Abstract

Statement of Purpose: Although 75% of breast cancer patients may use integrative medicine, only 11.5% early-stage breast cancer patients believe integrative medicine has anticancer activity. The gap in users of and firm believers in integrative medicine indicates a need to increase awareness of integrative medicine’s applicability to curative cancer treatment. As diet, nutraceuticals, and traditional Chinese medicine are used by up to 82% of cancer patients who use integrative medicine, this paper focuses on nutraceuticals. As female cancer patients are most likely to use integrative medicine, nutraceuticals specific to breast, cervical, endometrial, and ovarian cancer are reviewed.

Methods: PubMed searches in September 2016 and January 2017, and accompanying hand searches were performed for English language, free full text articles published from 2012 onwards. Search terms were combinations of the key words: Homeopathy, phytochemicals, breast cancer, cervical cancer, endometrial cancer, ovarian cancer, cancer, prevention, treatment. Curative nutraceutical treatments were taken from these searches. Supplemental hand searches were performed as needed.

Findings: Integrative nutraceutical therapies are based on biologic plausibility. Individual nutraceuticals are frequently comprised of numerous phytochemical types, and have multiple mechanisms of action. Agonist-antagonist mechanisms are not exclusive to phytoestrogens, complicating which derivative or whole nutraceutical to use. Research on breast and cervical cancer inhibitors has seemingly outpaced that for endometrial and ovarian cancer. Breast cancer inhibitors include terpenes, isoflavones, organosulphurs, organoselenium compounds, and withanolides.

Conclusion and Significance: Potential nutraceutical candidates for curative cancer treatment abound. Discerning which constituents, which extraction method, and which delivery method to use for an efficacious treatment is an essential, lengthy process. If nutraceuticals such as limonene and Kahalalide F move forward to Phase III trials a nutraceutical cancer treatment pipeline may be established. Wilthaferin-A and derivatives for breast and ovarian cancer appear to be the leading gynecologic cancer nutraceutical drug candidates.

Keywords: Cancer treatment; Integrative cancer treatment; Naturopathic medicine; Nutraceuticals; Phytochemicals; Phytonutrients; Traditional chinese medicine

Introduction

One-time integrative medicine use ranges from 42% in the United States to 75% in France [1]. Upon cancer diagnosis patients are interested in conventional and integrative therapies [2]. Post cancer diagnosis, 17% of Italian, 27% of Austrians, 35.9% of Europeans overall, 44% of German gynecologic cancer, 47.2% of Canadian, and 50% of German breast cancer patients use integrative medicine therapies [2-6]. Consistent with this, integrative gynecologic cancer treatment is regarded as reasonable by 44.5%, and practiced by 24.2% of 310 Hessian region ObGyns who responded to a survey at an educational meeting [7]. Nonetheless, extremes in integrative cancer treatment are evident. In one study, 75% of breast cancer patients used integrative medicine [8]. However, as few as 11.5% of early stage breast cancer patients may perceive integrative medicine as having anticancer activity [9]. Pediatric oncology integrative medicine ranges from 12% to 42% in Europe and 25% to 82% in the United States [1].

In Dutch pediatric oncology patients, female gender is predictive of integrative cancer use [10]. Similarly, a cross-sectional study of three Pennsylvanian oncology centers found that being female (p=0.005), having breast cancer (p=0.016), and being at 12 to 36 months post diagnosis (p=0.017) most associated with integrative cancer treatment [11]. Time from diagnosis had an n shaped association with integrative cancer treatment [11]. Herbal and traditional Chinese medicine (TCM; excluding acupuncture) are received by 74.5% of patients at European integrative oncology centers [6]. Among Italian cancer patients as few as 38.3% currently and as many as 84.2% previously used nutraceuticals comprised of diet, dietary supplements, and herbs [12].

Nutraceuticals

Nutraceuticals range from active constituent phytochemicals, minerals, and vitamins, through whole functional foods [13]. Integrative nutraceutical therapies are based on biologic plausibility and have played an historical role in the development of the prescription pharmaceutical industry [14]. Willow bark derived analgesic aspirin, Aspergillus terreus derived cholesterol-lowering lovastatin, *Camellia sinensis* (green tea) polyphenols derived...
Methods

Historically, nutraceuticals have been a starting point of new manufacturing processes including plant cell suspension cultures to Taxus baccata (European yew tree), topotecan derived from epigallocatechin-3-gallate (EGCG) [17]. Clinical trial EGCG dosing is based 10 mg/kg body weight, which can be achieved from green tea [18]. Pharmaceuticals may require enormous amounts of the biologic source: 1,520 Taxus chinensis (Chinese yew) trees, an endangered species, are used to produce 1 kg of paclitaxel [19]. Whereas nutraceuticals, being food or dietary supplement products with health modifying effects do not require a prescription [20]. Thus, nutraceuticals lie in a gray zone between prescription pharmaceuticals and open market household items [15]. Nonetheless, nutraceutical agriculture and manufacturing may have to comply with good agricultural practices, good manufacturing practices, and standard operating procedures [21]. Nutraceuticals undergo in vitro and in vivo pharmacokinetic, pharmacologic, therapeutic, safety, and head-to-head comparative effectiveness studies as do pharmaceuticals [20]. Clinical pharmacology studies of nutraceuticals are essential for nutraceutical development and delineation [22]. Nutraceuticals use as prophylaxis against and treatment of pharmaceuticals’ adverse effects is the subject of a separate paper, which also addresses nutraceuticals’ inherent adverse effects. This paper reviews nutraceuticals as curative (alternative) cancer treatment, focusing on gynecologic cancer treatment.

Nutraceuticals by phytochemical classification

This study does not review vitamins and minerals per se. However, whole fruits and vegetables, and fruit and vegetable derived phytochemicals may be vitamin and mineral rich. Fruit and vegetable derived phytochemicals may be studied based on mechanism of action, pathologies treated, or biochemical classification. By mechanism of action, the Matrix MetalloProtease (MMP) inhibitors aqueous cinnamon extract, green tea extract, curcumin, fenugreek derived cinnamon, and marine compound derived chitoooligosaccharides would be grouped together. However, each of these nutraceuticals has multiple mechanisms of action. Therefore, nutraceuticals and cancer treatment capabilities are initially presented based on chemical classification. There are numerous biochemical nutraceutical categories including alkaloids, lipids, organic acids and polysaccharides, organosulphurs, phenols, phytic acids, phytosterols, and terpenes, as indicated in Figure 2. Functional foods and other nutraceuticals comprised of constituent compounds will have constituent compounds from multiple biochemical classifications. The encompassing groups of alkaloids, organic acids, and polysaccharides will be briefly mentioned. Nutraceuticals from the secondary and tertiary groups will be briefly mentioned. Then attention will turn to those nutraceuticals pertinent to breast, cervical, and ovarian cancer treatment.

Alkaloids

Alkaloids are physiologically active, vegetable-based, organic, nitrogen-containing ring compounds. Alkaloids may belong to additional biochemical phytounitnutrient groups based on further chemical structure definition and resultant activity. Alkaloids can be co-constituents of a functional food, as is the case with Camellia sinensis tea leaves, which also contain flavonoids, steroids, gallic tannins and catecholic tannins (flavanols) [23]. There are numerous alkaloid sub-classifications: Coffee derives aroma from trigonelline, a bitter alkaloid, and bitter taste from caffeine, a purine-like alkaloid [24]. Alkaloids include colchicine from Colchicum autumnale (autumn crocus or meadow saffron), scopolamine/hyoscyamine from Hyoscyamus niger (henbane or stinking nightshade), physostigmine from Physostigma venenosa (Calabar bean), reserpine from...
**Rauvolfia serpentine** (Indian snakeroot or devil pepper), and taxol from *Taxus brevifolia* (Pacific yew) [24].

![Figure 2: Nutraceuticals biochemical classification.](image)

**Organic acids and polysaccharides**

Organic acids and polysaccharides are mentioned first as these are encompassing chemical composition groupings. Constituents thereof will belong to additional biochemical phytonutrient groups based on chemical structure and resultant activity. Organic acids are carbon containing acids. Polysaccharides are comprised of bonded sugar molecules.

**Organic acids**: Organic acids are inflammatory mediators with antioxidant, chemopreventive, and hepatoprotective properties [25]. Cinnamic acid in Aloe vera, ellagic acid (a polyphenol organic acid, found in berries, green tea, guava, pecans, and walnuts), ferulic acid in oats and rice, gallic acid in tea, oxalic acid in coffee, spinach, and tea, and salicylic acid in peppermint are a few examples [25]. Ellagic acid is a reactive epoxide scavenger that inhibits DNA methylation and DNA polymerase [26].

**Polysaccharides**: Mushrooms are immune boosting chemopreventive polysaccharides [25]. Fibrous polysaccharides also bind carcinogens, lower bile acids, and modulate estrogen metabolism [26]. Polysaccharides from the TCM adaptogen ginseng up regulates chemopreventive polysaccharides [25]. Fibrous polysaccharides also bind carcinogens, lower bile acids, and modulate estrogen metabolism [26]. Polysaccharides from the TCM adaptogen ginseng up regulates chemopreventive polysaccharides [25].

**Organosulphur compounds**

Sulphur’s pungent odor is a hallmark of the organosulphur compounds comprised of indoles, thiouisothionates, and isothiocyanates. Organosulphur compounds, which include indole-3-carbinol (I3C) and diindolylmethane (DIM), are commonly derived from cruciferous vegetables. Cephalosporins which were developed from Acremonium cellulare and penicillin from Penicillium fungi are also organosulphurs. Rauwolfia serpentine (Indian snakeroot or devil pepper), and taxol from Taxus brevifolia (Pacific yew) [24].

**Indoles and Thiocyanates**: Indoles and thiouisothionates result from the ingestion and digestion of some cruciferous vegetables [25,28]. 

Indoles induce and activate cytochrome P450 for phase I detoxification [25]. In A/J mice, I3C inhibits lung adenoma [28]. In BALB/c mice DIM inhibits lung metastasis [28]. While I3C is rapidly hydrolyzed in the stomach, hydro and lipophbic DIM requires combination with vitamin E and phosphatidylcholine for bioavailability [28].

**Isothiocyanates**: Allium compounds found in garlic induce glutathione s-transferase (GST) and microsomal monooxygenases [26]. Garlic, onion, and turnips are isothiocyanate sources. Cabbage, garden cress, and Indian cress derived benzyl isothiocyanate (BITC) and watercress derived phenethyl isothiocyanate (PEITC) may be less studied than the phenols, terpenes, and thiosulfonates that are also found in allium containing vegetables [29,30]. Nonetheless, BITC activates AP-1, caspase-3, MAPK, and NF-kβ while inhibiting other protein kinases [31]. BITC induces apoptosis, G2/M phase cell-cycle arrest, glutathione depletion, and reactive oxygen species (ROS) generation [31]. PEITC inhibits phase I enzymes but activates phase II enzymes [31].

**Lipids**

There are numerous lipid subgroups including fats, waxes, glycolipids, phospholipids, and polyphrenyl compounds. Steroids, fat-soluble vitamins, and isoflavonoids (terpenoids) are polyphrenyl compounds. Omega-3- and omega-6-polyunsaturated fatty acids (PUFAs) are essential fatty acids [25].

**Isoxiprenoid**: Isoxiprenoids enhance antioxidant power by improving receptor functionality [25]. Farnesol (from floral essential oil) and geraniol (from citronella, geranium, lemon, palmarosa, and rose oils) have *in vivo* cytotoxicity against murine liver cancer, leukemia, and melanoma [32]. Cell membrane phospholipid bi-layers are protected from free radicals by isoprenoids [25]. Isoxiprenoids production by microorganisms including *Escherichia coli*, and *Saccharomyces cerevisiae* reduces the likelihood of supply limitations [33,34]. Therefore, it is biologically plausible and ecologically reasonable that isofoxiprenoid-phospholipid conjugates be studied for potential medicinal purposes [32].

**Omega-3-PUFAs**: Omega-3-PUFAs are immune boosting, platelet aggregating, anti-inflammatory [25]. Omega-3-PUFAs are consumed as docosahexaenoic acid (DHA) and eicosapentaenoic acid (EPA) [35]. Omega-3-PUFAs reduce coronary heart disease risk; decrease triglycerides, blood pressure, and inflammatory markers; improve endothelial function; prevent some cardiac arrhythmias; reduce vasoconstriction; enhance fibrinolysis and reduce fibrin formation; and decrease the risk of microalbuninuria and sudden cardiac death [35]. Omega-3-PUFAs form resolvins and protectins, suppress COX-2, IL-β, and TNF-α [35]. Omega-3-PUFAs bind peroxisome proliferator receptor activator (PPAR) gamma, allowing omega-3-PUFAs to inhibit cell proliferation and induce cancer cell apoptosis [35]. DHA has greater dose dependent pro-apoptotic activity against DU145 prostate carcinoma cells than does EPA. DHA’s pro-apoptotic activity is mediated via MAPK, NF-kβ, p53, and PI3K-Akt signaling pathways [36].

**Phenols**

Phenols which may be best known for the fruit and vegetable rainbow have a phenylalanine base. Coumarins, flavonoids, lignans, polyphenols, quinones, stilbenes, tannins, and xanthones, comprise the phenol category. Flavonoids in turn are comprised of anthocyanidins, flavones, flavanones, flavonols, and isoflavones. Each
Substituted coumarins are antiproliferative against breast and liver carcinomas [47]. Cyanidin-3-O-β-glucopyranoside (C3G) from flavanonols, the primary active ingredient of the homeopathic treatment Ruta [44]. Delphinidin, malvidin, peonidin, and petunidin, which are found in rice bran oil, tricin arrests MDA MB 468 breast cancer cells in G2/M phase [47]. Delphinidin is an anthocyanidin that differentiates Smilax bockii and S. glabra (sarsaparilla) from true Ceylon cinnamon, C. verum that has minuscule coumarin content [38]. Coumarin's reduction of melanoma recurrence and inhibition of renal-cell carcinoma did not withstand corroborating trials [37]. Coumarin has dose-dependent cytotoxicity against the Hep2 cell line [39]. Carbon-4 substituted coumarins are antiproliferative against breast and liver carcinomas [39]. Carbon-4 substituted coumarins inhibit aromatase, protein kinase, quinone reductase induction, sulphatase, 17β-HSD3, Cdc25, DNA intercalation, HDACs, Hsp90, microtubulin, NF-κB, and TNF-α [40]. Carbon-4 substituted coumarins are also down regulating selective estrogen receptor modulators (SERMs) [40]. A synthetic coumarin with a 4-position methylene thiol linker to a 6-membered heteroaromatic ring (4-Hydrazinyl-2-[(6-methyl-2-oxo-2H-chromen-4-yl) methyl]sulfanyl)-6-phenylpyrimidine-5-carbonitrile) is cytotoxic to MCF-7 breast cancer and HepG2 hepatocarcinoma cells, IC50 5.5 mg/mL and 6.9 mg/mL, respectively [39]. Potential cytotoxicity is indicated with IC50 of 4 μg/mL or 10 μM or less [41]. Ferula narthex Bio (an Ayurvedic spice called "Rauw") derived sesquiterpene (C15H24) chloroform soluble coumarin, C24H30O4 is an in vitro IC50 of 14.074 ± 0.414 μg/mL against PC3 cells [42]. Prediction of Activity Spectra achieved a probability to be active score of 0.303 with human histone acetyltransferase that should correlate with anti-pancreatic cancer activity [42].

Flavanoids: Flavanoids have a broad mechanism of action array. Flavanoids are antioxidants, anti-angiogenic, anti-proliferative, active against oxygen species scavengers, electrophiles, metal chelators, hydrogen peroxide producers, nitrosation inhibitors, and phase I detoxification enzyme modulators [25,26,43]. Flavones, flavanones, flavonols, flavanols, isoflavones, flavanoids, anthocyanidins and anthocyanins comprise the flavonoid category. Flavanoids including apigenin, kaempferol, nobiletin, quercetin, and rutin are found in cocoa, edible fruits, leafy vegetables, herbs, spices, legumes, tea and red wine [25,26]. Rutin, a compound flavanol-disacharide found in citrus fruits is the primary active ingredient of the homeopathic treatment Ruta [44]. Rice bran oil tricin arrests MDA MB 468 breast cancer cells in G2/M phase [45].

Anthocyanidins and anthocyanins: This group includes cyanidin, delphinidin, malvidin, peonidin, and petunidin, which are found in blueberries [43,46]. Carob fruit contains delphinidin, pelargonidin and cyanidin [47]. Delphinidin is an anthocyanidin that affects NF-κB regulation [47]. Cyanidin-3-0-β-glucopyranoside (C3G) from pigmented oranges, berries, and grapes activates caspase-3 in DU145 cells, and induces p21 protein expression in LnCap cells, resulting in antiproliferative effects against DU145 and LnCap prostate cancer cell lines [48]. C3G is associated with increased concentration of P53NGFR tumor suppressor from normal-like cell phenotype [48]. C3G antioxidant effect may be specific to androgen-independent tumors [48]. C3G has dose dependent ROS activity, requiring 100 μM [48].

Flavanonol: Dihydrokaempferol has been isolated from the roots of Smilax bockii and from S. glabra (sarsaparilla) rhizomes [49]. Both Smilax bockii and S. glabra are used in TCM [49]. The leaves of S. glycyphylla (sweet sarsaparilla) contain (2R,3R)-dihydrokaempferol-3-O-β-D-glucopyranoside [49]. S. china yields dihydrokaempferol and dihydrokaempferol-3-O-α-L-rhamnoside, which are less apoptotic to MCF-7 and MDA-MB-231 than are resveratrol and oxyresveratrol [49]. Taxifolin (dihydroxyquercetin) is a constituent of milk thistle seeds' silymarin extract and S. glabra rhizome [49]. Taxifolin-3-O-glycoside is one of the major active constituents of the tuber of S. china [49]. Like dihydrokaempferol and dihydrokaempferol-3-O-α-L-rhamnoside, taxifolin is weakly cytotoxic [50].

Flavanones: The flavanones hesperetin and naringenin undergo transformation by neohesperidose and rutinoside becoming flavanone glycosides hesperidin, neohesperidin, narirutin, and naringin [51]. Hesperetin is apoptotic to A431 epidermoid skin carcinoma via cyclin and MAPK regulation [51]. Naringenin is effective against A431, B16-F10 melanoma, and in vivo HepG2 cells [51]. In murine models, Naringenin dosed at 50 mg/kg body weight suppresses lung cancer [51]. Naringenin induces G0/G1 and G2/M phase cell cycle arrest, ROS generation, and down regulates TGF-β1 [51]. Naringenin induces mitochondrial apoptosis via Bax, Bcl-2, caspase-3, and p53 [51]. Naringenin modulates CYP1A1, NF-κB and PCNA [51]. Hesperidin (IC50=150.43 μM), is more cytotoxic than naringin and neohesperidin against HepG2, and is also anti-proliferative and apoptotic against A549 lung cancer and NCI-H358 non-small cell lung cancer (NSCLC) cells [51]. Hesperidin activates Erk, MMPS, mitochondrial calcium overload, ROS production, and membrane potential loss, while modulating fibroblast growth factor and NF-κB pathways [51]. Naringenin activates caspase-8 and caspase-9 [51]. Naringen inhibits MAPK-AP1 and IκKs-1βC-NF-κB pathways [51].

Flavones: Citrus fruit derived flavones include apigenin, chrysoeriol, diosmetin, and luteolin. Apigenin and luteolin have other sources, including celery and parsley [43]. Polyphenol-based flavones (PMF) are unique to citrus species’ peels, thus used in TCM [52]. Tangeretin, a tangerine derived PMF is anti-inflammatory in microglial cells, suppresses adipoocyte storage of triglycerides, and induces G1 arrest or apoptosis in cancer cells [52]. Tangeretin and nobiletin (another PMF) are antiproliferative to A549, MCF-7 and MDA-MB-435 breast cancer, HT-29 colon cancer, and HL-60 leukemia cell lines [51,53,54]. Nobiletin decreases Bcl-2 and increases Bax, leading to G2/M phase cell cycle arrest and apoptosis [51]. Nobiletin modulates hypoxia-TGF-induced epithelial-mesenchymal transition, inhibits notch-1 and TGF-β1/Smad3 signaling, while promoting microRNA-200b re-expression [51]. In vitro, the tangeretin derivative 5-AcTMF is antiproliferative to MCF-7, the U226 myeloma and CL1-5 NSCLC cell lines [52]. The tangeretin derivative 5-demethyltangeretin is more cytotoxic than tangeretin against A549, H460, and HI299 cell lines, with 79, 57 and 56 fold lower IC50 respectively [51]. The nobiletin derivative 5-demethylnobiletin has demonstrated in vivo antiproliferative activity against human NSCLC [51].

Flavonols: Tea is the source of several flavanols: Epicatechin (EC), epigallocatechin (EGC), epicatechin-3-gallate (ECG), EGCG, galloyl polyphenol, thearubigin, and theaflavins [43]. Green tea derived galloyl polyphenol and EGCG decrease retinoblastoma protein (Rb) phosphorylation, arresting cells in G1 phase [55]. The green tea polyphenols reactivate tumor suppressor genes by demethylation [55]. Green tea has the most EGCG, then white tea, and least of all black tea. Black tea derived thearubigin and theaflavins are anti-angiogenic [43]. The constituent epicatechin of Maytenus buchananii (a small evergreen

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tree), a traditional Cameroonian anti-cancer treatment, has an excellent MIC of 32 to 128 μg/ml against Staphylococcus aureus [56].

Other flavonols include quercetin from apples, berries, and red onions, and kaempferol from citrus fruits [43]. Luteolin and quercetin induce apoptosis in TRAIL-resistant cancer cells. Kaempferol, luteolin, and quercetin have shown dose dependent reduction in VEGF production in ovarian cancer growth and VEGF production [25]. Quercetin is a lignin metabolite, enterolactone, is a dose dependent agonist-cytotoxic to papillary urothelial carcinoma T24 cells, p<0.05 [57]. M. angiosperms, topoisomerase, tyrosine kinase, EGFR, IGFR-1 and NF-κB signaling pathways [26,43,58]. Genistein and other isoflavones, acting as SERMs, potentially balancing estrogen receptor α (ER-α)’s hyperproliferative effects in adipose tissue, the bone, breast, liver, ovarian thecal cells, prostate, and uterus [59].

Isoflavones: Like lignans, isoflavones are also phytoestrogens [43]. Daidzein, genistein, and kievetone have structural similarities to 17β-estradiol (E2). Daidzein, genistein, and kievetone elevate sex hormone-binding globulin (SHBG), are anti-oxidants, apoptotic, and inhibit angiogenesis, topoisomerase, tyrosine kinase, EGFR, IGF-1 and NF-kB signaling pathways [26,43,58]. Genistein and other isoflavones preferentially bind the pro-angiogenic estrogen receptor β (ER-β), functioning as SERMs, potentially balancing estrogen receptor α (ER-α)’s hyperproliferative effects in adipose tissue, the bone, breast, liver, ovarian thecal cells, prostate, and uterus [59].

Lignans: Lignans are also phytoestrogens. Lignans are derived from berries, flaxseed, sesame seed, whole grains [43]. Matairesinol and secoisolariciresinol are weak estrogen agonists, elevate SHBG, and inhibit aromatase [26]. Like the isoflavone phytoestrogen genistein, the lignin metabolite, enterolactone, is a dose dependent agonist-antagonist cancer specific modulator [43].

Polyphenols: Polyphenols are antioxidants. Polyphenols affect cancer by inhibiting cyclooxygenase, DNA topoisomerase, and DNA methylation, inducing phase II detoxification, and influencing cell signaling [25,26]. Polyphenols include cocoa, curcumin (found in turmeric), rice bran oil phenolics, caffeic acid, ferulic acid, and ferulic acid esters [45].

Quinones: Thymoquinone and dithymoquinone are derived from Nigella sativa L. seed (blackseed) [60]. Nigella sativa was the biblical and the prophet Mohammed’s curative black cumin, Hippocrates and Dioscorides’ Melanthion [60,61]. Nigella sativa is also used in Ayurveda and TCM [61]. Nigella sativa seed melanoic extract inhibition of Erlich ascites carcinoma in mice led to patenting of thymoquinone and dithymoquinone for additional research [60]. Thymoquinone and dithymoquinone displayed in vivo cytotoxicity to multidrug resistant cells [60]. Thymoquinone has cytotoxic activity against MCF-7, Doxorubicin via PTEN, p53, and p21 upregulation; NF-κB, P3K-Akt, phase I enzymes, and PPAR inhibition, and phase II enzyme (GST and N-acetyl transferase) activation [61]. Thymoquinone induces caspase-3 apotosis against Hep-2 cells, whereas caspase-3, -8, and -9 are activated against HL60 p53-myceloid leukemia [61]. Thymoquinone inhibits COX2, which affects breast, lung, stomach, and pancreatic cancers [61]. Thymoquinone inhibits MMPs and VEGF and induces apoptosis via Bak-Bax induction and Bcl-2 and Bcl-XL inhibition [61].

Thymoquinone is also active against bladder, colon, bone, and skin cancers [61].

The Plum bago (leadwort), Drosera (sundews), Nepenthes (monkey cups or tropical pitcher plants), and Juglans nigra (black walnut drupe) derived quinone plumbagin and the Ardisia elliptica (shoebutton ardisia) derived rapanone are under investigation as chemotherapeutics [41]. In vitro, plumbagin has an IC50 less than 10 μM against A549 (1.14 ± 0.02 μM), Caco2 colorectal adenocarcinoma (0.07 ± 0.01 μM), DLD-1 human colon adenocarcinoma cells (0.98 ± 0.11 μM), HepG2, MCF-7 (0.06 ± 0.01 μM), and SPC212 mesothelioma cell lines (0.27 ± 0.01 μM) (Kuete et al., 2016) [41]. Of these, plumbagin shows greater cytotoxicity than doxorubicin against Caco-2 (0.72 ± 0.13 μM) and MCF-7 (0.35 ± 0.05 μM) [41]. Plumbagin had previously shown good cytotoxicity against Caski, Colo-38, HeLa, MiaPaCa-2, PF-382 leukemia T-cells, and U87MG glioblastoma-astrocytoma cell lines [41]. Rapanone achieved an IC50 of 2.27 ± 1.52 μM against SPC212 cells, 46.62 μM against DLD-1 cells [41].

Stilbenes: Resveratrol, a berry, grape, and peanut derived polyphenol with stilbene derivates, upregulates cancer cell line wild-type p53 [43,62], Rhaspandigenin, a synthetic stilbene derivative of resveratrol, which also degrades HIF-1α (thereby suppressing VEGF and MAPK), has shown greater anti-angiogenesis effect than resveratrol [43]. Stilbenes also activate forkhead box (FOX) O apotosis transcription factors, while inhibiting Akt, IGF1, the PI3K-Akt and Ras-MEK pathways [43].

Tannins: Fruits’ astringency may derive from tannins [47]. Plant derived tannins are subdivided into hydrolysable tannins that are ellagic acid or gallic acid esters and condensed, non-hydrolysable tannins that are oligo- or polymeric proanthocyanidins [47]. Brown alge derived tannins are a separate class, phlorotannins. Tannins may have 12 or more hydroxy groups and five or more phenyl groups. Gallic acid and flavan-3-ols may be classified as pseudo tannins. Condensed, non-hydrolysable tannins, which include flavan-3-ols are also classified as flavanols, in which category catechins were mentioned above.

Hydrolysable or pyrogallol-type tannins: Punicalagin, ellagic acid, punicin can are ellagitannins, while gallic acid is a gallotannin. Carob pods have 0.95 mg/gm of hydrolysable tannins, while carob fruit have 0.237 mg/gm to 1.647 mg/gm of gallic acids, which is the third highest gallic acid content after chestnuts and cloves [47]. Carob fruit fiber contains at least three sets of gallic acid based tannins: Epigallocatechin with four gallic acid units, hexose with two or three cereal tannins which are classed as flavanols, in which category catechins were mentioned above.

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Szzygium cumini (Indian blackberry) or Eugenia jambolana, and Phyllanthus emblica or Emblica officinalis (Indian gooseberry, one of the three fruits from which Ayurvedic triphala is made) contain ellagic acid and gallic acid [63]. Indian gooseberry also contains the ellagittannins chebulagic acid, corilagin, emblicanin (also found in Amla), furosin, geraniin, isocorilagin (the α-anomer of corilagin), and pyrogallol (also found in chestnut and oak bark) [63]. Chebulagic, ellagic, and gallic acids, and corilagin inhibit NF-κB [63]. Corilagin is apoptotic to SKOV3ip, Hey, and HO-8910PM ovarian cancer cell line by inducing G2/M phase arrest and TGF-β/Akt/Erk/Smad modulation [63]. Cdc2, Cdk2, Cdk4, Cdk6, cyclin B1 and E inhibition by gallic acid is pro-apoptotic [63].

Punicalagin, derived from Punica granatum (pomengrate) is metabolized to ellagic acid [46]. Pomengrate tannins have
antiaromatase activity against MCF-7 cells that have been transected with the aromatase gene [64]. Punicalagin, ellagic acid, and total pomegranate tannin are apoptotic to HT-29 cells [64]. Punicalagin and total pomegranate tannin inhibit NF-κB response element binding and TNF-α mediated COX-2 expression [64]. Punicalagin is apoptotic to U87MG cells via PARP_breakage and caspase-3 and -9 activation [64]. Punicalagin also triggers autophagy cell death in U87MG cells by increasing AMPK-p27 [64].

Condensed or Non-hydrolysable tannins: Proanthocyanidins including flavan-3-ol groups and their gallocatechin esters form 2.75 mg/gm of carob pods [47]. Litchi contains proanthocyanidins A1, A2, and A6, and procyanidins B2 and B4 [63]. Proanthocyanidins are polymeric tannins that are depolymerized to anthocyanidins.

Xanthones: Pericarps of Garcinia mangostana (Mangosteen Linn fruit) contain α-Mangostin and gartanin [65,66]. α-Mangostin has antitumor activity against A549 cells via ROS-mediated apoptosis induction, but these effects are negated by N-acetyl cysteine [66]. α-Mangostin dose-dependently increases the Bax/Bcl-2 ratio, inhibits A549 cell migration, down regulates A549 antioxidants - catalase, glutathione peroxidase, and GSH [66]. Gartanin is dose dependently anti-migratory to T98G glioma cells via PI3K/Akt/mTOR signaling pathway autophagy modulated G1 cell cycle arrest [65]. Garatanin is anti-migratory to T98G cells via MAPK signaling pathway suppression of MMP-2 and -9 [65].

Phytic Acids: Phytic acids include defatted rice bran phytate and inositols [45]. Inositols include inositol phosphates, inositol hexakisphosphate (InsP6), and myo-Inositol [67]. Cereals, legumes, and oilseeds yield InsP6, which is broken down to myo-Inositol and inositol phosphates [67]. Inositol hexaphosphate inhibits proliferation [26]. Inositols inhibit cell cycles by inhibiting pRB phosphorylation and increasing pRB/E2F complex formation [67]. Inositols reduce PI3K, disrupt FGF receptor binding and EGF-transduction, down regulate Akt and Erk, inhibit NF-κB, and decrease COX2 and PGE2 expression [67]. Inositols impair β-catenin translocation, N-cadherin, Notch-1, SNA1 release, and Wnt-activation [67]. Inositols inhibit MMPs and ROCKI/2 release, decrease cofilin and fascin, upregulate focal adhesion kinase (FAK) and e-cadherin, impairing invasiveness and contributing to cancer microenvironment alterations [67]. DU145 tumors are reduced almost 64% by InsP6 [67].

Phytosterols: Lettuce, nuts, capers, flaxseed, cucumbers, rice bran are phytosterol sources. Stigmastaen is a soy-lipid derived phytosterol, and campesterol is a saw palmetto derived phytosterol [45]. β-Sitosterol does not stimulate the endometrium, nor do plant stanols and stanol esters stimulate ER-positive MCF-7 [68]. β-Sitosterol may be more biologically active with liposomal delivery [68]. β-Sitosterol down regulates NF-κB, thereby sensitizing cancer cells to TNF-α induced apoptosis [68]. β-Sitosterol reduces MDA-MB-231 triple negative breast cancer (TNBC) and MCF-7 tumors by 66% and 87% respectively, and low density lipoprotein by 8.8% [68]. β-Sitosterol’s activity against MDA-MB-231 is significant as although TNBC comprises only 15% of breast cancer, TNBC is treatment resistant, being unresponsive to endocrine and other targeted treatments [69]. β-Sitosterol and lupeol are the main anticancer constituents of Nardostachys jatamansi DC, a traditional Himalayan medicine herb used for cancer treatment [70].

Terpenes: Terpenes are single or multiple hydrocarbon compounds, categorized as saponins, and mono- or higher terpenoids, including tetraterpenes, based on the number of carbon atoms and isoprene residues [25].

Saponins: Saponins are antioxidants, immune modulators, and cell proliferation regulators [25,26]. Saponins are found in legumes and ginseng [26].

Betulonic acid: Betulonic acid from Betula alba (birch) bark, a plant triterpenoid saponin is a known chemopreventive, antiviral, anti-inflammatory, and antioxidant [71]. Betulonic acid increases IL-2, TNF-α, and CD4+ lymphocyte subsets. Betulonic acid increases the CD4+/CD8+ ratio in a dose dependent manner. Betulonic acid dosed at 200 mg/kg decreased B-cell lymphoma 2 (Bc1-2) protein and cell proliferation marker (Ki-67) protein expression and increased caspase-8 protein expression more than did 25 mg/kg cyclophosphamide, p<0.05 [72].

Escin: Escin from Aesculus hippocastanum (horse chestnuts) inhibits extracellular signal regulated kinase (ERK) and p38 mitogen-activated protein kinase (MAPK) cell proliferation, motility, and apoptosis pathways for anti-angiogenic endothelial cell migration and motility inhibition [43]. Crude ethyl acetate and methanol extracts of Mangifera pajarang (bambangan, a noncommercial mango) kernel, have strong to moderate cytotoxic activity against the HT-29 colon cancer cell line at IC50 of 10 μg/ml [72]. Crude petroleum ether and chloroform extracts of M. pajarang stem bark have strong activity at 15 μg/ml [72]. A component flavonol glycoside, querctin had very strong cytotoxic activity against HT-29 at IC50 of 3.82 ± 0.91 μg/ml [72].

Tripterine: Tripterine, the major active component of Tripterigium wilfordii (thunder duke vine), should be anti-angiogenic and cytotoxic due to modulation of MMP-9, intercellular adhesion molecule (ICAM)-1, vascular endothelial growth factor (VEGF), cell survival inhibitor of apoptosis protein (IAP)-1, X-linked (X)-IAP, Bcl-2, Bcl-xL, fllice inhibitory protein (cFLIP), survivin and cell proliferation molecules (cyclin D1, COX-2) in tumor cells [43]. At doses 500-fold higher than its anti-angiogenic activity, withaferin A from Withania somnifera (ashwagandha, Indian ginseng), has demonstrated direct anti-tumor activity in breast, pancreatic, and prostate cancers [43,73].

Monoterpenoids: Monoterpenoids induce apoptosis, cell differentiation, and phase I and II detoxification enzymes, including GST, affect cellular energy, and inhibit cell proliferation [25,26]. Monoterpenoids include limonene found in cardamom and Perillyl alcohol derived from the essential oils of lavandin, peppermint, spearmint, cherries and celery seeds. Ferula species derived sesquiterpene coumarin feselol has anti-tumor activity against U937 but not M14, MCF-7, T98G, A549, Saos-2, and FRO cell lines [74].

Diterpenoids: In gallbladder cancer, the diterpenoid oridonin derived from anti-angiogenic Radenosia rubescens (Dong Ling Cao), activates caspase-3 and 9 mitochondrial pathway apoptosis, activates PARP1 cleavage, increases the Bax/Bcl-2 ratio, and inhibits NF-κB nuclear translocation [69,75]. In acute lymphoblastic leukemia, oridonin inhibits the Akt-mTOR and Raf-MEK-ERK pathways, but in in vitro human osteosarcoma cells shows Akt and MAPK inhibition [69]. Oridonin treated murine L929 fibrosarcoma cells indicate Erk-
p53 activation leading to G2/M phase cell cycle arrest [69]. GSH induction by coffee-derived compounds including Kahweol palmitate, a diterpene ester, inhibits DMBA model breast cancer initiation [26].

**Tetramerpenes:** Carotenoids are anti-oxidant tetramerpenes [25]. β-carotene found in apricots, carrot, pumpkins, sweet potato is also antiangiogenic, anti proliferative, and induces cellular differentiation [26]. Astaxanthin, β-carotene, cryptoxanthin, lycopene, lutein, and zeaxanthin have breast cancer animal model anti-cancer activity [43]. Lycopene is found in tomatoes and watermelon, lutein and zeaxanthin are found in broccoli, cayenne and red pepper, corn, leafy greens (collard, kale, romaine lettuce, spinach, and turnip), paprika, and peas [76].

**A Combination Herbal**

 Yangzheng xiaoji, a 16 herb TCM dose dependently inhibits adhesion of A549, MCF-7, HRT18 colorectal cancer cells, HGC27 gastric cancer cells, and MG-63 osteosarcoma cells via the PI3K pathway and anti-angiogenesis [77]. The Yangzheng xiaoji extract DME25 inhibits cancer growth by inhibiting FAK pathway phosphorylation [77].

**Whole Plant Nutraceuticals**

**Cardamon**

In female Swiss albino mice, cardamom, garlic, and saffron are chemopreventive against DMBA induced skin cancer [28]. Cardamom stimulates phase II detoxification and anti-oxidation in female Swiss albino mice with DMBA treated skin and livers [28]. Cardamom contains numerous phytochemicals, some of which are mentioned below. Phase I trials of limonene, a major constituent of cardamom suggest breast, colorectal, and prostate cancer inhibition with minimal toxicity at doses up to 100 mg/kg [28]. D-Limonene is metabolized to perillyl alcohol, which inhibits G-protein isoprenylation [26]. D-Limonene inhibits gastric and lung cancers and leukemia [28]. 1,8-cineole inhibits and induces apoptosis in SK-MEL-28 human melanoma cells, B16-F1 murine melanoma cells, and Molt 4B and HL-60 human leukemia cells [28]. Linalool uses p53 upregulation against leukemia, and inhibits renal adenocarcinoma and amelanotic melanoma [28]. α-Pinene and α-Terpenine are anti-inflammatory to oral buccal cells [28]. Myrcene is chemopreventive against DMPA in rats, and apoptotic in human hepatoma cell lines. Trans-nerolidol is cytotoxic to A-549 human lung carcinoma cells and DLD-1 cells [28].

**Artemisia capillaris** (wormwood), which is known to inhibit CNE-2 nasopharyngeal carcinoma, DMBA induced breast cancer, SMMC-7721 human hepatoma, HL-60 leukemia, HepG2, Huh-7 human liver cells, HeLa cervical cancer, and mouse liver cells, contains borneol, which is also a constituent of cardamom [28,78]. Borneol is anti-oxidative, cytotoxic, DNA-damaging, and DNA-protective to Caco-2 and VH10 [28].

**Marine compounds**

Ecteinascidia turbinata, the source of the active ingredient of Trabectedin, approved for the treatment of platinum-sensitive ovarian cancer and tumor soft tissue sarcoma, displays in vitro and in vivo transcription factor inhibition [79]. Elysia rufescens derived Kahalalide F, is the basis for synthetic PM02734, which is in Phase II clinical trials for evaluation of H322 and A549 cell line apoptosis [79]. Kahalalide F inhibited breast, liver, and pancreatic cancer, as well as melanoma in Phase I trials. Spisula polysperma derived ES-285-HC1 has inhibited solid hepatocellular, prostate, and renal cancer in vivo [79]. Dictaethis orbita containing the indole derivatives tyrindolenine, tyrindolione, 6-bromoisatin and 6,6'-dibromoiindirubin, demonstrated in vivo apoptotic activity against azoxymethane exposed rat distal colon cells [79]. Dictaethis orbita derived tyrindolenine and 6-bromoisatin are twice as apoptotic to the KGN tumor-derived granulosa cell line than to normal primary human granulosa cells (HGC), 66% to 31%, respectively [79]. The biological plausibility of 6-bromoisatin containing compounds will be presented in a separate article on oncologic homeopathic remedies.

**Effect on Breast, Cervical, and Ovarian Cancers**

As known nutraceutical inhibition of endometrial cancer focuses on I3C, this section focuses on breast, cervical, and ovarian cancers [26].

**Breast cancer**

Breast cancer inhibitors include allium derivatives, β-carotene and other carotenoids, catechin, coumarin, curcumin, dithiolthiones, dried green coffee, fiber, glucosinolates and indoles (DIM and 13C), d-Limonene, isoflavones and lignans, glycirrhizic acid, kauehol palmitate, orange oil, protease inhibitors/Bowman-Birk inhibitor (BBI), rosemary extract, and selenium/organoselenium compounds [26,80]. Annual actual integrative cancer treatment costs for an early stage breast cancer patient are USD 1,594 [81]. An idealized integrative treatment program for stage 4 breast cancer that includes bromelain, coenzyme Q10, curcumin, green tea, intravenous artesunate and ascorbic acid, melatonin, mistletoe injections, Triametes versicolor mushroom, vitamin D3, and Wobenzym TM digestive enzymes has an annual total cost of USD 27,137.16 [81].

In murine studies BITC has proven MCF7 and HBL-100 inhibition by p53-LKB1 and p73-LKB1 pathway activation, which enhances p53 signaling [82]. As LKB1 activation is associated with improved breast cancer outcomes this finding is significant [82]. BITC increases breast cancer cell p53, Erk, and cyclic adenine monophosphate-response-element-binding protein (CREB) phosphorylation, but decreases Akt phosphorylation [82]. In vitro studies against MDA-MB-231, MDA-MB-468, BT-474, T47D, Hs578T, HCT116p53+/-, and HCT116p53+/- cells show BITC induced p53-independent apoptosis [82]. Bitter gourd derived triterpenoids have demonstrated moderate cytotoxicity to MCF-7 and MDA-MB-231 breast cancer cells, IC50 of 19 and 23 M respectively at 72 hours [83]. Bitter gourd derived triterpenoids have several mechanisms of action. In some cases, caspase-dependent apoptosis occurs [83]. Akt-NF-κB signaling is down-regulated, p38 mitogen activated protein kinase and p53 are up-regulated, ROS generation and cytoprotective autophagy are increased, and histone deacetylases (HDACs) protein expression is decreased [83]. In other cases, apoptotic peroxisome proliferator-activated receptor (PPAR) γ-targeted gene products modulation occurs [83].

Caffeic acid phenethyl ester from propolis, which is from floral resins of buds and sprouts of green trees that is collected by bees, has dose and time dependent strong cytotoxicity against MDA-MB-231, IC50 of 14.08 μM and excellent cytotoxicity against Hs578T, IC50 of 8.01 μM [84]. Caffeic acid phenethyl ester is caspase-3 dependent apoptotic and anti proliferative by NF-κB inhibition, which also increases the Bax: Bcl-2 ratio [84]. Cephalotaxus griffithii Hook. E (Griffith’s plum yew) needle derived flavonoids via petroleum ether...
The quinones plumbagin and rapanone induce caspase independent, ROS and MMP mediated MCF-7 cell apoptosis [41].

Crude ethyl acetate and methanol extracts of M. pajang kernel, and crude petroleum ether and chloroform extracts of the stem bark have strong cytotoxic activity against MCF-7 cells at IC50 of 10 μg/ml and 15 μg/ml, respectively [72]. The cytochrome c trimeric component, 3β-hydroxy-cycloart-24-ene-26-oic acid had moderate cytotoxic activity towards MCF-7 at IC50 13.03 ± 0.81 μg/ml [72]. Pseudovaria monticola leaf methanol extracts (6E,10E) isopolycerasoidol and (6E, 10E) isopolycerasoidol methyl ester are selectively anti-proliferative and cytotoxic to MCF-7 and MDA-MB-231 cells via ROS induced changes to mitochondrial membrane permeability, dose dependent caspase 3, 7, and 9 activation, anti-apoptotic Bcl-2 down regulation, and nuclear accumulation of p38 MAPK [90]. At 48 hours, (6E,10E) isopolycerasoidol had IC50 59 ± 5.1 M against MCF-7 and IC50 76 ± 8.5 M against MDA-MB-231, while (6E,10E) isopolycerasoidol methyl ester is more potent at IC50 43 ± 2.4 against MCF-7 and IC50 58 ± 2.6 against MDA-MB-231 [90]. Normal human breast epithelial MCF-10A cells are comparatively spared with IC50 of 94 ± 5.9 M for (6E,10E) isopolycerasoidol, and IC50 90 ± 4.7 M for (6E,10E) isopolycerasoidol methyl ester [90].

Silibinin, the main active flavonoid component of Silymarin, derived from Silibium marianum (milk thistle) seeds, is anti-angiogenic by MMP-2 and capillary formation suppression, and inhibition of angiogenic signaling molecules [43,77]. Silibinin induces caspase cascade apoptosis in TRAIL-resistant cancers, down regulates Akt-mediated NF-kB1 [77]. Silibinin inhibits MDA-MB-231 breast cancer cell line metastases by inhibiting mRNA levels of GDP dissociation inhibitor (D4-GDI) expression and cell division cycle 42 [77].

Withaferin A has numerous mechanisms of action: Inactivation of Akt and NF-kB to achieve apoptosis, decrease in pro-survival protein B-cell lymphoma 2 (Bcl-2), gap-2-mitosis (G2/M) cell cycle arrest, generation of ROS, induction of protease-activated receptor-4 (Par-4), activation of caspase 3 and 9 activities, DNA damage, inhibition of molecular chaperone heat shock protein 90 (HSP90), regulation of forkhead box O 3a (FOXO3a) and Bcl-2 interacting mediator of cell death (Bim), and inhibition of Notch-1 [73,91,92]. For breast cancer inhibition, withaferin A activates the extracellular signal–regulated kinase (ERK)-ribosomal S6 kinase (RSK) and ETS-like transcription factor 1 (Elk1)-C-EBP homologous protein (CHOP) kinase pathways which upregulate DR5 transcription [92]. This mechanism also provides synergism with celecoxib, etoposide, and TRAIL [92].

Cervical Cancer

Betulinic acid dosed at 100 mg/kg and 200 mg/kg showed dose dependent in vivo anti-tumor activity in U14 cervical cancer bearing mice, which approximated to that of 25 mg/kg cyclophosphamide, p <0.05 [71]. Conium maculatum (hemlock) ethanolic extract is dose and time dependent anti-proliferative via sub-G stage cell cycle arrest by 48 hours [93]. At 24 hours ROS generation is increased. By 48 hours MMP depolarization, phosphatidyl serine externalization, morphological changes, DNA damage, decreased Bcl-2:Bax, and caspase-3 dependent apoptosis occur in HeLa cells [93]. C. maculatum ethanolic extract is also effective against A375, A549, and HepG2 cell lines, with WRL68 normal liver cells surviving treatment better than PBMCs, indicating limited harm to normal tissue [93].

Crude ethyl acetate and methanol extracts of M. pajang kernel, and crude petroleum ether and chloroform extracts of the stem bark have strong cytotoxicity to ZR75B breast cancer cells, IC50 22.3 ± 3.9 μg/ml [85]. Curcumin and sulforaphane, also modulate HDACs for breast cancer antiproliferation [83]. The ML-1 lectin containing mistletoe extracts Iscador M special and Iscador Qu, dosed at 15 g/dl have strong cytotoxic activity to the MAXF-401-NL breast cancer line [80].

In murine studies, CYD-6-28, a synthetic analog of oridonin, a diterpenoid terpene, dosed at 5 mg/kg, inhibited TNBC cell lines HCC1806 and HCC1937 [69]. CYD-6-28 induces Erk, p21, caspase-3, -7, -8 and PARP cleavage, G2/M-phase cell cycle arrest, and death receptor 5 (DR5) mediated apoptosis [69]. CYD-6-28 inhibits Akt, cyclin D1, FLIPL, STAT3, and XIAP [69], CYD-6-28’s efficacy may be due to improved aqueous solubility and cell permeability from a thiazole ring at A-ring C-1 and C-2 [69].

Edible Phaleria macrocarpa (Scheff.) Boerl fruit (mahkota dewa) of Indonesian traditional medicine has flavonoid, flavonol, phenol, saponin, and terpenoid derivatives [86]. Ethyl acetate and methanol extracts have strong cytotoxicity comparable to standard gallic acid, with IC50 8.15 ± 0.02 g/ml in DPPH antioxidant assay [86]. Gallic acid is cytotoxic to brain, breast, cervix, colon, esophageal, and gastric cancer cell lines [86]. The chloroform extract has excellent cytotoxicity against MDA-MB-231 breast cancer cells, IC50 7.80 ± 1.57 μg/ml at 48 hours [86]. The P. macrocarpa hexane extract has excellent cytotoxicity against MDA-MB-231, IC50 4.6 ± 1.32 μg/ml at 48 hours, while the ethyl acetate extract has good cytotoxicity against MDA-MB231, IC50 6.4 ± 1.09 μg/ml at 48 hours [86]. Ethanolic mango seed extract may be apoptotic to MCF-7 cells via ROS induction [87]. Ethanolic kernel extract of Mangifera indica L. (waterlily mango) has dose dependent strong cytotoxicity to MCF-7, IC50 15.6 μg/ml in DPPH antioxidant assay [86]. Gallic acid is cytotoxic to breast epithelial cells [89]. Luteolin was also found to be more active than the twice as available coumarin wedelolactone component of E. edule phytochemical profile [85]. Ethanolic mango seed extract of Mangifera indica L. (waterlily mango) has dose dependent cytotoxicity to MCF-7, IC50 15.6 μg/ml. Future studies should explore the efficacy of lower concentrations.

Nobiletin and tangerin are dose dependently antiproliferative to MCF-7 and MDA-MB-435 cells via G1 cell cycle phase arrest [51]. Naringin increases p21, modulates the β-catenin pathway, and induces G1 phase cell cycle arrest, resulting in TNBC apoptosis and anti-proliferation [51]. Herespin activates caspase-3 and p53 mediated apoptosis against MCF-7 cells [51]. Hesperitin is antiproliferative to MDA-MB-231 cells by GLUT1 and GLUT4 down regulation, and IR-β and Akt phosphorylation inhibition, which reduces basal and insulin-stimulated glucose uptake [51]. Naringenin inhibits MCF-7 basal and insulin-stimulated glucose uptake via PI3K and MEK inhibition [51]. Hesperitin suppresses in vivo aromatase expression; therefore, it is biologically plausible that hesperitin will be effective against ER+ breast cancers [51]. Hesperitin and naringenin have in vitro activity against HER2 positive cancers, which is biologically plausible as hesperitin and naringenin are HER2 tyrosine kinase inhibitors [51].

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strong cytotoxic activity against the HeLa cervical cancer cell line at IC50 of 10 μg/ml and 15 μg/ml, respectively [72]. 3β-hydroxy-cycloart-24-ene-26-oic acid had strong anti-cancer activity towards MCF-7 cells at IC50 6.67 ± 0.61 μg/ml [72]. Quercetin had moderate to strong cytotoxic activity at 11.93 ± 0.63 μg/ml [72]. Oridonin inhibits the HeLa cell line P13K-Akt pathway [69].

Curcumin and emodin down regulate TGF-β receptor II, P-Smad3, and Smad4, which inhibits TGF-β induced migration and invasion in HeLa and SiHa cervical cancer cell lines [94]. Curcumin and emodin up regulate the Bax:Bcl-2 ratio which inhibits p15, p16, p21, p27, CDK6, CyclinD1, and Pin1 expression, thereby inhibiting Wnt/β-catenin in HeLa cells [94]. SiHa cells were chosen for TGF-β resistance [94]. Curcumin and emodin are synergistic. Curcumin and emodin were dosed at IC50: For curcumin, 15 μM in SiHa and 25 μM in HeLa cells, representing strong cytotoxicity. For emodin, 40 μM in SiHa and HeLa cells [94]. Curcumin and emodin may not be as good candidates for cervical cancer inhibition as M. pajiang extracts, but, if tolerated, curcumin offers additional overall health benefits, such that curcumin and emodin for chemosensitization should not be overlooked.

The phenol, naringin modulates GM3 ganglioside, affecting EGFR signaling, resulting in HeLa suppression [51]. Taxifolin is synergistic with andrographolide (a diterpenoid lactone) against HeLa cells by reducing andrographolide induced protective autophagy, while increasing mitochondrial outer-membrane permeabilization, caspase dependent, and independent cell death [50].

In vivo athymic nude mice models show that withaferin A, dosed at 8 mg/kg intraperitoneally on alternate days, can reduce CaSkii human papilloma virus (HPV) types 16 and 18 positive, cervical cancer cell line tumors by 70%, p<0.01 at 6 weeks of treatment [73]. The withaferin A derivative, 3-azido Withaferin A (3-azidoWA), irreversibly inhibits MMP-2 via dose dependent pro-apoptotic extracellular par-4 secretion, resulting in suppressed motility and invasion of HeLa cervical and PC-3 prostate cancer cells lines (p<0.01) [95]. In vivo mice studies indicate 3-azidoWA, dosed at 30 mg/kg/day intraperitoneally, dosed on day 8 or days 8 to 10, dose dependently activates capase-3 mediated apoptosis and is dose dependently anti-angiogenic [95]. Withaferin A down regulates human papilloma virus (HPV) E6 and E7 oncoproteins’ expression, in turn reactivating p53-dependent tumor suppression [73,91].

Ovarian Cancer

The 71 and 87 percentage point survival advantages between Stage 1 and Stages IIIC and IV ovarian cancer respectively indicates a need for more effective treatment modalities for advanced stage ovarian cancer and mechanisms for ovarian cancer prevention [62]. However, a brief literature search suggests that much more could be done to find nutraceutical therapies targeted to ovarian cancer treatment.

The P. macrocarpa methanol extract has excellent cytotoxicity against SKOV-3 ovarian cancer cells, IC50 7.75 ± 2.56 μg/ml at 72 hours [86]. The P. macrocarpa hexane extracts have good cytotoxicity against SKOV-3, IC50 10.15 ± 2.71 μg/ml at 24 hours [86]. The P. macrocarpa ethyl acetate extracts have good cytotoxicity against SKOV-3, IC50 8.1 ± 1.81 μg/ml at 72 hours [86]. Withaferin A inhibits ovarian adenocarcinoma cell lines CaOV3 and SKOV3 by down regulating Notch1, Notch3, Cdc25C, total and phosphorylated Akt, and bcl-2 proteins, which in turn induces early apoptosis (p<0.05) and G2/M phase cell cycle arrest [96].

Future Research

Research is needed to determine which of the whole plant or organism, the natural phytochemical component, the semi-synthetic phytochemical, or the completely synthetic phytochemical, is the most efficacious form of a nutraceutical. Crude extracts can derive anti-cancer activity from phytochemical components that are removed with additional processing. This is the case with Ferula extracts that lose phenolic components when purified to terpenoid coumarins [74]. However, components of the whole plant or intact portions of the plant can have antagonistic effects, necessitating separation for maximal beneficial effect. For instance, quercetin appears to antagonize M. pajiang from which it is derived [72]. Trials of I3C should evaluate the concentration of the I3C metabolite DIM present following administration and metabolism of I3C [28]. Studies on cineole should determine the equivalence, if any, between 1,4-Cineole, the major phytochemical in cardamom, and 1,8-cineole, the most studied cineole [28]. Similarly, clarity is needed as to the activity of each isomer of limonene. All studies on isomeric phytochemicals should specify the isomer used for clarity, efficacy, and adverse effect ascertainment.

Similarly, delivery modality effectiveness for nutraceuticals displays enormous variation. Improving oral nutraceutical bioavailability is an emerging discipline [97]. While nano-encapsulated EGCG is 10-fold more efficacious than non-encapsulated EGCG, and berberine and curcumin act similarly, kaempferol is significantly less efficacious when encapsulated [43]. Nutraceutical microbial fermentation can improve bioavailability and antioxidant efficacy [45]. Therefore, the optimal delivery modality for each nutraceutical’s optimal efficacy should be determined. In this light, quercetin is being studied as an enhancer for green tea polyphenol bioavailability and activity [98]. Pro-drug derivatives may need to be developed [52]. Similarly, with synthetic analogs, bioavailability enhancement may be possible without adverse effects as displayed by CYD-6-28 [69]. Then, the focus can shift to making manufacturers supply pharmaceutical grade nutraceuticals in the most efficacious delivery modality.

In vivo trials of withaferin A for ovarian adenocarcinoma treatment are a logical progression from successful in vitro CaOV3 and SKOV3 cell line studies [96]. Additional trials of M. pajiang may indicate if a breast, cervix, or colon cancer treatment can be realistically pursued [72]. The P. macrocarpa extracts can be further studied for breast and ovarian cancer [86]. The main cytotoxic constituents of M. indica L’s ethanolic kernel extract could be determined and studied in vivo [88]. C. maculatum ethanolic extract should undergo in vivo murine trials against A375, A549, HeLa, HepG2, and WRL68 cell lines, to assess efficacy and tolerability. Tangeretin, nobiletin and 5-AcTMSF should be further trialed for breast, colon, leukemia, myeloma, and NSCLC treatment [52-54]. BITC should be further studied for breast cancer treatment [82].

Clinical trials of sillinbin, G. lucidum polysaccharides and ganoderic acids, cordycepin from C. militaris, EPA, γ-linolenic acid, Yangzheng Xiaoji and DME25 may lead to antimitastatic drug candidates [77]. Comparative effectiveness trials of EGCG, sillinbin, curcumin, melatonin, oleanolic acid, resveratrol, withaferin A, enterolactone, kaempferol, and triterpene, listed from theoretically most anti-angiogenic activity to least anti-angiogenic activity, could determine which of these polyphenol nutraceuticals is the most anti-angiogenic for a given cancer [43]. These nutraceuticals can also be trialed with conventional chemoradiation to determine if synergism will permit chemoradiation dose reduction.
Nutraceutical treatment of TNBC is worthy of further investigation. β-Sitosterol, bitter gourd, caffic acid phenethyl ester from propolis, chloroform extracted luteolin from Eclipta alba, Phaleria macrocarpa chloroform and hexane nobiletin extracts, Pseudovaria monticola leaf menthol extracts (6E,10E) isopolycerasoidol and (6E,10E) isopolycerasoidol methyl ester, silibinin, and the tangeretin derivative 5-AcTMF may be potential drug candidates. Microenvironmental efficacy limitations of thymoquinone and its synthetic analogs should be performed to ascertain if microenvironmental effects limit thymoquinone’s in vivo efficacy against bladder, bone, breast, lung, stomach, and pancreatic cancer cell lines, in addition to known limitations of thymoquinone in ID8-NGL mouse ovarian cancer [61,99].

Conclusion

Some nutraceuticals such as the Ferula species derived phytoestrogenic monoterpene umbelliprenin have very limited curative anti-cancer effects that can be limited to a single cancer. Other nutraceuticals, such as the natural MMP inhibitors, 3-azido WA, aqueous cinnamon extract, green tea extract, curcumin, fenugreek derived steroidal saponin, and marine compound derived chitooligosacharides should have a broader applicability. When the incidence and severity of a cancer, for example TNBC is considered, there is benefit in developing a nutraceutical that initially seems to be exclusively of single indication benefit. The breadth and depth of the nutraceutical clinical pharmacology literature suggests that an increased range of curative nutraceutical cancer treatments will become available. However, endometrial cancer treatment should be given as much consideration as have breast, cervical, and ovarian cancers.

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