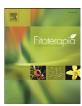


Contents lists available at ScienceDirect

Fitoterapia

journal homepage: www.elsevier.com/locate/fitote



Review

Chemical study and medical application of saponins as anti-cancer agents

Shuli Man^a, Wenyuan Gao^{a,*}, Yanjun Zhang^b, Luqi Huang^c, Changxiao Liu^d

- ^a School of Pharmaceutical Science and Technology, Tianjin University, Weijin Road, Tianjin 300072, China
- ^b Tianjin University of Traditional Chinese Medicine, Tianjin, China
- c Institute of Chinese Materia Medica, China Academy of Chinese Medical Sciences, Institute of Pharmaceutical Research, Beijing, China
- ^d The State Key Laboratories of Pharmacodynamics and Pharmacokinetics, Tianjin, China

ARTICLE INFO

Article history: Received 20 January 2010 Accepted in revised form 28 May 2010 Available online 13 June 2010

Keywords: Saponins Distribution Mechanism Structure–activity

ABSTRACT

Saponins are a group of naturally occurring plant glycosides, characterized by their strong foam-forming properties in aqueous solution. The presence of saponins has been reported in more than 100 families of plants out of which at least 150 kinds of natural saponins have been found to possess significant anti-cancer properties. There are more than 11 distinguished classes of saponins including dammaranes, tirucallanes, lupanes, hopanes, oleananes, taraxasteranes, ursanes, cycloartanes, lanostanes, cucurbitanes and steroids. Due to the great variability of their structures, saponins always display anti-tumorigenic effects through varieties of antitumor pathways. In addition, there are a large amount of saponins that still either remain to be trapped or studied in details by the medicinal chemists. This article reviews many such structures and their related chemistry along with the recent advances in understanding mechanism of action and structure-function relationships of saponins at the molecular and cellular levels. These aglycones have been described and their classification and distribution have been listed in the review. Some special saponins with strong antitumor effects have also been exhibited. Ginsenosides, belonging to dammaranes, have been found beneficial targeted on inhibition of tumor angiogenesis by suppressing its inducer in the endothelial cells of blood vessels, and then on prevention of adhering, invasion, and metastasis of tumor cells. Dioscin, one of the steroidal saponins, and its aglycone diosgenin also have been extensively studied on its antitumor effect by cell cycle arrest and apoptosis. Other important molecules discussed include oleanane saponins such as avicins, platycodons, saikosaponins, and soysaponins along with tubeimosides.

© 2010 Elsevier B.V. All rights reserved.

Contents

	Introduction
	Mechanisms of the antitumor effect of saponins
٥.	3.1. Cycloartanes
	3.2. Dammaranes
	3.3. Oleananes
	3.4. Spirostanes
	3.5. Furostanes
4	Structure–function relationship of saponins with the antitumor properties 70

E-mail address: pharmgao@tju.edu.cn (W. Gao).

^{*} Corresponding author.

4	4.1.	Influenc	e of the aglycone on the antitumor activities of saponins
		4.1.1.	The site of the hydroxyl group
		4.1.2.	The number of the hydroxyl group in aglycone
		4.1.3.	Else
4	4.2.	Influenc	e of the sugar side chain on the antitumor activities of saponins
		4.2.1.	The sugar linkage
		4.2.2.	The lipophilicity of the sugar
		4.2.3.	The number of the sugar
		4.2.4.	The kinds of sugar sequence
5. (Conclu	ısion and	perspective
Ackno	wledg	gement.	
Refere	ences		

1. Introduction

Saponins are common in a variety of higher plants and usually found in roots, tubers, leaves, blooms or seeds. Based on the carbon skeletons, saponins were classified into triterpenes and steroids. Their glycone parts were mostly oligosaccharides, arranged either in a linear or branched fashion, attached to hydroxyl groups through an acetal linkage [1].

Modern research found that saponins have antitumor effect on many cancer cells. Several saponins inhibit tumor cell growth by cell cycle arrest and apoptosis with IC50 values up to 0.2 mM. Meanwhile, saponins in combination with conventional tumor treatment strategies, result in improved therapeutic success. Furthermore, a much clearer understanding of how the various saponin structures are related to each other is obtained with the use of the classification presented [2]. The objective of this review is to provide a timely update on the sources, classification, and applications

of saponins with special focus on their mechanism of antitumor effect and structure–function relationship.

2. Source and classification

The percentage of saponins has been reported in more than 100 families of plants, out of which at least 150 kinds of natural saponins have been found to possess significant anticancer properties (Tables 1 and 2).

The steroidal saponins are mainly found in Agavaceae, Dioscoreaceae, Liliaceae, Solanaceae, Scrophulariaceae, Amaryllidaceae, Leguminosae and Rhamnaceae; while triterpene saponins are predominantly present in Acanthopanax, Leguminosae, Araliaceae, Scrophulariaceae, Campanulaceae and Caryophyllaceae. There are more than 11 mainly distinguished classes of saponins including dammaranes, tirucallanes, lupanes, hopanes, oleananes, taraxasteranes, ursanes, cycloartanes, lanostanes, cucurbitanes, and steroidals. Among

```
Notes to Table 1:
```

^aAra α-L-arabinofuranosyl, Fuc β-D-fucosyl, Gal β-D-galactosyl, Gla α-galactosidase, Glc β-D-glucopyranosyl, Rha α-L-rhamnopyranosyl, Xyl β-D-xylosyl.

bHuman promyelocytic acute myelogenous leukemia: HL-60.

Human chronic myelogenous leukemia cells: K562, THP-1, U937.

Lymphocytic leukemia cell: P388. Murine leukemic: L1210, P388.

Human oral squamous cell carcinoma: HSC-2. Central nervous system (CNS) cancer line; U251.

Nasopharynx: CNE-2Z, KB.

Non-small cell lung cancer (NSCLC): A549, LA795, LL/2.

Human breast ductal carcinoma: BT-549.

Breast adenocarcinoma: Bcap37, MCF7, MDA-MB-231, NCI-ADR-RES.

Human gastric adenocarcinoma: MK-1. Stomach cancer cells: SGC-7901.

Pancreas carcinoma: MIA PaCa-2. Hepatoma: BEL-7402, HA22T, SK-Hep-1.

Colon carcinoma: Colo-205, 26-L5, DLD-1, HT-29, KM-12.

Laryngeal epidermoid: Hep-2. Human oral epidermoid carcinoma: KB. Glioblastoma: U251MG, U373, U87MG.

Malignant melanoma: A375, B16BL6, B16F10, H1477, HTB-140, LOX, MALME-3M, M14, M4 Beu, SK-MEL.

Glioma: GBM8401/TSGH.

Human fibrosarcoma: HT-1080.

Non-cancer mouse 3T3 fibroblasts; human skin fibroblasts: HSFs.

Human monocytic: THP-1.

Renal carcinoma cell: 786-0 and UO-31.

Human ovary carcinoma: A2780, HO-8910, OVCAR3, SK-OV-3.

Ovarian teratocarcinoma: PA 1.
Uterine cervix cancer: HeLa.
Prostatic adenocarcinoma: PC 3.
References cited in this table [3–43].

these saponins, cycloartanes, dammaranes, oleananes, lupanes and steroids showed strong antitumor effect on kinds of cancers.

3. Mechanisms of the antitumor effect of saponins

Some special saponins with strong antitumor effects have been exhibited.

Table 1The steroid saponins in the natural plants.

3.1. Cycloartanes

Cycloartane saponins displayed slight anti-cancer effect but they could be used as chemotherapeutic agent in the treatment of tumors. For example, total Astragalus saponins (AS) (Fig. 1 and Table 3) [58] possess antitumor properties in human colon cancer cells and tumor xenografts. They down-regulated expression of the HCC tumor marker α -fetoprotein and suppressed HepG2 cell growth by inducing apoptosis and

Family	Species	Saponinsa	Kinds of cancer ^b	Ref.
	Agave utahensis		HL-60	[3]
Agavaceae	Agave fourcroydes	Chlorogenin hexasaccharide	HeLa	[4]
	Agave jourcroydes	26- <i>O</i> -Glc- 5α -furost-25(27)-ene-3 β ,22,26-tet		[4]
		raol-3-0-Glc($1\rightarrow 2$)[Glc($1\rightarrow 3$)]-Glc($1\rightarrow 4$)-Gal	SF-268	
		26-O-Glc-5β-furost-20(22)-25(27)-dien-3		[5]
Alliaceae	Allium macrostemon Bunge	β ,12 β ,26-triol-3- θ -Glc(1 \rightarrow 2) Gal	NCI-H460, SF-268	
		p,p,	HepG2, MCF-7, NCI-H460,	[6]
		Macrostemonoside O, P, Q and R	SF-268, R-HepG2	[6]
			51 250, K 11cpG2	
	Allium leucanthum C. Koch	Leucospiroside A	A549, DLD-1	[7]
Asclepiadac	Cynanchum auricula		A549	[8]
iscreptuduci	wiyriopteron extensu		Eight cancer cell lines	[9]
	Asparagus racemosus V		HepG2	[10]
	Asparagus filicinus	Filiasparosides A-D	A549. MCF-7	[11]
		Aspafiliosides A and B	110 10, 11101	1
	Asparagus oligoclono	S Aspaoligonins A and B	Five human tumor cell lines	[12]
Asparagaceo	ae	Asparanin A		
1 0	Asparagus gobicus	3-O-[Xyl(1-4)-Glc(1-2)-Glc]- (25S)-5β-spirostan-3β-ol HC		[13]
		Methyl protodioscin (NSC-698790)	HepG2	[14–1
	Dioscorea collettii	Methyl protogracillin (NSC-698792)	KM12,U251,MALME-3M&M14,7	[17
	var. hypoglauca		86-0&UO-31,MDA-MB-231	
		Protoneodioscin, protodioscin, protoneogracillin, protogra	acillin, K562	[18
		Prosapogenin A of dioscin, dioscin and gracillin	K562	[19
		Prosapogenin B of dioscin	K562	[20
Dioscoreace		(25S)-spirost-5-en-3β,		
	futschauensis	27-diol-3-0-[Rha(1 \rightarrow 2)-Glc(1) \rightarrow 3]-Glc, prosapogenin A	ts-FT210	[21
	R.Kunth	of dioscin, dioscin and gracilin		
	Dioscorea	dioscoreside E, protodioscin	Many	[22
	panthaica	dioscoresides C, D, pseudoprotodioscin, pregnadienolone	A375, L929, HeLa	[23]
	parientica	3- <i>O</i> -β-gracillimatriose, pregnadienolone 3- <i>O</i> -β-chacotrioside		
	Dracaena draco	Icodeside	HL-60, A-431	[24
Dracaenace	ae	Draconins A-C	HL-60	[25]
	Nam ginseng (roots and		26-L5, HT-1080, B-16 BL6	[26
	of Dracaena angustifoli Paris polyphylla var. yunnanei		Lung liver cancer	[27,2
	Paris formosana Hayata	Formosanin C	Lung, liver cancer HT-29	[27,2
	Anemarrhena asphodeloides	Timosaponin A-III	H1-29 HeLa	[30]
	Polygonatum sibiricum	Neosibiricosides A-D,	MCF-7	[31]
	Sansevieria ehrenbergii	Sansevistatin 1, 2	P388	[32]
	Camassia cusickii	TGHS-1, TGHS-2	L1210	[33]
Convallaria		Smilacinoside A, funkioside D, aspidistrin K562	L1210	[34]
- Convananta	atropurpurea		SK-MEL, KB, BT-549, SK-OV-3,	
	ut op ut pur eu	Atropurosides B, F; dioscin	HepG2, non-cancerous Vero cells	[35]
	Convallaria majalis L	. Convallamaroside	Human kidney tumor cells	[36]
	Solanum torvum	Torvosides M	Human cancer cell lines	[37]
	Solanum nigrum	Degalactotigonin	HepG2, NCI-H460, MCF-7, SF-268	[38]
Solanaceae	3	protodioscin, methyl protodioscin, methyl	Colo-205, KB, HeLa, HA22T,	
	Solanum indicum L.	protoprosapogenin A of dioscin, dioscin	Hep-2, GBM8401/TSGH, H1477	[39]
	Cestrum	(25R)-2α,17α-dihydroxyspirost-5-en-3β-yl	HSC-2 cells and normal	[40]
	nocturnum	O -Glc- $(1\rightarrow 2)$ - O -[Xyl- $(1\rightarrow 3)$] - O -Glc- $(1\rightarrow 4)$ -Gal	Human gingival fibroblasts	[40]
		(25R)-26-O-Glc-5-furostan-3β,22α,26-triol 3-O-	U937, MCF7,	[41]
	Tribulus parvispinus	$\{Gal-(1\rightarrow 2)-O-[Xyl-(1\rightarrow 3)]-O-Glc-(1\rightarrow 4)-Gal\}$	and HepG2	[41]
Zygophyllac	reae	Gitonin	-	
	Tribulus terrestris	Saponins	SK-MEL, KB, BT-549, SK-OV-3	[42]
	Balanites aegyptiaca ke		A549, U373	[43]

modulating an ERK-independent NF-KB signaling pathway [113]. In addition, AS could be used as an adjuvant in combination with other orthodox chemotherapeutic drugs to reduce the side effects of the latter compounds [114]. It would target at NSAID-activated gene (NAG-1) to reduce the additive effects when used along with PI3K-Akt inhibitors. The information obtained could facilitate future development of a novel target-specific chemotherapeutic agent with known molecular pathways [115].

3.2. Dammaranes

Most dammarane saponins showed anti-cancer effect. The naturally occurring compound OSW-1 (Fig. 2) is found in the bulbs of Ornithogalum saudersiae and is highly cytotoxic against tumor cell lines. Nonmalignant cells were statistically significantly less sensitive to OSW-1 than cancer cells, with concentrations that cause a 50% loss of cell viability 40-150fold greater than those observed in malignant cells. Electron microscopy and biochemical analyses revealed that OSW-1 damaged the mitochondrial membrane and cristae in both human leukemia and pancreatic cancer cells, leading to the loss of transmembrane potential, increase of cytosolic calcium, and activation of calcium-dependent apoptosis [116].

3.3. Oleananes

Oleananes own most kinds of saponins in the nature. Their antitumor effect worked through various pathways, such as anticancer, anti-metastasis, immunostimulation, chemoprevention and so forth. Avicins, tubeimoside, saikosaponins, platycodigenins, soybean saponin and Pulsatilla koreana saponins showed anti-cancer effect through different signaling transductions. In addition, all of them except tubeimoside and Pulsatilla koreana saponins displayed immunostimulation. Saikosaponins, platycodigenins and soybean saponin also have anti-metastatic activity. The detailed mechanisms of saponins listed as following.

Avicins (Fig. 3), derived from the Cactus plant Acacia victoriae found in Australia's deserts [117], can dephosphorylate Stat3 in a variety of human tumor cell lines and lead to a decrease in the transcriptional activity of Stat3, which regulated proteins such as c-myc, cyclin D1, Bcl2, survivin and VEGF [69]. Avicins D and G, as the main components of avicins, induced growth inhibition of human T lymphocytes, promoted apoptosis [118] and triggered autophagic cell death [119]. Meanwhile, they decreased respiratory activity [120] and induced ATP efflux after inhibition of the voltage dependent anion channel in the outer mitochondrial membrane [121].

Tubeimoside I (Fig. 4), one of the triterpenoid saponins from the bulb of Bolbostemma paniculatum (Maxim) Franquet, appears to be a promising agent for cancer chemoprevention [122]. It exerts cytotoxicity in HeLa cells through both mitochondrial dysfunction and ER stress cell death pathways [69]. As an anti-microtubule agent, it can bind to the colchicine binding site of tubulin [123].

Saikosaponin A (Fig. 5) activates ERK together with its downstream transcriptional machinery mediated p15(INK4b) and p16(INK4a) expression that led to HepG2 growth inhibition [124]. It inhibited the proliferation or viability of the MDA-MB-231 and MCF-7 cells in a dose-dependent manner and caused an obvious increase in the sub-G1 population of cell cycles [125]. Treatment with saikosaponin D (Fig. 5) decreased the cell proliferation of Hep G2 and Hep 3B cells in a dosedependent manner. It therefore decreased the cell proliferation and inducted apoptosis both in p53-positive Hep G2 and p53negative Hep 3B cells [126]. In addition, it inhibited the proliferation of A549 by inducing apoptosis and blocking cell cycle progression in the G1 phase [127].

Notes to Table 2:

"Ara α -L-arabinofuranosyl, Fuc β -D-fucosyl, Gal β -D-galactosyl, Gla α -galactosidase, Glc β -D-glucopyranosyl, Rha α -L-rhamnopyranosyl, Xyl β -D-xylosyl.

^bHuman promyelocytic acute myelogenous leukemia: HL-60.

Human chronic myelogenous leukemia cells: K562, THP-1, U937.

Lymphocytic leukemia cell: P388. Murine leukemic: L1210, P388.

Human oral squamous cell carcinoma: HSC-2. Central nervous system (CNS) cancer line; U251.

Nasopharynx: CNE-2Z, KB.

Non-small cell lung cancer (NSCLC): A549, LA795, LL/2.

Human breast ductal carcinoma: BT-549.

Breast adenocarcinoma: Bcap37, MCF7, MDA-MB-231, NCI-ADR-RES.

Human gastric adenocarcinoma: MK-1. Stomach cancer cells: SGC-7901.

Pancreas carcinoma: MIA PaCa-2. Hepatoma: BEL-7402, HA22T, SK-Hep-1.

Colon carcinoma: Colo-205, 26-L5, DLD-1, HT-29, KM-12.

Larvngeal epidermoid: Hep-2. Human oral epidermoid carcinoma: KB. Glioblastoma: U251MG, U373, U87MG,

Malignant melanoma: A375, B16BL6, B16F10, H1477, HTB-140, LOX, MALME-3M, M14, M4 Beu, SK-MEL.

Glioma: GBM8401/TSGH.

Human fibrosarcoma: HT-1080.

Non-cancer mouse 3T3 fibroblasts; human skin fibroblasts: HSFs.

Human monocytic: THP-1.

Renal carcinoma cell: 786-0 and UO-31.

Human ovary carcinoma: A2780, HO-8910, OVCAR3, SK-OV-3.

Ovarian teratocarcinoma: PA 1.

Uterine cervix cancer: HeLa.Prostatic adenocarcinoma: PC 3.

Macrophage-like cell line, J-774.1. References cited in this table [30,44-112].

Table 2 Triterpenoid saponins in plants.

Family	Species	Saponin ^a	Kinds of cancer ^b	Ref.
	Alternanthera philoxeroide		SK-N-SH, HL60	[44]
Amaranthacaa		Achyranthoside H methyl ester	MCF-7, MDA-MB-45	[45]
Amaranthaceae	Achyranthes fauriei	Chikusetsusaponin IV a	SK-Hep-1	[46]
	Physospermum	Saikosaponin a	эк-пер-1	[40]
Apiaceae	verticillatum	Buddlejasaponin IV, songarosaponin D	COR-L23	[47]
(Umbelliferae)		Potundifoliosidos A I	MK-1, HeLa,	[4/]
Ombenijerae)	Bupleurum rotundifoliun	Rotundiosides A-J	B16F10	[48]
			DLD-1, PA 1, A549, MCF7, PC 3,	1500000
Araliaceae	Hedera colchica K. Koch	Hederacolchisid A1	M4 Beu vs normal human fibroblasts	[49]
arrace are	Hedera helix	α-hederin	B16, 3T3 fibroblasts	[50]
	Meryta denhamii	Echinocystic acid as the aglycone	J774.A1, HEK-293, and WEHI-164	[51]
l»	eris sonchifolia	Ixeris saponins B, C	A375, L929, and HeLa	[52]
	ilphium radula Nutt.	Urs-12-ene-3β,6β-triol-3-Gal-(1→2)-Glc	MDA-MB-231	[53]
	iguiera decurrens	Monodesmoside oleanolic acid saponins	P388, COLON	[54]
S	olidago virgaurea L.	O-glycosylation pattern at carbon atom 3 and 28		
	eteropappus altaicus	Acylglycosidic carbohydrate sequence 1-fucose-2		
	Willd.) Novopokr.	<1-xylose-3 <1-rhamnose of these bisdesmoside		[55]
Н	. biennis (Ldb.) Echinocystic acid glycosides the acylglycosidic ca	arbohydrate P-815	[55]
T	amamsch.	sequence 1-arabinose-2<1-rhamnose-4<1-gluco		
Н	elianthus annuus L.			
	Platycodon 3-O-Gl	c-2 β ,12 α ,16 α ,23,24-pentahydroxyoleanane- 28(13)-lac	ctone ECA- 109	9
Campanulaceae		c- $(1\rightarrow 3)$ -Glc- 2β , 12α , 16α , 23α - tetrahydroxyoleanane-2		[56]
	Platyco	odin D	U937, THP-1, K562	[57-59
Caprifoliaceae	Lonicera macranthoides	Macranthoside B	Many	[60]
	Dianthus versicolor	dianversicosides A-G	Many	[61]
	Vaccaria 3-0-[G	al-(1→2)-Glc] quillaic acid	LNcap,	
Caryophyllaceae	segetalis 28-0-0	$lc-(1\rightarrow 3)$ -Xyl- $(1\rightarrow 4)$ -Rha- $(1\rightarrow 2)$ -[Fuc- $(1\rightarrow 4)$]-Fuc	P-38,	[30]
curyophymaccac	VacA-5	49 caroside E, vaccaroside G, vaccaroside B, segetoside	H and segetoside I	
	Gypsophila	Gypoldoside A	Many	[62]
	oldhamiana	Jenisseensosides A, B, C, D	Human colon tumor cells	[63]
	Chenopodium	3-[(O-Glc-(1→3)-Ara)oxy]-23- oxo-olean-12-en- 28-oi	c acid Glc, HeLa	[64]
Chenopodiaceae		B-[(O-Glc-(1→3)-Ara)oxy]-27-oxo-olean-12-en-28-oic	acid Glc,	[01]
	quinoa	3-O-Ara serjanic acid 28-O-Glc ester, 3-O-Glc serjanic a	cid 28-O-Glc ester Caco-2	[64]
Combretaceae	Terminalia tropophylla	H. Perrier Terminaliaside A	A2780	[65]
	Gymnocladus chinensis	Baillon GC-1	Hepatoma cells	ICC1
		Gypenoside (Gyp) XLIX	THP-1	[66]
		(23S)-3β,20ξ,21ξ-trihydroxy-19-oxo-21,23-		
	Gynostemma	poxydammar-24-ene 3-0-[Rha(1)][Xyl(1 \rightarrow 3)]-Ara	666 7004	
	pentaphyllum	(23S)-21ξ-O-ethyl-3β,20ξ,21-trihydroxy-19-oxo-21,2	23-epoxyd SGC-7901,	[67]
	ć	mmar-24-ene 3-0-[Rha(1→2)][Xyl(1→3)]-Ara	BEL-7402	
Cucurbitaceae		$3-O-\{[Rha(1\rightarrow 2)][Xyl(1\rightarrow 3)]-6-O-acetyl\ Glc$		
		Bisdesmoside	K-562, BEL-7402	[60]
		Tubeimoside V	U87MG	[68]
	Bolbostemma paniculat	um Tubeimoside-1	HeLa	[69]
		Tubelmoside-1	CNE-2Z	[70]
	Cucurbita foetidissimo	Foetidissimoside B	Human colon cancer	[71]
	Momordica dioica	α-spinasterol-3-0-Glc	L1210	[72]
	Assenhus navia I	Aesculiosides Ia-Ie, leukemia, NSCL, col	on, CNS, melanoma,	[73]
Hippocastanaceae	Aesculus pavia L.		t ovarian, renal, prostate	
	Aesculus hippocasta	num β-escin	HT-29	[74]
Lardizabalaceae	Craniotome furcat	a Craniosaponin A	Many	[75]
	Acacia tenuifolia	Saponins	Many	[76]
	Trigonella foenum graecu		HT-29	[77]
Leguminosea	Trigonella foenumgraecu		HL-60	[78]
ocouninioseu	Gleditsia sinensis		402, BGC-823, HeLa, HL-60, MCF-7	[79]
	Gleditsia sinensis Lam.	Saponin	K562, HL-60	[79]
	Archidendron ellipticum	Elliptoside A	LOX	[80]
Molluginaceae	Glinus lotoides L.	lotoidoside D and lotoidoside E	HeLa	[81]
onagmaceae	Junus totolues L.	lotoidosides A, C, lotoidosides G	J774.A1, HEK-293, WEHI-164	[82]
	62 3223 B) 1979 N 1979	ardisiacrispin A, B		
	Ardisia pusilla A. DC.	13,28-epoxyoleanane type	U251MG	[83]
	800200000000000000000000000000000000000	olean-12-ene		
	Ardisia crenata	ardisiacrispin (A+B)	Bel-7402	[84]
Myrsinaceae		Cyclamiretin A 3 β -O-Rha-(1 \rightarrow 3)-[Xyl-(1→2)]	
ny isinaceae		-Glc- $(1\rightarrow 4)$ -[Glc- $(1\rightarrow 2)$]-Ara		
	Ardisia gigantifolia Stapf.	Cyclamiretin A 3 β -O-Rha-(1 \rightarrow 3)-[Glc-(1	1→3)-Xyl- Many	[85]
		$(1\rightarrow 2)$]-Glc- $(1\rightarrow 4)$ -Glc- $(1\rightarrow 2)$]-Ara		
		ardisiacrispin A		
		•	A375, B16	[86]
Pittosporaceae	R. parvifolius	Total saponins	7575, 510	[00]

(continued on next page)

Table 2 (continued)

Family	Species	Saponin ^a		Kinds of cancer ^b	Ref.
	Muraltia heisteria			Human colon cancer	[88]
Polygalaceae		trimethoxycinnamoyl derivative	es		
	Securidaca inappend			J-774.1	[89]
	Lysimachia thyrsif			HTB-140, HSFs	[90]
	Lysimachia	3-O-Glc oxyuronic acid-(1→2)-Xyl-cyclamiretin A		A-2780 cells	[91]
	davurica	3-O-Glc-(1→2)-Ara-cyclamiretin A			
Primulaceae		$3-0-[Xyl-(1\to 2)-Glc-(1\to 4)-[Glc-(1\to 2)]-Ara]$		Human hepatoma	[92]
	Androsace	-3β-hydroxy- 13β,28-epoxy-16-oxo-oleanan-30-al		cells	
	umbellata	$3-O-Xyl-(1\rightarrow 2)-Glc-(1\rightarrow 4)-Ara-$			
		3β-hydroxy-13β,28-epoxy-16-oxo-oleanan-30-al			
	Anemone flaccida Fr	: Flaccidoside II > anhuienoside E	> hederasaponin	HeLa cells	[93]
	Schmidt	B > glycoside St-I4a > glycoside S	St-J		
D	Pulsatilla koreana N	akai Pulsatilla saponin D		LLC in BDF1 mice	[94]
Ranunculaceae	Clematis chinensis	Monodesmosidic saponins		Many	[95]
	Clematis parvilobo	a Clematoside S, alpha-hederin	HCT-8, Bel-7402, BG	C-823, and A-2780	[96]
	Nigella glandulifer	ra Freyn et Sint. Kalopanaxsaponins A and I		HepG2, R-HepG2	[97]
	Nigella sativa	α-hederin	LL/2 in BDF1 n	nice, P388	[98]
	rigena sativa	u-nederm	A549, HEp-2, I	HT-29, MIA PaCa-2	[99]
	Pulsatilla olea	anolic acid 3-0-Rha-(1→2)-[Glc-(1→4)]-Ara A549,SKO	OV-3,SK-MEL-2,HCT15		
	koreana	hederagenin 3-O-Rha- $(1\rightarrow 2)$ -[Glc- $(1\rightarrow 4)$]-Ara		BDF1 mice bearing	[100]
	hed	lragenin 3-O-Glc-(1→4)-O-Glc- (1→3)-O-Rha-(1→2)-A	Ara	LLC	
Rubiaceae	Cimicifuga specie	Tritarnana alucacidas		MDA-MB-453r	2007
	Cimicijuga specie:	Actein			2009
		Asiaticoside A, 25-anhydrocimigenol-β-0-Xyl, asia	iticoside	B, HepG2,	2006
	Actaea asiatica	25-0-ethylcimigenol-3-0-Xyl, 25-0-acetylcimigeno	ol-3-β-O-Xyl	MCF-7	
		Trisaccharide chain attached at C3 of the aglyco	ne and	OVCAR3	[101]
		two angeloyl groups acylated at C21 and C22			
	Xanthoceras	3-0-[(3-0-Ara-2-0-Gal)-Glc]-21,		Six human tumour	[102]
	sorbifolia Bunge x	22-di-O-angeloyl-R1-barrigenol		cell lines	
	sorbijona bange x	xanifolia-Y0, -Y2, -Y3, and -Y7		OVCAR3	[103]
Sapindaceae		16-O-α-21-O-(4-angeloyl)-Rha barringtogenol C	С,	A375-S2, HeLa	[101]
баришисеце		28-O-Glc 16-deoxybarringtogenol C		A373-32, HeLa	[101]
	Thevetia peruvian	a Thevefolin, cardenolide glycosides		Human gastric	[104]
	Sapindus emargin	atus Four triterpenoid saponins		adenocarcinoma cells	
	Sapindus	Sapinmusaponins A, C, D, E	Hepa59T/V	GH, NCI, HeLa, Med	[105]
	mukorossi	Sapinmusaponin K, L, M, N		many	[106]
	Dodonaea viscosa	Dodoneasides A and B		A2780	[107]
Scrophulariaceae	Buddleja officina	lis Mimengosides C-G		HL-60	[108]
	Tribulus			MDR-cancer xenograft	[400]
Zygophyllaceae	terrestris Me	thylprototribestin, the mixture of the 3 acetylated isor	mers	transplanted mice	[109]
	Sa	ponins		786-0	[110]
				BEL-7402	[111]
				Bcap37	[112]

Saponins derived from *Platycodon grandiflorum* may suppress tumor invasion and migration by inhibiting MMP-2 and MMP-9 activation [57]. Platycodon D (Fig. 6) as one of the platycodigenins, is a potentially interesting candidate for use in cancer chemotherapy. Its exposure induced apoptosis through caspase-3 dependent PARP, lamin A cleavage and ROS induced through Egr-1 activation [128]. The primary antileukemia activity is induction of endoreduplication and mitotic arrest, as a consequence of suppressing spindle MT dynamics and promoting apoptosis in human leukemia cells [59]. Furthermore, it has direct cytotoxic effect on human leukemia cells and suppresses telomerase activity through transcriptional and post-translational suppression of hTERT [58].

Soybean saponin (Fig. 7) inhibits tumor cell metastasis by suppressing MMP-2 and MMP-9 productions, and stimulating TIMP-2 secretion [129]. At physiologically relevant doses, it can suppress HCT-15 colon cancer cell proliferation through S-phase cell cycle delay, and induce macroautophagy, the hallmark of Type II PCD. B-group soyasaponins may be another colon cancer suppressive component of soy that warrants further examination as a potential chemopreventive

phytochemical [130,131]. It significantly increased activity of raf-1 by a maximal 200%, suggesting that this enzyme in part modulates the enhanced ERK1/2 activity [132].

Pulsatilla koreana saponins (Fig. 8) were examined for their in vitro cytotoxic activity against the human solid cancer cell lines, A-549, SK-OV-3, SK-MEL-2, and HCT15, using the SRB assay method, and their in vivo antitumor activity using BDF1 mice bearing Lewis lung carcinoma (LLC) [100]. Pulsatilla saponin D as an antitumor component showed potent inhibition rate of tumor growth (IR, 82%) at the dose of 6.4 mg/kg on the BDF1 mice bearing LLC cells [133].

3.4. Spirostanes

Polyphyllin D (PD), formosanin C and dioscin belonging to the diosgenyl saponins, showed strong anti-cancer and immunostimulative activity.

With the ascertained chemical structure and the improved synthesis of polyphyllin D, both *in vitro* and *in vivo* studies were performed on its effect. Recent research showed that PD

Fig. 1. The structure of astragalosides.

Table 3 Astragalosides.

	R1	R2	R3
Astragaloside I	Xyl (2,3-diAc)	Glc	H
Astragaloside IV	Xyl	H	H
Astragaloside VII	Xyl	Glc	Glc

is a potent apoptosis inducer through mitochondrial dysfunction and ER stress [134–136].

Meanwhile, dioscin [137–141] is a preclinical drug showing potent antiproliferative activities against most cell lines from leukemia and solid tumors. Proteomic analysis revealed that it induced apoptosis via the mitochondrial and some other pathway (Fig. 9 and Table 4) [142].

Formosanin C, mainly a constituent in *Rhizoma Paris* saponins either, had some effect on the immune responses. Intraperitoneal treatment with 1–2.5 mg/kg of formosanin C would retard the growth of subcutaneously transplanted MH134 mouse hepatoma. The mechanism of its antitumor effect might be associated with its modification of the immune system [143]. It can also enhance the antitumor effect of 5-fluorouracil. Activation of caspase-2 and the

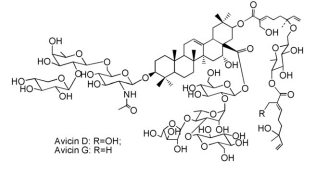


Fig. 3. Structure of avicins.

dysfunction of mitochondria maybe also contributed to its antitumor effect in human colorectal cancer HT-29 cells [144].

3.5. Furostanes

Most of furostanes only showed some anti-cancer activity. Protoneodioscin, protodioscin, protoneogracillin, and protogracillin, along with their corresponding artifacts: methyl protoneodioscin, methyl protodioscin, methyl protoneogracillin, and methyl protogracillin showed cytotoxic activities against K562 cancer cell as antineoplastic agents [18] (Fig. 10).

Methyl protogracillin was cytotoxic against all the tested cell lines from leukemia and solid tumors in the NCI's human cancer panel; it showed particular selectivity against one colon cancer line (KM12), one central nervous system (CNS) cancer line (U251), two melanoma lines (MALME-3M and M14), two renal cancer lines (786-0 and UO-31) and one breast cancer line (MDA-MB-231) [17].

4. Structure-function relationship of saponins with the antitumor properties

Differences in saponin structure which include the type, position, and number of sugar moieties attached by a

Fig. 2. Structures of ginsenosides and OSW-1.

Fig. 4. Structure of tubeimosides.

glycosidic bond at different positions of the rings can characteristically influence biological responses, especially for the antitumor activity. We could draw the following structure–activity relationships in the succeeding sections.

4.1. Influence of the aglycone on the antitumor activities of saponins

Comparing different kinds of saponins, it shows that small changes such as different positions or the number of the hydroxyl groups, R/S configuration on the aglycone led to slight changes in activity, and more sizable changes diminished the activity.

4.1.1. The site of the hydroxyl group

Changes on the agycone could change the antitumor activity of saponins. C-16 hydroxyl group of tubeimoside II plays an important role in enhancing the biological activity of tubeimoside II and in decreasing its toxicity [145] (Fig. 4). C-

Fig. 5. Structure of saikosaponins.

Fig. 6. Structure of platycodigenin.

 17α -hydroxyl group to the aglycone of the active saponins slightly reduced their cytotoxicities, such as pennogenyl saponins and diosgenyl saponins [146] (Fig. 9). C-27 of the aglycone of the furostanol saponins, which also bore an additional monosaccharide at C-27 (compared to the spirostan saponins mentioned above), showed less antitumor effect (Fig. 9).

4.1.2. The number of the hydroxyl group in aglycone

Ginsenosides with a dammarane structure have two main classes: panaxadiols (PPD) and panaxatriols (PPT). The activities of PPD compounds are greater than those of the PPT compounds. And the aglycones are more effective than the ginsenosides Rh1 (PPT type) and Rh2 (PPD type), which possess sugar moieties at C-6 and C-3, respectively. All the ginsenosides have similar chemical structures, but their effects on B16 melanoma cells were remarkably different (Fig. 2) [147].

4.1.3. Else

Based on structure–activity relationship, C-25 R/S configuration was critical for leukemia selectivity between methyl protoneogracillin and methyl protogracillin. Meanwhile, Fring was critical to selectivity between furostanol (methyl

Fig. 7. Structure of soyasaponin.

Fig. 8. Structure of Pulsatilla koreana saponin.

protoneogracillin and methyl protogracillin) and spirostanol (gracillin) saponins. Methyl protoneogracillin has been selected as a potential anti-cancer candidate for hollow fiber assay to nude mice, which is slightly better than methyl protogracillin, but gracillin would not be pursued due to the lack of selectivity against human cancer diseases (Fig. 9) [148].

4.2. Influence of the sugar side chain on the antitumor activities of saponins

In the comparison with the saponins bearing saccharide groups, different characteristics (sugar linkage, the number, lipophilicity, or different kinds) of sugar side chain play important roles in their antitumor effect.

4.2.1. The sugar linkage

With the same aglycone and length of sugar chain, the sugar linkage determines the antitumor potency. This point is clearly demonstrated by the four disaccharide congeners. $1\rightarrow 3$ linkage had much lower activity than $1\rightarrow 2$ and $1\rightarrow 4$ linkages, respectively [149].

4.2.2. The lipophilicity of the sugar

Some saponins showed no activity with the exceptions of those possessing some acyl groups at the glycosyl moiety. Meanwhile, two deoxypyranoses, including D-fucose and L-rhamnose, were also cytotoxic. These data led us to assume that the presence of a certain degree of lipophilicity in the sugar moiety is essential for exhibiting the cytotoxic activity [146].

Diosgenyl and penogenyl saponins

Fig. 9. Structures of diosgenyl and pennogenyl saponins.

Table 4 Diosgenyl saponins (R' H).

R	Name(R'=H)
-H -3-0-Glc -3-0-Rha $(1\rightarrow 2)$ -Glc -3-0-Rha $(1\rightarrow 4)$ -[Rha $(1\rightarrow 2)$]-Glc -3-0-Rha $(1\rightarrow 4)$ -[Rha $(1\rightarrow 2)$]-Glc -3-0-Rha $(1\rightarrow 2)$ -[Glc $(1\rightarrow 3)$]-Glc -3-0-Rha $(1\rightarrow 4)$ -Rha $(1\rightarrow 4)$ -[Rha $(1\rightarrow 2)$]-Glc	Diosgenin Trillin ParisV Polyphyllin D Dioscin Gracillin Formosanin C

4.2.3. The number of the sugar

The number of the sugar also influences the antitumor effect of saponins. The activity of the various ginsenosides has been demonstrated to be in the order: monosaccharide glycoside>disaccharide glycoside>trisaccharide glycoside>tetrasaccharide glycoside, indicating that increasing the number of sugar moieties reduces the potency of the compound [147]. In the contrast, diosgenyl saponins showed the contrary rule. Diosgenin β-D-glucoside showed no cytotoxic activity against HL-60 cells (IC50 20 mg/ml), and the attachment of an α -L-rhamnosyl group at C-2 of the glucosyl moiety led to the appearance of considerable activity. Further addition of an α -L-rhamnosyl, an α -Larabinofuranosyl or a β-D-glucosyl, with the exception of a β-D-galactosyl, to C-3 or C-4 of the inner glucosyl moiety either gave no influence on the activity or slightly increased the activity; the attachment of a β-D-galactosyl at C-3 of the glucosyl residue led to a decrease in the activity [150].

4.2.4. The kinds of sugar sequence

C-3 of oleanolic acid and hederagenin linked with a sugar sequence $O-\alpha$ -L-rhamnopyranosyl- $(1\rightarrow 2)-\alpha$ -L-arabinopyranoside showed a good effect, suggesting that the two elements are essential factors for the antitumor activity [100]. Meanwhile, in some disaccharide derivatives, it was used as a nontoxic carrier moiety to enhance the activity of anti-cancer drugs [151].

5. Conclusion and perspective

The identification and development of saponins have greatly contributed to medical treatment of cancer and many of these compounds are now being used in clinical practice. Almost all saponins induce apoptosis in tumor cells; they are

Fig. 10. Structure of proto-type saponins.

preferable drugs for the treatment of cancer, because eliminating tumor cells by apoptosis is helpful to lower side effects in patients by avoiding necrosis. A good understanding of the antitumor mechanisms of saponins is necessary for a directed improvement of saponin-based tumor therapies in the future. Meanwhile, special attention should be given to combinations of saponins and other anticarcinogenic drugs, since these offer very efficient treatment regimens against cancer. The most important is the saponin-mediated potentiation of tumor growth inhibition and the possibility to circumvent drug resistance. Furthermore, the elucidation of structure-activity relationships between different saponins in combination with conventional drugs is much more complicated than for saponins alone. Thus, it is not surprising that no mechanistic processes for these effects are known, however, detailed information on this basis is necessary for a directed improvement of saponin-based tumor therapies in the future.

It is hoped that the information collated here will provide the reader with information regarding the potential applications of saponins and stimulate further research into these compounds.

Acknowledgement

This work was supported by a grant 30873378 from the National Natural Science Foundation of China and a Drug Creation Project 2009ZX09103-362 from the Science and Technology in China.

References

- [1] Sparg SG, Light ME, van Staden J. Biological activities and distribution of plant saponins. J Ethnopharmacol 2004;94:219–43.
- [2] Vincken JP, Heng L, de Groot A, Gruppen H. Saponins, classification and occurrence in the plant kingdom. Phytochemistry 2007;68:275–97.
- [3] Yokosuka A, Jitsuno M, Yui S, Yamazaki M, Mimaki Y. Steroidal glycosides from Agave utahensis and their cytotoxic activity. J Nat Prod 2009;72:1399–404.
- [4] Ohtsuki T, Koyano T, Kowithayakorn T, et al. New chlorogenin hexasaccharide isolated from Agave fourcroydes with cytotoxic and cell cycle inhibitory activities. Bioorg Med Chem 2004;12:3841–5.
- [5] Xu Y, Chiu JF, He QY, Chen F. Tubeimoside-1 exerts cytotoxicity in HeLa cells through mitochondrial dysfunction and endoplasmic reticulum stress pathways. J Proteome Res 2009;8:1585–93.
- [6] Chen H, Wang G, Wang N, et al. New furostanol saponins from the bulbs of *Allium macrostemon* Bunge and their cytotoxic activity. Pharmazie 2007;62:544–8.
- [7] Mskhiladze L, Legault J, Lavoie S, et al. Cytotoxic steroidal saponins from the flowers of *Allium leucanthum*. Molecules (Basel, Switzerland) 2008;13:2925–34.
- [8] Yao N, Gu X, Li Y. Effects of three C21 steroidal saponins from Cynanchum auriculatum on cell growth and cell cycle of human lung cancer A549 cells. China Journal of Chinese Materia Medica 2009;34: 1418–21 Zhongguo Zhong yao za zhi = Zhongguo zhongyao zazhi.
- [9] Yang MF, Li YY, Gao XP, Li BG, Zhang GL. Steroidal saponins from Myriopteron extensum and their cytotoxic activity. Planta Med 2004;70:556-60.
- [10] Liu W, Huang XF, Qi Q, et al. Asparanin A induces G(2)/M cell cycle arrest and apoptosis in human hepatocellular carcinoma HepG2 cells. Biochem Biophys Res Commun 2009;381:700–5.
- [11] Zhou LB, Chen TH, Bastow KF, et al. Filiasparosides A-D, cytotoxic steroidal saponins from the roots of Asparagus filicinus. J Nat Prod 2007;70:1263-7.
- [12] Kim GS, Kim HT, Seong JD, et al. Cytotoxic steroidal saponins from the rhizomes of Asparagus oligoclonos. J Nat Prod 2005;68:766–8.
- [13] Yang CX, Huang SS, Yang XP, Jia ZJ. Nor-lignans and steroidal saponins from Asparagus gobicus. Planta Med 2004;70:446–51.

- [14] Liu MJ, Yue PY, Wang Z, Wong RN. Methyl protodioscin induces G2/M arrest and apoptosis in K562 cells with the hyperpolarization of mitochondria. Cancer Lett 2005;224:229–41.
- [15] Wang G, Chen H, Huang M, et al. Methyl protodioscin induces G2/M cell cycle arrest and apoptosis in HepG2 liver cancer cells. Cancer Lett 2006;241:102–9.
- [16] Hu K, Yao X. The cytotoxicity of methyl protodioscin against human cancer cell lines in vitro. Cancer Investig 2003;21:389–93.
- [17] Hu K, Yao X. Methyl protogracillin (NSC-698792): the spectrum of cytotoxicity against 60 human cancer cell lines in the National Cancer Institute's anticancer drug screen panel. Anticancer Drugs 2001;12:541–7.
- [18] Hu K, Dong A, Yao X, Kobayashi H, Iwasaki S. Antineoplastic agents. II. Four furostanol glycosides from rhizomes of *Dioscorea collettii* var. hypoglauca. Planta Medica 1997;63:161–5.
- [19] Hu K, Dong A, Yao X, Kobayashi H, Iwasaki S. Antineoplastic agents; I. three spirostanol glycosides from rhizomes of *Dioscorea collettii* var. hypoglauca. Planta medica 1996;62:573–5.
- [20] Wang SL, Cai B, Cui CB, et al. Apoptosis of human chronic myeloid leukemia k562 cell induced by prosapogenin B of dioscin (P.B) in vitro. Chinese Journal of Cancer 2003;22:795–800 Ai zheng = Aizheng.
- [21] Liu HW, Hu K, Zhao QC, et al. Bioactive saponins from Dioscorea futschauensis. Pharmazie 2002;57:570–2.
- [22] Dong M, Feng XZ, Wang BX, Ikejima T, Wu LJ. Steroidal saponins from Dioscorea panthaica and their cytotoxic activity. Pharmazie 2004;59:294–6.
- [23] Dong M, Feng XZ, Wu LJ, Wang BX, Ikejima T. Two new steroidal saponins from the rhizomes of *Dioscorea panthaica* and their cytotoxic activity. Planta Med 2001;67:853–7.
- [24] Hernandez JC, Leon F, Estevez F, Quintana J, Bermejo J. A homoisoflavonoid and a cytotoxic saponin from *Dracaena draco*. Chem Biodivers 2006;3:62–8.
- [25] Gonzalez AG, Hernandez JC, Leon F, et al. Steroidal saponins from the bark of *Dracaena draco* and their cytotoxic activities. J Nat Prod 2003;66:793–8.
- [26] Tran QL, Tezuka Y, Banskota AH, et al. New spirostanol steroids and steroidal saponins from roots and rhizomes of *Dracaena angustifolia* and their antiproliferative activity. J Nat Prod 2001;64:1127–32.
- [27] Man S, Gao W, Zhang Y, et al. Antitumor and antimetastatic activities of Rhizoma Paridis saponins. Steroids 2009;74:1051–6.
- [28] Cheng ZX, Liu BR, Qian XP, et al. Proteomic analysis of anti-tumor effects by *Rhizoma Paridis* total saponin treatment in HepG2 cells. [Ethnopharmacol 2008;120:129–37.
- [29] Lee JC, Su CL, Chen LL, Won SJ. Formosanin C-induced apoptosis requires activation of caspase-2 and change of mitochondrial membrane potential. Cancer Sci 2009;100:503–13.
- [30] Sy LK, Yan SC, Lok CN, Man RY, Che CM. Timosaponin A-III induces autophagy preceding mitochondria-mediated apoptosis in HeLa cancer cells. Cancer Res 2008;68:10229–37.
- [31] Ahn MJ, Kim CY, Yoon KD, et al. Steroidal saponins from the rhizomes of *Polygonatum sibiricum*. J Nat Prod 2006;69:360–4.
- [32] Pettit GR, Zhang Q, Pinilla V, et al. Antineoplastic agents. 534. Isolation and structure of sansevistatins 1 and 2 from the African Sansevieria ehrenbergii. J Nat Prod 2005;68:729–33.
- [33] Candra E, Matsunaga K, Fujiwara H, et al. Two steroidal saponins from Camassia cusickii induce L1210 cell death through the apoptotic mechanism. Can J Physiol Pharmacol 2001;79:953–8.
- [34] Yang SL, Liu XK, Wu H, Wang HB, Qing C. Steroidal saponins and cytoxicity of the wild edible vegetable—Smilacina atropurpurea. Steroids 2009:74:7–12.
- [35] Cheng G, Zhang Y, Zhang X, et al. Tubeimoside V (1), a new cyclic bisdesmoside from tubers of *Bolbostemma paniculatum*, functions by inducing apoptosis in human glioblastoma U87MG cells. Bioorg Med Chem Lett 2006;16:4575–80.
- [36] Nartowska J, Sommer E, Pastewka K, Sommer S, Skopinska-Rozewska E. Anti-angiogenic activity of convallamaroside, the steroidal saponin isolated from the rhizomes and roots of *Convallaria majalis* L. Acta Pol Pharm 2004;61:279–82.
- [37] Lu Y, Luo J, Huang X, Kong L. Four new steroidal glycosides from *Solanum torvum* and their cytotoxic activities. Steroids 2009;74:95–101.
- [38] Zhou X, He X, Wang G, et al. Steroidal saponins from *Solanum nigrum*. | Nat Prod 2006;69:1158–63.
- [39] Chiang HC, Tseng TH, Wang CJ, Chen CF, Kan WS. Experimental antitumor agents from Solanum indicum L. Anticancer Res 1991;11:1911–7.
- [40] Mimaki Y, Watanabe K, Ando Y, et al. Flavonol glycosides and steroidal saponins from the leaves of *Cestrum nocturnum* and their cytotoxicity. J Nat Prod 2001;64:17–22.
- [41] Perrone A, Plaza A, Bloise E, et al. Cytotoxic furostanol saponins and a megastigmane glucoside from tribulus parvispinus. J Nat Prod 2005;68: 1549–53.
- [42] Bedir E, Khan IA, Walker LA. Biologically active steroidal glycosides from *Tribulus terrestris*. Pharmazie 2002;57:491–3.

- [43] Gnoula C, Megalizzi V, De Neve N, et al. Balanitin-6 and -7: diosgenyl saponins isolated from *Balanites aegyptiaca* Del. display significant anti-tumor activity in vitro and in vivo. Int J Oncol 2008;32:5–15.
- [44] Fang JB, Yao Z, Chen JC, et al. Cytotoxic triterpene saponins from *Alternanthera philoxeroides*. J Asian Nat Prod Res 2009;11:261–6.
- [45] Fukumura M, Ando H, Hirai Y, et al. Achyranthoside H methyl ester, a novel oleanolic acid saponin derivative from Achyranthes fauriei roots, induces apoptosis in human breast cancer MCF-7 and MDA-MB-453 cells via a caspase activation pathway. J Nat Med 2009;63:181–8.
- [46] Yoo HH, Kwon SW, Park JH. The cytotoxic saponin from heatprocessed Achyranthes fauriei roots. Biol Pharm Bull 2006;29:1053-5.
- [47] Tundis R, Bonesi M, Deguin B, et al. Cytotoxic activity and inhibitory effect on nitric oxide production of triterpene saponins from the roots of *Physospermum verticillatum* (Waldst & Kit) (Apiaceae). Bioorg Med Chem 2009;17:4542–7.
- [48] Fujioka T, Yoshida K, Fujii H, et al. Antiproliferative constituents from Umbelliferae plants VI. New ursane-type saikosaponin analogs from the fruits of *Bupleurum rotundifolium*. Chemical & Pharmaceutical Bulletin 2003;51:365–72.
- [49] Barthomeuf C, Debiton E, Mshvildadze V, Kemertelidze E, Balansard G. In vitro activity of hederacolchisid A1 compared with other saponins from *Hedera colchica* against proliferation of human carcinoma and melanoma cells. Planta Med 2002;68:672–5.
- [50] Danloy S, Quetin-Leclercq J, Coucke P, et al. Effects of alpha-hederin, a saponin extracted from *Hedera helix*, on cells cultured in vitro. Planta Med 1994;60:45–9.
- [51] Cioffi G, Dal Piaz F, Vassallo A, et al. Antiproliferative oleanane saponins from Meryta denhamii. Journal of natural products 2008;71:1000–4.
- [52] Feng XZ, Dong M, Gao ZJ, Xu SX. Three new triterpenoid saponins from *Ixeris sonchifolia* and their cytotoxic activity. Planta Med 2003;69: 1036–40
- [53] Calabria LM, Piacente S, Kapusta I, et al. Triterpene saponins from *Silphium radula*. Phytochemistry 2008;69:961–72.
- [54] Marquina S, Maldonado N, Garduno-Ramirez ML, et al. Bioactive oleanolic acid saponins and other constituents from the roots of Viguiera decurrens. Phytochemistry 2001;56:93–7.
- [55] Bader G, Plohmann B, Hiller K, Franz G. Cytotoxicity of triterpenoid saponins. Part 1: activities against tumor cells in vitro and hemolytical index. Pharmazie 1996;51:414–7.
- [56] Son IH, Park YH, Lee SI, Yang HD, Moon HI. Neuroprotective activity of triterpenoid saponins from *Platycodi radix* against glutamate-induced toxicity in primary cultured rat cortical cells. Molecules (Basel, Switzerland) 2007;12:1147–52.
- [57] Lee KJ, Hwang SJ, Choi JH, Jeong HG. Saponins derived from the roots of Platycodon grandiflorum inhibit HT-1080 cell invasion and MMPs activities: regulation of NF-kappaB activation via ROS signal pathway. Cancer Lett 2008;268:233–43.
- [58] Dastager SG, Lee JC, Ju YJ, Park DJ, Kim CJ. Microbacterium kribbense sp. nov., isolated from soil. Int J Syst Evol Microbiol 2008;58:2536–40.
- [59] Kim MO, Moon DO, Choi YH, et al. Platycodin D induces mitotic arrest in vitro, leading to endoreduplication, inhibition of proliferation and apoptosis in leukemia cells. Int J Cancer 2008;122:2674–81.
- [60] Wang S, Li J, Huang H, et al. Anti-hepatitis B virus activities of astragaloside IV isolated from radix Astragali. Biol Pharm Bull 2009;32:132–5.
- [61] Ma L, Gu YC, Luo JG, et al. Triterpenoid saponins from *Dianthus versicolor*. J Nat Prod 2009;72:640–4.
- [62] Bai H, Zhong Y, Xie YY, et al. A major triterpenoid saponin from *Gypsophila oldhamiana*. Chem Biodivers 2007;4:955–60.
- [63] Gaidi G, Correia M, Chauffert B, et al. Saponins-mediated potentiation of cisplatin accumulation and cytotoxicity in human colon cancer cells. Planta Med 2002;68:70–2.
- [64] Kuljanabhagavad T, Thongphasuk P, Chamulitrat W, Wink M. Triterpene saponins from *Chenopodium quinoa* Willd. Phytochemistry 2008;69:1919–26.
- [65] Cao S, Brodie PJ, Callmander M, et al. Saponins and a lignan derivative of *Terminalia tropophylla* from the Madagascar Dry Forest. Phytochemistry 2009.
- [66] Huang TH, Tran VH, Roufogalis BD, Li Y. Gypenoside XLIX, a naturally occurring gynosaponin, PPAR-alpha dependently inhibits LPS-induced tissue factor expression and activity in human THP-1 monocytic cells. Toxicol Appl Pharmacol 2007;218:30–6.
- [67] Yin F, Zhang YN, Yang ZY, Hu LH. Nine new dammarane saponins from *Gynostemma pentaphyllum*. Chem Biodivers 2006;3:771–82.
- [68] Tang HF, Zhang SY, Yi YH, et al. Isolation and structural elucidation of a bioactive saponin from tubers of *Bolbostemma paniculatum*. China Journal of Chinese Materia Medica 2006;31:213–7 Zhongguo Zhong yao za zhi = Zhongguo zhongyao zazhi.
- [69] Haridas V, Nishimura G, Xu ZX, et al. Avicin D: a protein reactive plant isoprenoid dephosphorylates Stat 3 by regulating both kinase and phosphatase activities. PLoS ONE 2009;4:e5578.

- [70] Weng XY, Ma RD, Yu LJ. Apoptosis of human nasopharyngeal carcinoma CNE-2Z cells induced by tubeimoside I. Chinese Journal of Cancer 2003;22:806-11 Ai zheng = Aizheng.
- [71] Gaidi G, Marouf A, Hanquet B, et al. A new major triterpene saponin from the roots of *Cucurbita foetidissima*. J Nat Prod 2000;63:122–4.
- [72] Luo L, Li Z, Zhang Y, Huang R. Triterpenes and steroidal compounds from *Momordica dioica*. Acta Pharmaceutica Sinica 1998;33:839–42 Yao xue xue bao.
- [73] Zhang Z, Li S. Cytotoxic triterpenoid saponins from the fruits of Aesculus pavia L. Phytochemistry 2007;68:2075–86.
- [74] Patlolla JM, Raju J, Swamy MV, Rao CV. Beta-escin inhibits colonic aberrant crypt foci formation in rats and regulates the cell cycle growth by inducing p21(waf1/cip1) in colon cancer cells. Mol Cancer Ther 2006:5:1459–66.
- [75] Fan CQ, Sun HF, Chen SN, et al. Triterpene saponins from *Craniotome furcata*. Nat Prod Lett 2002;16:161–6.
- [76] Seo Y, Hoch J, Abdel-Kader M, et al. Bioactive saponins from Acacia tenuifolia from the suriname rainforest. | Nat Prod 2002;65:170–4.
- [77] Raju J, Patlolla JM, Swamy MV, Rao CV. Diosgenin, a steroid saponin of Trigonella foenumgraecum (Fenugreek), inhibits azoxymethane-induced aberrant crypt foci formation in F344 rats and induces apoptosis in HT-29 human colon cancer cells. Cancer Epidemiol Biomark Prev 2004:13:1392–8.
- [78] Hibasami H, Moteki H, Ishikawa K, et al. Protodioscin isolated from fenugreek (*Trigonella foenumgraecum* L.) induces cell death and morphological change indicative of apoptosis in leukemic cell line H-60, but not in gastric cancer cell line KATO III. Int J Mol Med 2003;11:23–6.
- [79] Zhong L, Qu G, Li P, Han J, Guo D. Induction of apoptosis and G2/M cell cycle arrest by Gleditsioside E from *Gleditsia sinensis* in HL-60 cells. Planta Med 2003;69:561–3.
- [80] Beutler JA, Kashman Y, Pannell LK, et al. Isolation and characterization of novel cytotoxic saponins from Archidendron ellipticum. Bioorg Med Chem 1997;5:1509–17.
- [81] Yan MC, Liu Y, Chen H, et al. Synthesis and antitumor activity of two natural N-acetylglucosamine-bearing triterpenoid saponins: lotoidoside D and E. Bioorg Med Chem Lett 2006;16:4200–4.
- [82] Hamed AI, Piacente S, Autore G, et al. Antiproliferative hopane and oleanane glycosides from the roots of *Glinus lotoides*. Planta Med 2005;71:554–60.
- [83] Tian Y, Tang HF, Qiu F, et al. Triterpenoid saponins from Ardisia pusilla and their cytotoxic activity. Planta Med 2009;75:70–5.
- [84] Zheng ZF, Xu JF, Feng ZM, Zhang PC. Cytotoxic triterpenoid saponins from the roots of Ardisia crenata. J Asian Nat Prod Res 2008;10:833–9.
- [85] Wen P, Zhang XM, Yang Z, Wang NL, Yao XS. Four new triterpenoid saponins from Ardisia gigantifolia Stapf. and their cytotoxic activity. Journal of Asian natural products research 2008;10:873–80.
- [86] Zheng ZX, Zhang LJ, Huang CX, et al. Antitumour effect of total saponins of Rubus parvifolius on malignant melanoma. China Journal of Chinese Materia Medica 2007;32:2055–8 Zhongguo Zhong yao za zhi = Zhongguo zhongyao zazhi.
- [87] Kim YJ, Wang P, Navarro-Villalobos M, et al. Synthetic studies of complex immunostimulants from *Quillaja saponaria*: synthesis of the potent clinical immunoadjuvant QS-21Aapi. J Am Chem Soc 2006;128:11906–15.
- [88] Elbandy M, Miyamoto T, Chauffert B, Delaude C, Lacaille-Dubois MA. Novel acylated triterpene glycosides from *Muraltia heisteria*. J Nat Prod 2002;65:193–7.
- [89] Yui S, Ubukata K, Hodono K, et al. Macrophage-oriented cytotoxic activity of novel triterpene saponins extracted from roots of Securidaca inappendiculata. Int Immunopharmacol 2001;1:1989–2000.
- [90] Galanty A, Michalik M, Sedek L, Podolak I. The influence of LTS-4, a saponoside from *Lysimachia thyrsiflora* L., on human skin fibroblasts and human melanoma cells. Cell Mol Biol Lett 2008;13:585–98.
- [91] Liang B, Tian JK, Xu LZ, Yang SL. Triterpenoid saponins from *Lysimachia davurica*. Chem Pharm Bull 2006;54:1380–3.
- [92] Wang Y, Zhang D, Ye W, et al. Triterpenoid saponins from Androsace umbellata and their anti-proliferative activities in human hepatoma cells. Planta Med 2008;74:1280-4.
- [93] Lu J, Xu B, Gao S, et al. Structure elucidation of two triterpenoid saponins from rhizome of *Anemone raddeana* Regel. Fitoterapia 2009;80:345–8.
- [94] Bang SC, Seo HH, Yun HY, Jung SH. Facile synthesis of trisaccharide moiety corresponding to antitumor activity in triterpenoid saponins isolated from *Pullsatilla* roots. Chem Pharm Bull 2007;55:1734–9.
- [95] Mimaki Y, Yokosuka A, Hamanaka M, et al. Triterpene saponins from the roots of *Clematis chinensis*. J Nat Prod 2004;67:1511–6.
- [96] Yan LH, Xu LZ, Lin J, Yang SL, Feng YL. Triterpenoid saponins from the stems of Clematis parviloba. J Asian Nat Prod Res 2009;11:332–8.
- [97] Tian Z, Liu YM, Chen SB, et al. Cytotoxicity of two triterpenoids from Nigella glandulifera. Molecules (Basel, Switzerland) 2006;11:693–9.
- [98] Ali BH, Blunden G. Pharmacological and toxicological properties of Nigella sativa. Phytother Res 2003;17:299–305.

- [99] Rooney S, Ryan MF. Effects of alpha-hederin and thymoquinone, constituents of Nigella sativa, on human cancer cell lines. Anticancer Res 2005;25:2199–204.
- [100] Bang SC, Lee JH, Song GY, et al. Antitumor activity of *Pulsatilla koreana* saponins and their structure–activity relationship. Chem Pharm Bull 2005;53:1451–4.
- [101] Chan PK. Acylation with diangeloyl groups at C21–22 positions in triterpenoid saponins is essential for cytotoxicity towards tumor cells. Biochem Pharmacol 2007;73:341–50.
- [102] Li ZL, Yang BZ, Li X, et al. Triterpenoids from the husks of Xanthoceras sorbifolia Bunge. J Asian Nat Prod Res 2006;8:361–6.
- [103] Chan PK, Zhao M, Che CT, Mak E. Cytotoxic acylated triterpene saponins from the husks of Xanthoceras sorbifolia. J Nat Prod 2008;71: 1247–50.
- [104] Miyagawa T, Ohtsuki T, Koyano T, Kowithayakorn T, Ishibashi M. Cardenolide glycosides of *Thevetia peruviana* and triterpenoid saponins of *Sapindus emarginatus* as TRAIL resistance-overcoming compounds. J Nat Prod 2009;72:1507–11.
- [105] Kuo YH, Huang HC, Yang Kuo LM, et al. New dammarane-type saponins from the galls of Sapindus mukorossi. J Agric Food Chem 2005;53:4722-7.
- [106] Huang HC, Wu MD, Tsai WJ, et al. Triterpenoid saponins from the fruits and galls of Sapindus mukorossi. Phytochemistry 2008;69:1609–16.
- [107] Cao S, Brodie P, Callmander M, et al. Antiproliferative triterpenoid saponins of *Dodonaea viscosa* from the Madagascar dry forest. J Nat Prod 2009;72:1705–7.
- [108] Guo H, Koike K, Li W, et al. Saponins from the flower buds of Buddleja officinalis. J Nat Prod 2004;67:10–3.
- [109] Ivanova A, Serly J, Dinchev D, et al. Screening of some saponins and phenolic components of *Tribulus terrestris* and *Smilax excelsa* as MDR modulators. In vivo (Athens, Greece) 2009;23:545–50.
- [110] Yang HJ, Qu WJ, Sun B. Experimental study of saponins from *Tribulus terrestris* on renal carcinoma cell line. China Journal of Chinese Materia Medica 2005;30:1271–4 Zhongguo Zhong yao za zhi = Zhongguo zhongyao zazhi.
- [111] Sun B, Qu WJ, Zhang XL, et al. Investigation on inhibitory and apoptosis-inducing effects of saponins from *Tribulus terrestris* on hepatoma cell line BEL-7402. China Journal of Chinese Materia Medica 2004;29: 681–4 Zhongguo Zhong yao za zhi = Zhongguo zhongyao zazhi.
- [112] Sun B, Qu W, Bai Z. The inhibitory effect of saponins from *Tribulus terrestris* on Bcap-37 breast cancer cell line in vitro. Journal of Chinese medicinal materials 2003;26:104–6 Zhong yao cai = Zhongyaocai.
- [113] Auyeung KK, Law PC, Ko JK. Astragalus saponins induce apoptosis via an ERK-independent NF-kappaB signaling pathway in the human hepatocellular HepG2 cell line. Int J Mol Med 2009;23:189–96.
- [114] Tin MM, Cho CH, Chan K, James AE, Ko JK. Astragalus saponins induce growth inhibition and apoptosis in human colon cancer cells and tumor xenograft. Carcinogenesis 2007;28:1347–55.
- [115] Auyeung KK, Cho CH, Ko JK. A novel anticancer effect of Astragalus saponins: transcriptional activation of NSAID-activated gene. Int J Cancer 2009;125:1082–91.
- [116] Zhou Y, Garcia-Prieto C, Carney DA, et al. OSW-1: a natural compound with potent anticancer activity and a novel mechanism of action. J Natl Cancer Inst 2005;97:1781–5.
- [117] Jayatilake GS, Freeberg DR, Liu Z, et al. Isolation and structures of avicins D and G: in vitro tumor-inhibitory saponins derived from Acacia victoriae. J Nat Prod 2003;66:779–83.
- [118] Haridas V, Higuchi M, Jayatilake GS, et al. Avicins: triterpenoid saponins from Acacia victoriae (Bentham) induce apoptosis by mitochondrial perturbation. Proc Natl Acad Sci USA 2001;98:5821–6.
- [119] Xu ZX, Liang J, Haridas V, et al. A plant triterpenoid, avicin D, induces autophagy by activation of AMP-activated protein kinase. Cell Death Differ 2007;14:1948–57.
- [120] Lemeshko VV, Haridas V, Quijano Perez JC, Gutterman JU. Avicins, natural anticancer saponins, permeabilize mitochondrial membranes. Arch Biochem Biophys 2006;454:114–22.
- [121] Haridas V, Li X, Mizumachi T, et al. Avicins, a novel plant-derived metabolite lowers energy metabolism in tumor cells by targeting the outer mitochondrial membrane. Mitochondrion 2007;7:234–40.
- [122] Yu LJ, Ma RD, Wang YQ, et al. Potent anti-tumorigenic effect of tubeimoside 1 isolated from the bulb of *Bolbostemma paniculatum* (Maxim) Franquet. Int J Cancer 1992;50:635–8.
- [123] Zhang C, Li B, Gaikwad AS, et al. Avicin D selectively induces apoptosis and downregulates p-STAT-3, bcl-2, and survivin in cutaneous T-cell lymphoma cells. J Investig Dermatol 2008;128:2728–35.
- [124] Wen-Sheng W. ERK signaling pathway is involved in p15INK4b/ p16INK4a expression and HepG2 growth inhibition triggered by TPA and Saikosaponin a. Oncogene 2003;22:955–63.
- [125] Chen JC, Chang NW, Chung JG, Chen KC. Saikosaponin-A induces apoptotic mechanism in human breast MDA-MB-231 and MCF-7 cancer cells. Am J Chin Med 2003;31:363–77.

- [126] Hsu YL, Kuo PL, Chiang LC, Lin CC. Involvement of p53, nuclear factor kappaB and Fas/Fas ligand in induction of apoptosis and cell cycle arrest by saikosaponin d in human hepatoma cell lines. Cancer Lett 2004:213:213–21.
- [127] Hsu YL, Kuo PL, Lin CC. The proliferative inhibition and apoptotic mechanism of saikosaponin D in human non-small cell lung cancer A549 cells. Life Sci 2004;75:1231–42.
- [128] Shin DY, Kim GY, Li W, et al. Implication of intracellular ROS formation, caspase-3 activation and Egr-1 induction in platycodon D-induced apoptosis of U937 human leukemia cells. Biomedicine & Pharmacotherapy 2009;63:86–94 Biomedecine & pharmacotherapie.
- [129] Kang JH, Han IH, Sung MK, et al. Soybean saponin inhibits tumor cell metastasis by modulating expressions of MMP-2, MMP-9 and TIMP-2. Cancer Lett 2008:261:84–92.
- [130] Ellington AA, Berhow M, Singletary KW. Induction of macroautophagy in human colon cancer cells by soybean B-group triterpenoid saponins. Carcinogenesis 2005;26:159–67.
- [131] Zhang W, Popovich DG. Effect of soyasapogenol A and soyasapogenol B concentrated extracts on HEP-G2 cell proliferation and apoptosis. J Agric Food Chem 2008;56:2603–8.
- [132] Ellington AA, Berhow MA, Singletary KW. Inhibition of Akt signaling and enhanced ERK1/2 activity are involved in induction of macroautophagy by triterpenoid B-group soyasaponins in colon cancer cells. Carcinogenesis 2006;27:298–306.
- [133] Gao XD, Ye WC, Yu AC, et al. Pulsatilloside A and anemoside A3 protect PC12 cells from apoptosis induced by sodium cyanide and glucose deprivation. Planta Med 2003;69:171–4.
- [134] Cheung JY, Ong RC, Suen YK, et al. Polyphyllin D is a potent apoptosis inducer in drug-resistant HepG2 cells. Cancer Lett 2005;217:203–11.
- [135] Siu FM, Ma DL, Cheung YW, et al. Proteomic and transcriptomic study on the action of a cytotoxic saponin (Polyphyllin D): induction of endoplasmic reticulum stress and mitochondria-mediated apoptotic pathways. Proteomics 2008;8:3105–17.
- [136] Lee MS, Yuet-Wa JC, Kong SK, et al. Effects of polyphyllin D, a steroidal saponin in *Paris polyphylla*, in growth inhibition of human breast cancer cells and in xenograft. Cancer Biol Ther 2005;4:1248–54.
- [137] Zhang Y, Li HZ, Zhang YJ, et al. Atropurosides A–G, new steroidal saponins from *Smilacina atropurpurea*. Steroids 2006;71:712–9.
- [138] Nguyen VT, Darbour N, Bayet C, et al. Selective modulation of P-glycoprotein activity by steroidal saponines from *Paris polyphylla*. Fitoterapia 2008.
- [139] Yun H, Li JC, Wen HZ, et al. Separation and identification of steroidal compounds with cytotoxic activity against human gastric cancer cell lines in vitro from the rhizomes of *Paris polyphylla* var. *chinensis*. Chemistry of Natural Compounds 2007;43:672–7.
- [140] Wang T, Liu Z, Li J, et al. Determination of protodioscin in rat plasma by liquid chromatography-tandem mass spectrometry. J Chromatogr 2007;848:363–8.
- [141] Cai J, Liu M, Wang Z, Ju Y. Apoptosis induced by dioscin in Hela cells. Biol Pharm Bull 2002;25:193–6.
- [142] Wang Y, Cheung YH, Yang Z, et al. Proteomic approach to study the cytotoxicity of dioscin (saponin). Proteomics 2006;6:2422–32.
- [143] Wu RT, Chiang HC, Fu WC, et al. Formosanin-C, an immunomodulator with antitumor activity. Int J Immunopharmacol 1990;12:777–86.
- [144] Nam DH, Jeon HM, Kim S, et al. Activation of notch signaling in a xenograft model of brain metastasis. Clin Cancer Res 2008;14: 4059-66.
- [145] Yu TX, Ma RD, Yu LJ. Structure–activity relationship of tubeimosides in anti-inflammatory, antitumor, and antitumor–promoting effects. Acta Pharmacol Sin 2001;22:463–8.
- [146] Mimaki Y, Yokosuka A, Kuroda M, Sashida Y. Cytotoxic activities and structure-cytotoxic relationships of steroidal saponins. Biol Pharm Bull 2001;24:1286–9.
- [147] Wang W, Zhao Y, Rayburn ER, et al. In vitro anti-cancer activity and structure–activity relationships of natural products isolated from fruits of *Panax ginseng*. Cancer Chemother Pharmacol 2007;59:589–601.
- [148] Hu K, Yao X. The cytotoxicity of methyl protoneogracillin (NSC-698793) and gracillin (NSC-698787), two steroidal saponins from the rhizomes of *Dioscorea collettii* var. hypoglauca, against human cancer cells in vitro. Phytother Res 2003;17:620–6.
- [149] Chwalek M, Lalun N, Bobichon H, Ple K, Voutquenne-Nazabadioko L. Structure-activity relationships of some hederagenin diglycosides: haemolysis, cytotoxicity and apoptosis induction. Biochim Biophys Acta 2006;1760:1418–27.
- [150] Wang Y, Zhang Y, Zhu Z, et al. Exploration of the correlation between the structure, hemolytic activity, and cytotoxicity of steroid saponins. Bioorg Med Chem 2007;15:2528–32.
- [151] Bang SC, Seo HH, Shin HR, et al. A convenient preparation of a disaccharide motif and its role in the cytotoxicity of the triterpenoid saponin, alpha-hederin. Arch Pharm Res 2008;31:555–61.