²⁰⁶ gastroprotective,²⁰⁹ and anti-cancer activities.⁸⁶ Investigation on leaves of A. *flabellata* Ridley provided 478-480, with strong antibacterial activities against Staphylococcus aureus.^{161,162,180} Compounds 481-489 were nine phenolic acids isolated from several Alpinia species.^{52,89,122,180,198,213,214} Protocatechuic acid (489) showed potent neuroprotective effect on MPP⁺-induced neurotoxicity and H_2O_2 induced oxidative damage in PC12 cells.²¹⁵⁻²¹⁹ In addition, it also exerted anti-aging effect on spleen and liver antioxidative system of senescent mice.³¹ 4-Hydroxybenzaldehyde (490), isolated from A. sichuanensis Z. Y. Zhu, A. blepharocalyx K. Schum., A. bracteata, and A. galanga (L.) Willd, $52,116,117,166,170$ didn't show any DPPH radical-scavenging activity. Instead, it exhibited inhibitory activity on xanthine oxidase (IC₅₀ = 19.6 µM).^{170,206} Compounds 491-496 were provided by several Alpinia plants.^{52,84,137,167,220} BSC Arbaness Securities are article in A gelaxie Common Common Common A System Common Co

Ethyl 4-O-feruloyl-b-glucopyranoside (497) and 4-hydroxy-3 methoxyphenyl 4-O-feruloyl-β-glucopyranoside (498) were two new glucoside esters of ferulic acid from rhizomes of A. speciosa, both of which showed antioxidant activities.¹⁹⁷ Investigation on rhizomes of A. officinarum Hance yielded 499-506.⁵⁵ While from rhizomes of A. bracteata, a new phenolic glycoside (507) was isolated and showed moderate antioxidant activity on scavenging DPPH free radicals ($EC_{50} = 169 \pm 4.8 \,\mu\text{M}$).¹⁷⁰ Leaves of A. speciosa K. Schum. provided coniferin (508) and syringin $(509).$ ¹⁶⁸

Dihydro-5,6-dehydrokawain (510) and 5,6-dehydrokawain (511) were major chemical constituents in several Alpinia species.^{81,100,128,175,188,221} They showed antiulcerogenic, antithrombotic,¹⁹⁵ antifungal,¹⁹¹ anti-obesity,²²² and plant growth inhibitory activities.²²³ Recently, it was reported that they could strongly inhibit HIV-1 integrase with respective IC_{50} values of 4.4 and 3.6 μ g mL⁻¹. In addition, they exhibited mixed type of inhibition against neuraminidase with both IC_{50} values of 25 μ M.⁹⁵ Furthermore, 511 was also reported as a slow and timedependent reversible inhibitor of neuraminidase, a moderated antioxidant, a strong inhibitor of skin diseases-related enzymes, and strong antiplatelet inhibitor.^{95,127,224} Interestingly, a dimer of 5,6-dehydrokawain, AS-II (511a), was an artifact formed by photo-irradiation during the isolation procedure of A. speciosa

K. Schum. leaves.²²³ 4-Hydroxy-5,6-dehydrokawain (512) was an a-pyrone isolated from A. blepharocalyx K. Schum. It displayed antiproliferative activity against murine colon 26-L5 carcinoma and human HT-1080 fibrosarcoma with ED_{50} 20.7 and 20.1 μ M, respectively.116,117 It also showed inhibitory effect on platelet aggregation induced by collagen, arachidonic acid (AA), adenosine diphosphate, and ristocetin.⁹⁶

7. Steroids

Seven steroids (Fig. 9) were isolated from Alpinia species including four cholestanes (513–516) and three sitosterol glycosides (517–519).^{27,52,89,116,118,225} As it is the same in plants of the other genera, β -sitosterol (513) and stigmasterol (514) were also widely distributed in Alpinia species.^{52,82,89,118,178,191,226-228} β -Sitosterol-3-O-β-D-6-palmitoylglucoside (518) showed potent antiemetic activity induced by $CuSO₄.²⁷$

8. Alkaloids

Officinaruminane A (520) and officinine B (521), two alkaloids of bi-diarylheptanoid connecting by a pyridine ring were contributed by rhizomes of A. officinarum Hance.^{131,157} A study on seeds of A. katsumadai Hayata afforded another six alkaloids (522–527) (Fig. 9).108,196

9. Stilbenes

Six stilbenes, 528–533 (Fig. 9), were all isolated from aerial parts of A. katsumadai Hayata.^{29,121}

10. Others

One esters (534) and three fatty acids, 535–537, were isolated from several *Alpinia* species.^{64,227-229} (S)-2-Pentanol-2-O-β-D-glucopyranoside (538), which showed inhibitory effect on NO production from LPS-activated RAW264.7 macrophage cells, was obtained from fruits of A. oxyphylla.⁸⁹ Two glycosides known as 3methyl-but-2-en-1-yl- β -D-glucopyranoside (539) and *n*-butyl- β -Dfructopyranoside (540) were isolated from A. officinarum Hance.^{55,230} While 541-544 were found in different Alpinia species (Fig. 9).24,51,108,196,201 Interestingly, 5-hydroxymethylfurfural (544) exerted memory improvement activity against Alzheimer's disease (AD) by mitigating the degree of neuronal damage.²³¹

11. Conclusions

The number of publications on the chemical constituents and their bioactivities for Alpinia species from 1955 to 2015 are shown in Fig. 10. Before 1999, fewer investigations (less than five per year, except six in 1987) were performed on this genus. However, after 2009, there were more than 10 papers published for each year. In 2013, the number of published articles reached 26, indicating a growing interest in the genus of Alpinia.

Till 2015, investigations on chemical constitutes of the Alpinia species afforded a total of 544 compounds, including 207 terpenoids, 143 diarylheptanoids, 25 phenylpropanoids, 71

flavanones, 66 phenolics, seven steroids, eight alkaloids, six stilbenes, and 11 others (Fig. 11). Among 207 terpenoids, 17 are monoterpenoids, 132 are sesquiterpenoids, 57 are diterpenoids, and the rest one is a triterpenoid. For sesquiterpenoids, eudesmanes and eremophilanes are undoubtedly predominant with 44 and 21 components, respectively. While for diterpenoids, almost all are labdanes.

Amongst 544 isolated compounds from the genus of Alpinia, 247 are new ones (Table 1), including 96 diarylheptanoids and 106 terpenoids. Obviously, diarylheptanoids, especially diarylheptane–flavonoids conjugates, are characteristic components for the genus of Alpinia.¹⁴⁹

The crude extracts of Alpinia species and their chemical constituents were found to possess various biological activities. Mainly reported were antiemetic,^{26,27} antibacterial,^{29-31,37,82,232-236} antioxidant,^{127,237–239} anticancer,^{32–34,240–245} anti-inflammatory,^{189,246,247} insecticidal,^{36,164} and neuroprotective bioactivities.^{38,39,231,248-250}

In addition, they also showed antiulcer,²⁵ antiplatelet, $117,183$ hepatoprotective,²⁵¹ and hypolipidemic effects.²⁵² Meanwhile, evidences showed that ethanol extract of A. galangal can retard lipid oxidation for minced beef, indicating a great potential utility for food storage.⁸ What should be aroused considerable interest was the promising anticancer and hepatoprotective properties, which could be a great potential to be developed as herbal medicines.

Although there are about 230 species for the Alpinia genus, only 35 were investigated for their chemical constituents and bioactivities (Fig. 12), because A. jianganfeng T. L. Wu includes Alpinia sichuanensis Z. Y. Zhu, and A. zerumbet (Pers.) B. L. Burtt & R. M. Sm. includes A. speciosa K. Schum. according to The Plant List. Among these species, A. galanga, A. oxyphylla, A.

officinarum, and A. katsumadai are four most studied plants with referenced papers of 43, 40, 32, and 23, respectively. While for the rest of 31 species, only very fewer articles were published, most of which were less than five. As a matter of fact, there was even only one paper published for 18 species. Although this genus contributed a diverse array of bioactive compounds, the potential of Alpinia species remains virtually untapped. Thus, much attention should be paid to Alpinia species on further phytochemical and pharmacological studies, which would produce structurally interesting and biologically active compounds with potential use in agricultural and medicinal applications. In addition, although most of *Alpinia* species were also used as edible plants, the nutritious components and their effects were seldom investigated, which could be a hotspot in the near future.

Acknowledgements

The project was supported by National Natural Science Foundation of China (41176148, 21372233, 21202080).

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