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Anticancer Properties of Phytochemicals Present in Medicinal Plants of North America

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Additional information is available at the end of the chapter

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1. Introduction

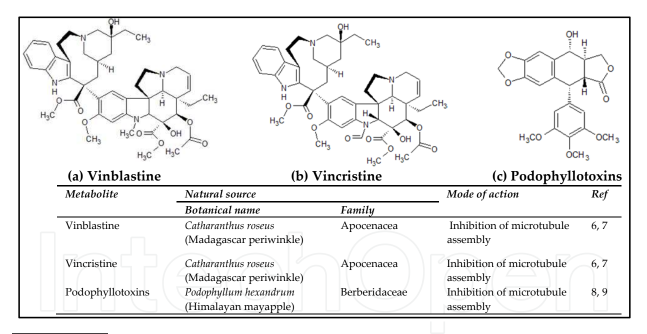
Cancer is one of the most severe health problems in both developing and developed countries, worldwide. Among the most common (lung, stomach, colorectal, liver, breast) types of cancers, lung cancer has continued to be the most common cancer diagnosed in men and breast cancer is the most common cancer diagnosed in women. An estimated 12.7 million people were diagnosed with cancer across the world in 2008, and 7.6 million people died from the cancer during the same year [1]. Lung cancer, breast cancer, colorectal cancer and stomach cancer accounted for two-fifths of the total cases of cancers diagnosed worldwide [1]. More than 70% of all cancer deaths occurred in low- and middle-income countries. Deaths due to cancer are projected to continuously increase and it has been estimated that there will be 11.5 million deaths in the year 2030 [1] and 27 million new cancer cases and 17.5 million cancer deaths are projected to occur in the world by 2050 [2]. According to Canadian cancer statistics, issued by the Canadian Cancer Society, it is estimated that 186,400 new cases of cancer (excluding 81,300 non-melanoma skin cancers) and 75,700 deaths from cancer will occur in Canada in 2012 [1]. The lowest number of incidences and mortality rate is recorded in British Columbia. Both incidence and mortality rates are higher in Atlantic Canada and Quebec [3].

More than 30% of cancers are caused by modifiable behavioral and environmental risk factors, including tobacco and alcohol use, dietary factors, insufficient regular consumption of fruit and vegetable, overweight and obesity, physical inactivity, chronic infections from *Helicobacter pylori*, hepatitis B virus (HBV), hepatitis C virus (HCV) and some types of human papilloma virus (HPV), environmental and occupational risks including exposure to ionizing and non-ionizing radiation [4].



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Conventional treatment of cancer includes interventions such as psychosocial support, surgery, radiotherapy and chemotherapy [4]. Currently, the most commonly use cancer chemotherapy includes mainly alkylating agents, antimetabolites, antitumor antibiotics, platinum analogs and natural anticancer agents. However, due to the increasing rate of mortality associated with cancer and adverse or toxic side effects of cancer chemotherapy and radiation therapy, discovery of new anticancer agents derived from nature, especially plants, is currently under investigation. Screening of medicinal plants as a source of anticancer agents was started in the 1950s, with the discovery and development of vinca alkaloids, vinblastine and vincristine and the isolation of the cytotoxic podophyllotoxins [5] (Figure 01). The cool temperate climate of North America supports the growth of an enormous number of plant species which are important sources of unique phytochemicals having anticancer properties (Table 01). In this chapter, selected medicinal plants grown in the cool climate of North America (mainly Canada and USA) are discussed. The major bioactive phytochemicals and their mechanisms of action are also reviewed.



(a) Vinblastine – [dimethyl $(2\beta,3\beta,4\beta,5\alpha,12\beta,19\alpha)$ - 15-[(55,9S)- 5-ethyl- 5-hydroxy- 9-(methoxycarbonyl)- 1,4,5,6,7,8,9,10-octahydro- 2H- 3,7-methanoazacycloundecino[5,4-b]indol- 9-yl] - 3-hydroxy- 16-methoxy- 1-methyl- 6,7-didehydroaspidospermidine- 3,4-dicarboxylate] (b) Vincristin – [(3aR,3a1R,4R,5S,5aR,10bR)-methyl 4-acetoxy-3a-ethyl-9-((5S,7S,9S)-5-ethyl-5-hydroxy-9-(methoxycarbonyl)-2,4,5,6,7,8,9,10-octahydro-1H-3,7-

methano[1]azacycloundecino[5,4-b]indol-9-yl)-6-formyl-5-hydroxy-8-methoxy-3a,3a1,4,5,5a,6,11,12-octahydro-1Hindolizino[8,1-cd]carbazole-5-carboxylate] (c) Podophyllotoxin – [(10*R*,11*R*,15*R*,16*R*)-16-hydroxy-10-(3,4,5trimethoxyphenyl)-4,6,13-trioxatetracyclohexadeca-1,3(7),8-trien-12-one]

Figure 1. Some selected currently used phytochemical-based anticancer agents

Plant	Family	Parts used	Major bioactive compounds		Ref	
Achyranthes	Amaranthaceae	Leaf	Triterpenoid saponins	regions USA	14	
aspera						
(Devil's Horsewhip)						
Annona glabra	Annonaceae	Leaf and	Acetogenins	USA	15	
(Pond apple)		fruit		0071	10	
Aralia nudicaulis	Araliacea	Whole	Steroids, sarsasapogenin, smilagenin,	Mainly	16	
(Wild sarsaparilla)		plant	sitosterol, stigmasterol, pollinastrenol,	Canada		
(Wha saisapanna)			glycosides, saponins, sarsasaponin			
			parillin, smilasaponin, smilacin,			
			sarsaparilloside, and sitosterol			
			glucoside			
Aster brachyactis	Asteraceae	Aerial	Not known	North	17	
(Rayless aster)	Asteraceae		NOT KHOWH	America	17	
(Rayless aster) Carduus nutans	Asteraceae	parts Aerial	Linalool derivatives, aliphatic acids,	North	18 10	
(Nodding	ASIEIALEAE		diacids, aromatic acids, and phenols	America	18, 19	
-		parts	diacids, aromatic acids, and phenois	America		
plumeless thistle)	Liliaceae	Whole		North	20 21	
Erythronium	Lillaceae		Alpha-methylenebutyrolactone		20, 21	
americanum		plant		America		
(Adder's tongue)	A .			NL	20.24.40	
Eupatorium	Asteraceae	Whole	Sesquiterpene lactone, pyrrolizidine	North	20, 21,19	
cannabinum		plant	alkaloid, and flavonoid	America	22	
(Bonesets)					22	
Foeniculum	Apiaceae	Seed	α-pinene, anisic aldehyde, cineole,	North	23	
vulgare			fecchone, limonene, and myrcene	America		
(Wild pepper						
fennel)						
Hydrastis	Ranunculaceae	Whole	Isoquinoline alkaloids (hydrastine,	Canada, USA	20, 21	
canadensis		plant	berberine, berberastine, candaline),			
(Orange root)			resin and lactone			
Hypericum	Clusiaceae	Flower	Hypericin and hyperforin	USA, Canada	24	
perforatum				(British		
(St. John's wort)		$\langle \nabla \rangle$		Columbia)		
Lactuca sativa	Asteraceae	Leaf	Sesquiterpene lactone	USA, Canada	25	
(Garden lettuce)						
Lantana camara	Verbenaceae	Whole	Alkaloids (camerine, isocamerine,	USA	26, 27, 28	
(Wild sage)		plant	micranine, lantanine, lantadene),			
			phenols, flavonoids, tannins, saponins,			
			and phytosterols			
Larrea tridentate	Zygophyllaceae	Whole	Resins and lignans	Southwester	18, 29, 30	
(Creosote bush)		plant		n USA		
Linum	Linaceae	Seed	Enterodiol, enterolactone, lignans, and	Canada, USA	31, 32	
usitatissimum			omega-3 fatty acids			
(Common Flax)						

Plant	Family	Parts used	Major bioactive compounds	Growing regions	Ref	
Olea europrae	Oleaceae	Leaf and	Oleuropein, hydroxytyrosol,	USA	33, 34,	
(Olive)		oil	hydroxytyrosol acetate, luteolin-7-O-		35, 36, 3	
			glucoside, luteolin-4'-O-glucoside and			
			luteolin, oleic acid and polyphenol			
Panax	Araliaceae	Root,	Ginsenosides and saponins	Eastern North	20, 21	
quinquefolius		Leaf		America		
(North American						
Ginseng)						
Plantago	Plantaginaceae	Aerial	Phenolics and flavonoids	Canada, USA	38	
lanceolata		parts				
(Ribwort plantain)						
Podophyllum	Berberidaceae	Rhizome	Podophyllotoxins	Eastern North	39	
peltatum				America		
(Mayapple)						
Polygonatum	Polygonaceae	Whole	Saponin and flavonoid and vitamin A	USA	20, 21, 4	
multiflorum		plant				
(Tuber fleece						
flower)						
Pyrus malus	Rosaceae	Bark and	Quercetin, catechin, flavonoid,	North	21	
(Apple)		fruit	coumaric and gallic acids, phloridzin	America		
			and procyanidin			
Rhodiola rosea	Crassulaceae	Rhizome	Monoterpene alcohols and their	Eastern	41, 42	
(Golden root)			glycosides, cyanogenic glycosides, aryl	Canada		
			glycosides, phenylethanoids,			
			phenylpropanoids and their			
			glycosides, flavonoids, flavonlignans,			
			proanthocyanidins and gallic acid			
			derivatives			
Saponaria vaccaria	Caryophyllaceae	Seed	Flavonoids, cyclopeptides, and	Western	43	
(Cowherb)			bisdesmosidic saponins	Canada		
Silybum marianum	Asteraceae	Dried	Silymarin-polyphenoic flavolignans	Canada, USA	44, 45	
(Milk thistle)		fruit,	(silybin, isosilybin, silychristin,			
		seed	silydianin and taxifoline)			
Sonchus arvensis	Asteraceae	Whole	Alkanes, n-alkenes, n-aldehydes and n-	Canada	46, 47	
(Perennial sow-		plant	alcohols, shikimate metabolites,			
thistle)			carotenoid-derived compounds,			
			terpenoids, steroids, and phenols			
Tanacetum vulgare	Asteraceae	Aerial	Monoterpenes, sesqueterpenes, and	Canada, USA	48	
(Tansy)		parts	oxygenated sesqueterpenes			
Taraxacum	Asteraceae	' Root and	Sesquiterpene lactones, triterpenoids,	North	49	
officinale		leaf	sterols, tannins, alkaloids, inulin,	America		
Officinale		icui	sterois, tarinins, antaroras, maini,			

Plant	Family	Parts	Major bioactive compounds	Growing	Ref	
Flant	Failing	used	Major bloactive compounds	regions	Nei	
Taxus brevifolia	Тахасеае	Bark	Taxol (diterpene)	Pacific	50	
(Pacific yew tree)				Northwest		
Thuja occidentalis	Cupressaceae	Whole	Flavonoid, tannin, and volatile oil	Northeastern	13, 51	
(White cedar)		plant		USA,		
				Eastern		
				Canada		
Xanthium	Asteraceae	Fruit	Sesquiterpene lactones (Xanthatin	Canada	17	
strumarium			and Xanthinosin)			
(Cocklebur)						

Table 1. Medicinal plants with potential anticancer properties grown in North America

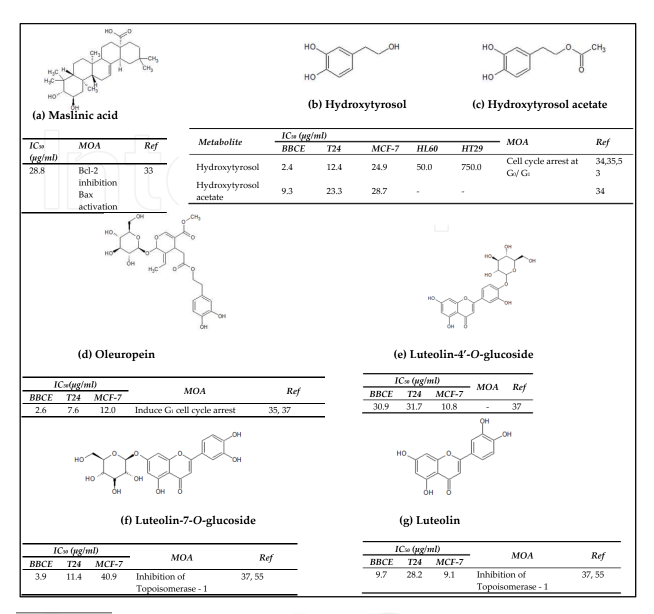
2. Pathophysiology of cancer

Cancer is a population of abnormal cells which divide without control, with the ability to invade other tissues. Cancer and some of the other chronic diseases share common pathogenesis mechanisms, such as DNA damage, oxidative stress, and chronic inflammation [10]. It is understood that both environmental factors and chemical carcinogens play a key role in the initiation and progression of cancer. Among the major environmental factors are asbestos, polluted air near industrial emission sources, exposure to secondary tobacco smoke, indoor air pollution such as radon, drinking water containing arsenic, chlorination by-products, and other pollutants [11]. Chemicals with carcinogenic activity can be classified as DNA reactive (e.g.: nitrogen mustards, chlorambucil, epoxides, aliphatic halides, aromatic amines), epigenetic (e.g.: chlordane, pentachlorophenol, hormones, cyclosporin, purine analogs), dichlorodiphenyltrichloroethane, phenobarbital, minerals (e.g.: asbestos), metals (e.g.: arsenic, beryllium, cadmium) and unclassified carcinogens (e.g.: acrylamide, acrylonitrile, dioxane) [12]. DNA-reactive carcinogens act in the target cells of tissue(s) of their carcinogenicity to form DNA adducts that are the basis for neoplastic transformation [12]. Epigenetic carcinogens lack chemical reactivity and hence, do not form DNA adducts. These carcinogens are produced in the target cells of tissue(s) of their carcinogenicity. Effects of epigenetic carcinogens indirectly lead to neoplastic transformation or enhance the development of tumors from cryptogenically transformed cells [12].

Carcinogenesis is a multi-step process consisting of tumor initiation, promotion and progression [13]. Cancer initiation can be blocked by activating protective mechanisms, either in the extracellular environment or intracellular environment by modifying trans-membrane transport, modulating metabolism, blocking reactive oxygen and nitrogen species, maintaining DNA structure, modulating DNA metabolism and repair, and controlling gene expression [10]. Tumor promotion is the second stage of carcinogenesis and is followed by tumor progression. Both stages can be suppressed by inhibiting genotoxic effects, favoring antioxidant and anti-inflammatory activity, inhibiting proteases and cell proliferation, inducing cell differentiation, modulating apoptosis and signal transduction pathways and protecting intercellular communications [10]. In addition, tumor progression can also be inhibited by affecting the hormonal status and the immune system in various ways and by inhibiting tumor angiogenesis [10].

3. In vitro anticancer activity of phytochemicals and extracts of medicinal plants

Cultured cancer-derived cell lines with comparison to normal healthy cell lines are commonly used to assess the anticancer properties of isolated phytochemicals and extracts of medicinal plants (Table 2). The anticancer properties of ethanolic extract of leaves, pulp and seeds of, Annona glabra (L.), commonly known as pond apple, were shown, using human drug-sensitive leukemia (CEM) and its multidrug-resistant-derived (CEM/VLB) cell lines [52]. The most potent anticancer activity was shown in the seed extract of A. glabra [52]. Both dried rhizome hexane extract and dried fruit hexane extract, partitioned from total methanol extract, of Aralia nudicaulis (L.) caused death of cancer cell lines such as human colon cancer cell (WiDr), human leukemia cell (Molt) and human cervix cancer cell (HeLa) at a lower concentration, than that of required for the death of normal cells [53]. Eupatoriopicrin, a sesquiterpene lactone isolated from Eupatorium cannabinum (L.) (Bonesets), indicated anticancer properties on FIO 26 (fibrosarcoma) cells with an IC₅₀=1.5 μ g/ml [22]. Methanolic extracts of Hypericum perforatum (L.) (St. Johns wort) possessed strong antiproliferative activity in the human prostate cell line (PC-3) and the major constituents, hyperforin and hypericin, synergistically contributed to the reduction of the PC-3 cells proliferation [24]. Maslinic acid, (Figure 2(a)) a triterpene from Olea europaea (L.) (Olive), has shown to be significantly inhibitory in cell proliferation of the human colorectal adenocarcinoma cell line (HT29) in a dose dependent manner [33]. The major components in the extract were identified to be oleuropein, hydroxytyrosol, hydroxytyrosol acetate, luteolin-7-O-glucoside, luteolin-4'-O-glucoside and luteolin [34] (Figure 2). All these phytochemicals inhibited the proliferation of cancer and endothelial cells with IC_{50} , at the low micromolar range [37]. Methanolic leaf extract of *Plantago lanceolata* (L.) (Ribwort plantain) inhibited the growth of three different cell lines; human renal adenocarcinoma (TK-10), the human breast adenocarcinoma (MCF-7) and the human melanoma (UACC-62) cell lines and the MCF-7 was totally inhibited [54]. Further, the ethanolic extract of P. lanceolata (L.), produced by maceration with ethanol : water, showed significant antiproliferative activity on cervix epithelioid carcinoma (HeLa), breast adenocarcinoma (MCF-7), colon adenocarcinoma (HT-29) and human fetal lung carcinoma (MRC-5) [38]. Several chemical constituents (Figure 3) from Silybum marianum (Milkthistle) have been isolated and their cytotoxic and anticancer potential has been investigated, in vitro, using both cancer and normal healthy cell lines. Silymarin, isolated from seeds of S. marianum, is a mixture of series of flavolignans, major constituents being: silybin A and B, (also known as silibinin), isosilybin A and B, silychristin, and silvdianin [55, 56].



IC₅₀: Concentration which inhibited 50% of cell proliferation; **MOA**: Mode of Action; **BBCE**: Bovine Brain Capillary Endothelial cells; **T24**: Human Urinary Bladder Carcinoma cells; **MCF-7**: Human Breast Adenocarcinoma cells; **HL60**: Human Promyelocytic Leukaemia cells; **HT29**: Colon Adenocarcinoma cells

(a) Maslinic acid - [(2a, 3b)-2,3-dihydroxyolean-12-en-28-oic acid]: (b) Oleuropein – [(45,5E,6S)-4-[2-[2-(3,4-dihydroxy-phenyl)ethoxy]-2-oxoethyl]- 5-ethylidene-6-[[(25,3R,4S,5S,6R)-3,4,5-trihydroxy-6-(hydroxymethyl)- 2-tetrahydropyranyl]oxy]-4H-pyran-3-carboxylic acid, methyl ester] : (c) Hydroxytyrosol – [4-(2-Hydroxyethyl)-1,2-benzenediol]: (d) Hydroxytyrosol acetate – [2-(3,4-dihydroxy)Phenyl ethyl acetate;4-[2-(Acetyloxy)ethyl]-1,2-benzenediol]: (e) Luteolin-7-O-glucoside - [2-(3,4-dihydroxyphenyl)-5-hydroxy-4-oxo-4H-chromen-7-yl beta-D-glucopyranoside; 4H-1-benzopyran-4-one, 2-(3,4-dihydroxyphenyl)-7-(beta-D-glucopyranosyloxy)-5-hydroxy-; 2-(3,4-Dihydroxy-phenyl)-5-hydroxy-7-((25,3R,4S,5S,6R)-3,4,5-trihydroxy-6-hydroxymethyl-tetrahydro-pyran-2-yloxy)-chromen-4-one] (f) Luteolin-4'-Oglucoside – [3',5,7-Trihydroxy-4'-(β-D-glucopyranosyloxy)flavone;2-(4-β-D-Glucopyranosyloxy-3-hydroxyphenyl)-5,7dihydroxy-4H-1-benzopyran-4-one;2-[3-Hydroxy-4-(β-D-glucopyranosyloxy)phenyl]-5,7-dihydroxy-4H-1benzopyran-4-one] (g) Luteolin - [2-(3,4-Dihydroxyphenyl)-5,7-dihydroxy-4-chromenone]

Figure 2. Major bio-active compounds present in Olea europaea (a,b,c,d,e,f and g) and Plantago lanceolata (f and g)

Silybin possessed a dose-dependent growth inhibitory effect on parental ovarian cancer cells (OVCA 433), drug-resistant ovarian cancer cells (A2780 WT) and doxorubicin (DOX)-resistant breast cancer cells (MCF-7) [55]. Both L and D diastereoisomers of silvbin inhibited A2780 WT cell growth at low IC₅₀ reported with L-diastereoisomer [55]. Furthermore, silybin potentiated the effect of Cisplatin (CDDP, a platinum analog; cis-diamminedichloroplatinum [II]) in inhibiting A2780 WT and CDDP-resistant cell growth. Cisplatin is an inorganic metal complex which acts as an alkylating agent [57]. Similar results recorded with doxorubicin (DOX) on MCF-7 DOX-resistant cells when silvbin associated with doxorubicin. Doxorubicin ((7S,9S)-7-[(2R,4S,5S,6S)-4-amino-5-hydroxy-6-methyloxan-2-yl]oxy-6,9,11-trihydroxy-9-(2-hydroxyace tyl)-4-methoxy-8,10-dihydro-7H-tetracene-5,12-dione) is an anthracycline antibiotic isolated from Streptomyces peucetius var caesius [57]. The effect of silvbin-CDDP and silvbin-DOX combinations resulted in a synergistic action, as assessed by the Berembaum isobole method [55]. Silymarin demonstrated to have marked inhibition of cell proliferation with almost 50% inhibition in a time dependent manner on the human breast cancer cell line (MDA-MB 468), at 25 µg/mI concentration, after five days of treatment. Its potential anticancer activity was dose dependent and showed a complete inhibition of cancer cells at 50 and 75 µg/mI concentrations at the beginning of Day 2 of exposure [56]. Induction of apoptotic cell death of human prostate cancer (DU145) treated with silibinin is shown to be due to activation of caspase 9 and caspase 3 enzymes [58].

4. Evidence from animal studies for anticancer activity of North American medicinal plants

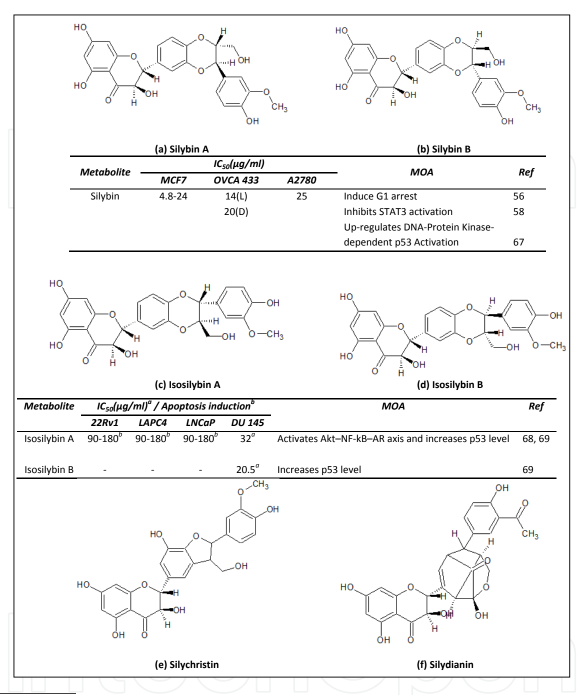
Anticancer and antiproliferative potential of some North American medicinal plants has also been studied in animal studies (Table 3). In vivo antitumor activities of Achyranthes aspera (L.) (Devil's Horsewhip) on athymic mice, with are subcutaneous xenograft, harboring human pancreatic tumor were demonstrated, using the leaf extract. The leaf extract significantly reduced both tumor weight and volume in mice treated with leaf extract intraperitonealy [14]. Intravenous administration of 40 mg/kg body weight eupatoriopicrin, a sesquiterpene lactone present in E. cannabinum, significantly delayed the growth of tumor in Lewis lung tumour-bearing syngeneic C57B1 female mice [22]. A 70% inhibition of tumor growth in PC-3 cells, orthopedically implanted into the dorsal prostatic lobe in athymic nude mice, was observed, upon their receiving 15 mg/kg intraperitoneal H. perforatum methanolic extract [24]. Lantadene A is a pentacyclic triterpenoid, isolated from the weed, Lantana camara (L) [59]. Feeding of female Swiss albino mice (LACCA) with a dose of 50 mg/kg body weight of Lantadene A twice a week for 20 weeks, showed potential chemopreventive activity. This chemopreventive activity could be linked to the expression of transcriptional factors and a significant decrease in the mRNA expression of AP-1 and c-fos), NF-kB (p-65) and p53 was observed in Lantadene A treated mice skin tumors [59]. Silibinin decreased tumor multiplicity by 71% (P < 0.01) in wild type mice, but did not show any such considerable effect in iNOS-/- mice upon oral feeding of 742 mg/kg body weight silibinin for 5 days per week for 18 weeks [60]. Lesser effects of silibinin in iNOS^{-/-} mice suggested that most of its chemopreventive and angiopreventive effects were through its inhibition of iNOS expression in lung tumors [60]. Treatment of a purified diet, containing 0.5% to 1.0% silibinin on a transgenic adenocarcinoma of are mouse prostate (TRAMP) model, decreased the weight of the tumor in both the prostate and seminal vesicle, when compared with control mice [61].

Treatment of silibinin significantly decreased tumor angiogenesis and proliferation and also there was increased apoptosis in prostate tumor tissue samples in the TRAMP model [61]. The protective effect of silibinin was also demonstrated in mouse skin with tumors caused by acute and chronic UVB-exposure-caused mitogenic and survival signaling and associated biological responses [62]. Mice were treated with silibinin, either topically (9 mg in 200 ml acetone/mouse) or orally (1% of diet), and both administrations strongly inhibited UVB-induced skin tumorigenesis in a long-term study [62]. Thymine dimers are formed in DNA, immediately after UVB irradiation, and are considered as an early and important biomarker for UVB induced DNA damage [62]. A noticeable, 71% reduction (P < 0.001) of thymine positive cells was obtained in the mice treated with 1% (w/w) silibinin before the UVB exposure, compared with the UVB alone group [62]. Oral feeding of 200 mg/kg of silibinin for 5 days per week, for 33 days, significantly inhibited human non-small-cell lung cancer cells (NSCLC A549) tumor xenograft growth in nude mice, in a time-dependent manner [63]. This accounted for 58% (P = 0.003) reduction in tumor weight per mouse and intraperitoneal administration of 4 mg/kg doxorubicin, once a week for four weeks, showed 61% (P = 0.005) reduction in tumor weight. However, interestingly, in silibinin-doxorubicin combination, 76% (P = 0.002, versus control) decrease in tumor weight per mouse was observed, that which was significantly different from either treatment alone, showing enhanced efficacy [63].

5. Mode of action of selected phytochemicals of North American medicinal plants

Apoptosis (programmed cell death) is the principal mechanism through which unwanted or damaged cells are safely eliminated from the body. This programmed cell death is mediated via either an extrinsic apoptotic pathway or an intrinsic apoptotic pathway [65]. These two apoptosis signaling pathways differ in the origin of their apoptosis signal, but converge upon a common pathway [66].

The extrinsic pathway is initiated by the stimulation of the cell surface 'death receptor' due to the binding of death ligand and the intrinsic pathway is also known as the mitochondrial pathway in which an intracellular apoptotic signal initiates the process [68]. Various natural extracts, obtained from medicinal plants grown in North America, have been found to induce apoptosis pathways at different levels (Figure 4 and Table 04). Leaf extract of *Achyranthes aspera* activated caspase-3 and induced caspase-3 mRNA in tumor cells. It also decreased Akt-1 transcription, as well its phosphorylation. Suppression of pAkt-1 and a corresponding activation of caspase 3 by the leaf extract, induced apoptosis of tumor cells [14]. It was also found that maslinic acid, isolated from *O. europaea*, inhibited considerably the expression of Bcl-2 (B-cell lymphoma 2), whilst increasing that of Bax. Maslinic acid stimulated the release of mitochondrial cytochrome-c and activated caspase-9 and caspase-3 [33]. These results showed the activation of the mitochondrial apoptotic pathway, in response to the treatment



IC₅₀: Concentration which inhibited 50% of cell proliferation; MOA: Mode of Action; MCF-7: Doxorubicin-resistant breast cancer cells; OVCA 433: Parental ovarian cancer cells; A2780: Drug-resistant ovarian cancer cells; 22Rv1, LAPC4, LNCaP, DU 145: Human prostate cancer cells

(a) Silybin A – [(2R,3R)-3,5,7-trihydroxy-2-[(2R,3R)-3-(4-hydroxy-3-methoxyphenyl)-2-(hydroxymethyl)-2,3-dihydroben-zo[b] [1,4]dioxin-6-yl]chroman-4-one]: (b) Silybin B – <math>[(2R,3R)-3,5,7-trihydroxy-2-[(2S,3S)-3-(4-hydroxy-3-methoxy-phenyl)-2-(hydroxymethyl)-2,3-dihydrobenzo[b] [1,4]dioxin-6-yl]chroman-4-one]: (c) Isosilybin A –<math>[(2R,3R)-3,5,7-Trihydroxy-2-[(2R,3R)-2-(4-hydroxy-3-methoxyphenyl)-3-(hydroxymethyl)-2,3-dihydrobenzo[1,4]dioxin-6-yl]-4-

chromanone]: (d) (c) Isosilybin B -[(2R,3R)-3,5,7-Trihydroxy-2-[(2R,3R)-2-(4-hydroxy-3-methoxyphenyl)-3-(hydroxymethyl)-2,3-dihydrobenzo[1,4]dioxin-6-yl]-4-chromanone]: (e) Silychristin – [(2R,3R)-3,5,7-trihydroxy-2-[(2R, 3S)-7-hydroxy-2-(4-hydroxy-3-methoxyphenyl)-3-(hydroxymethyl)-2,3-dihydro-1-benzofuran-5-yl]-2,3-dihydrochromen-4-one]:

Figure 3. Major bio-active flavonolignans present in Silybum marianum

of HT29 colon-cancer cells with maslinic acid. The major flavonoid present in *P. lanceolata*, luteolin-7-*O*-β-glucoside, as well as aglycon luteolin, acted as potent poisons for DNA topoisomerase I on cancer cell lines [54]. Silibinin (major bioactive component from *S. maria-num*) markedly activated the DNA-PK-p53 pathway for apoptosis, in response to UVB-induced DNA damage [69]. DNA-PK pull-down assay showed that silibinin pre-treatment strongly increased binding of DNA protein kinase with p53 [69].

Plant	Extraction solvent and concentration	Type of cancer cell line	IC ₅₀ or growth reduction	Key findings	Ref
Annona glabra (Pond apple)	Ethanolic extract of lyophilized plant material in powder form	Human drug- sensitive leukemia (CEM) and its multidrug- resistant-derived (CEM/VLB) cell lines	Leaf-1.00 (CEM/ VLB) Pulp-0.65 (CEM/VLB), Seed-0.10 (CEM/VLB) and Leaf-0.30 (CEM), Pulp-0.35 (CEM), Seed-0.07 (CEM) µg/ml	IC ₅₀ values were significantly lower than Adriamycin (Doxorubicin) (CEM=0.13 μg/ml and CEM/VLB=13.4 μg/ml) indicates its potential for cancer drug discovery programs	52
Aralia nudicaulis (Wild sarsaparilla)	Methanol extracts of rhizome, stem, leaf and fruit were further partitioned with hexane, ethyl acetate, butanol and water	WiDr (colon), Molt (leukemia), HeLa (cervix)	Hexane rhizome extract 30.1 (WiDr), 7.0 (Molt), 33.33 (HeLa) µg/ml	The concentrations of Rhizome hexane and Fruit hexane required for normal cell death was significantly higher than those required for the cancer cells	53
Eupatorium cannabinum (Bonesets)	Eupatoriopicrin concentrations of 0.1 - 50 µg/ml in 96% ethanol	FIO 26 (Fibrosarcoma)	1.5 μg/ml	Possess significant anticancer activity	22
Foeniculum vulgare (Wild pepper fennel)	Not specified	Breast (MCF-7), liver (HepG2)	-	Remarkable anticancer potential	23
Hypericum perforatum (St. John's wort)	Methanolic extract	Prostate (PC-3)	0.42 mg/ml	Extract components synergistically contribute to the	24

Plant	Extraction solvent and concentration	Type of cancer cell line	IC ₅₀ or growth reduction	Key findings	Ref
Linum Usitatissimum	Ethanol extract	Breast (MCF-7, MDA-MB-231)	Growth reduction of	reduction of the PC-3 cells proliferation Significantly reduced cell growth and induced apoptotic	31
(Common flax)			15.8% in MCF-7 and 11.4% in MDA-MB-231	cell death	
Olea europrae (Olive)	maslinic acid 0– 100 μg/mL	Colon (HT29)	28.8 µg/ml	Cell proliferation inhibition in a dose-dependent manner and causes apoptotic death	33
	Aqueous extract and methanol artificial mixture	Breast (MCF-7), Human urinary bladder (T-24), Bovine brain (BBCE)	72 (MCF-7), 100 (T-24), and 62 (BBCE) for aq. 565 (MCF-7), 135 (T-24), and 42 (BBCE) for methanol μg/ml	Antiproliferative activity of the extracts should mainly be attributed to its identified phytochemicals	34
Plantago lanceolata (Ribwort plantain)	Methanolic extract	Renal (TK-10), breast (MCF-7), melanoma (UACC-62)	"/>250 (ТК-10), 47.2 (МСF-7), 50.6 (UACC-62) µg/ml	Growth of MCF-7 was totally inhibited	54
	Extracted by maceration with ethanol/water during 72 hr at room temperature	cervix epitheloid (HeLa),breast (MCF-7), colon (HT-29), fetal lung (MRC-5)	172.3 (HeLa), 142.8 (MCF-7), 405.5 (HT-29), 551.7 (MRC-5) μg/ml	Showed significant antiproliferative activity	38
Rhodiola rosea (Golden root)	Not specified	Urinary bladder (RT4, UMUC-3, T24, 5637, J82)	264 (RT4),100 (UMUC-3), 71 (T24), 151 (5637), 165 (J82) μg/ml	Selectively inhibit the growth of cancer cell lines with minimal effect on nonmalignant cells	41
Saponaria vaccaria (Cowherb)	70% Methanol extract	colon (WiDr), breast (MDA- MB-231), lung (NCI-417), prostate (PC-3),	3.8-9.4 (WiDr), 11.4-19.6 (MDA- MB-231), 12.6-18.4	Dose-dependent growth inhibitory and selective apoptosis-inducing activity. Strong in a breast and a prostate cancer cell lines	43

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Plant	Extraction solvent and concentration	Type of cancer cell line	IC₅₀ or growth reduction	Key findings	Ref
		nontumorigenic fibroblast BJ (CRL-2522)	(NCI-417) mg/ml		
Silybum marianum (Milkthistle)	silybin, a flavonoid	Doxorubucin resistant breast (MCF-7), Parental ovarian (OVCA 433), Drug- resistant ovarian (A2780)	4.8-24 μM (MCF-7), 14 & 20 μM - L & D diastereoisomer s respectively (A2780) 25 μg/ml	Dose-dependent growth inhibitory effect on all three cell lines	55
	Silymarin at a dosages of 10- 75 μg/ml in ethanol	Breast (MDA-MB 468)	-	Inhibits the cell proliferation in a dose- and time dependent manner	56
	Silibinin in DMSO	Prostate (DU145)	-	Strongly inhibited activation of Stat3 and causes caspase activation and apoptotic death	58
	Isosilybin A and B	Prostate (LNCaP, 22Rv1) Prostate (DU 145)	lso A:32 μM (DU 145) lso B:20 μM (DU 145)	Anti-prostate cancer activity mediated via cell cycle arrest and apoptosis induction	69
Taraxacum officinale (Dandelions)	Water (lyophilized or reconstituted)	Acute T-cell leukemia (Jurkat clone E6-1), dominant- negative FADD Jurkat cells (clone I 2.1)		Effectively and selectively induced apoptosis in human leukemia cell lines in a dose and time dependent manner	49

Table 2. Anti-cancer properties of phytochemicals and extracts of medicinal plants revealed from *in vitro* studies using cancer cell lines

Plant	Preparation	Animal model	Dosage	Key findings	Ref.
		used			
Achyranthes	5% suspension in	Athymic nude	50, 100 and 200	The tumor weight and	14
aspera	hexane followed by	mice	mg/kg extract in	volume was significantly	
(Devil's	extraction in		1 ml PBS	reduced in the mice treated	
Horsewhip)	acetone overnight		administered IP	for 36 days with 50 mg/kg.	

Plant	Preparation	Animal model	Dosage	Key findings	Ref
		used			
	and residue was			In one treated mouse tumor	
	dissolved in			completely disappeared	
	methanol				
Eupatorium	Eupatoriopicrin, a	Syngeneic C57B1	i.v. injection of 20	Significantly stronger	22
cannabinum	sesquiterpene	female mice	or 40 mg/kg	growth delay of both lung	
(Bonesets)	lactone			tumours and fibrosarcoma	
Hypericum	Methanolic extract	Human prostatic	ip with a dose of	Inhibited tumor growth by	24
perforatum		carcinoma cell	15 mg/kg	70% with no observed side	
Orange		line	dissolved in 1%	effects	
root)		orthotopically	DMSO		
		implanted			
		athymic male			
		nude mice			
Lantana	Lantadene A,	Female Swiss	50 mg/kg body	Activity could be linked to	59
camara	pentacyclic	albino mice	weight twice a	the expression of	
(Wild sage)	triterpenoid	(LACCA)	week for 20	transcriptional factors	
	·	. ,	weeks	·	
Silybum	Silibinin	Lung - Male	742 mg/kg body	Significantly decreases	60
marianum		B6/129-	weight for 5	urethane-induced tumor	
(Milkthistle)		Nos2tm1Lau	d/wk for 18	number and size in WT mice.	
, , , , , , , , , , , , , , , , , , ,		(iNOS ^{-/-}) and	weeks	Decreased tumor	
		B6/129PF2 WT		multiplicity in WT mice, but	
		mice		not in iNOS ^{-/-} mice	
	Silibinin	Prostate - A	Purified diet	Decreased the weight of	61
		transgenic	containing 0%	tumor + prostate + seminal	0.
		adenocarcinoma	and 1%	vesicle. Significantly	
		of mouse	(w/w) silibinin	decreased tumor	
		prostate (TRAMP)	until	angiogenesis and	
		model	until	proliferation and increased	
				apoptosis also.	
	Topically applied	Skin – mouse	9 mg in 200 ml	silibinin (both topical and	62
	silibinin in acetone	Skin mouse	acetone/mouse	oral) strongly inhibited UVB-	02
	or oral feeding of		or 1% in diet	induced skin tumorigenesis	
	silibinin		of 176 in diet	in long-term study	
	Silibinin	Chip CKU 1	10/(100/100)	0	62
		Skin - SKH-1	1% (w/w)	Strong suppression of UVB-	63
		hairless mouse	silibinin in diet for 2 weeks	induced damage by dietary	
	Cilibinin	Athymaic (DALD)		feeding of silibinin	C A
	Silibinin	Athymic (BALB/	200 mg/kg body	Significantly inhibits human	64
		c,nu/nu) male	weight, 5 d/wk	NSCLC A549 tumor	
		nude mice	for 33 days	xenograft growth in a time	
				dependent manner	

Table 3. Anti-cancer properties of medicinal plants revealed from in vivo studies using experimental animals

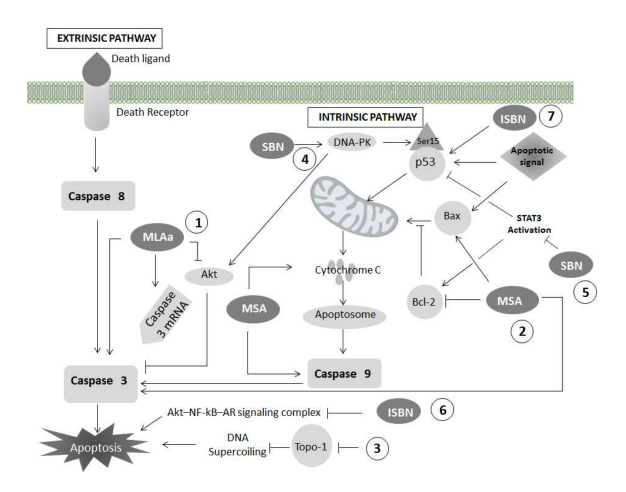


Figure 4. Schematic representation of current knowledge of mode of action of some selected anticancer phytochemicals in North America (in a hypothetical cancer cell).

Akt (Protein kinase B); Bcl-2 (Protein kinase B); Bax (Bcl-2–associated X protein); Topo-1 (Topoisomerase 1); p53 (tumor protein 53); Ser15 (Serine 15); NF-kB (nuclear factor kappa-light-chain-enhancer of activated B cells); AR (Androgen Receptor)

- **1.** Methanolic leaf extract of *Achyranthes aspera* (MLAa) induces caspase -3 mRNA and suppress expression of the kinase Akt-1. Apoptosis is induced by activation of caspase-3 and inhibiting Akt-phosphorylation.
- 2. The mechanism of Maslinic acid (MSA) (isolated from *Olea europaea*) is regulated via Bcl-2 inhibition and Bax induction, producing mitochondrial disruption, cytochrome-c release, leading finally to the activation of caspases 9 and caspase 3.
- **3.** Luteolin-7-*O*-β-glucoside and its aglycon, luteolin (major bio-active constituents of *Plantago lanceolata*) showed DNA topoisomerase I poison activities and Topoisomerase mediated DNA damage might be the possible mechanism which induce apoptosis.
- **4.** Silibinin (SBN) (extracted from *Silybum marianum*) pretreatment enhance DNA-PK (DNA Protein kinase) associated kinase activity as well as the physical interaction of p53 with DNA-PK and it preferentially activates the DNA-PK-p53 pathway for apoptosis.
- 5. SBN inhibits active Stat3 phosphorylation, and causes caspase activation and apoptosis.

- **6.** Isosilybin A (ISBN) (extracted from *Silybum marianum*) activates apoptotic machinery in human prostate cancer cells via targeting Akt–NF-kB–AR axis.
- 7. ISBN increases p53 protein levels.

Plant	Mode of action	References
Achyranthes aspera	Significantly induced caspase-3 mRNA and suppressed expression of the pro	14
(Devil's Horsewhip)	survival kinase Akt-1. Apoptosis was induced by activation of caspase-3 and	
	inhibiting Akt phosphorylation.	
Olea europaea	Activation of the mitochondrial apoptotic pathway	33
(Olive)	Significant block of G_1 to S phase transition manifested by the increase of cell	37
	number in G ₀ /G ₁ phase	
Plantago lanceolata	The topoisomerase-mediated DNA damage seems to be a candidate	54
(Ribwort plantain)	mechanism, by which some flavonoids may exert their cytotoxic potential	
Silybum marianum	Induces G1 arrest in cell cycle progression	56
(Milkthistle)	Up-regulates DNA-protein kinase-dependent p53 activation to enhance UVB-	67
	induced apoptosis	68
	Activates apoptotic machinery in human prostate cancer cells via targeting Akt-	58
	NF-kB–AR axis	69
	Inhibits active Stat3 phosphorylation, and causes caspase activation	
	Increases total p53 levels	
Podophyllum peltatum	Inhibition of microtubule assembly	70
(Mayapple)		

Table 4. Mode of action of anticancer activity of phytochemicals present in selected North American medicinal plants

6. Conclusion

Currently, natural products, especially plant secondary metabolites such as isoprenoids, phenolics and alkaloids, have been demonstrated to be the leading providers of novel anticancer agents. Thiese important groups of phytochemicals represent a vast majority of chemical groups, including alkaloids, flavonoids, flavonols, flavanols, terpenes and terpenoids, phenols, flavonolignans and steroids. Potential anticancer properties of these phytochemicals have been shown by both cell culture (*in vitro* methods) and animal (*in vivo* methods) studies. However, *in vitro* and *in vivo* findings should be strengthened by valid human clinical trial data before introducing to the medicine cabinet as natural therapeutics or drugs.

Abbreviations

CEM, Human drug-sensitive leukemia cells; CEM/VLB, Human multidrug-resistant-derived leukemia cells; Jurkat clone E6-1, Acute T-cell leukemia cells; WiDr and HT29, Human colon

cancer cells; Molt, Human leukemia cancer cell; FIO 26, Human fibrosarcoma cells; MCF-7 and MDA-MB-231, Human breast cancer cells; HepG2, Human hepatocarcinoma cells; LNCaP, 22Rv1, PC-3 and DU145, Human prostate cancer cells; RT4, UMUC-3, T24, 5637 and J82, Human urinary bladder cancer cells; BBCE, Bovine brain capillary endothelial cells, TK-10, Human renal cancer cells; UACC-62, Human melanoma cells; HeLa, Human cervical epithelioid cells; MRC-5, Fetal lung cancer cells; NCI-417, Human lung cancer cells; CRL-2522, Human nontumorigenic fibroblast cells; OVCA 433, Parental ovarian cancer cells; A2780, Drug-resistant ovarian cancer cells.

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