Phytochemicals for Controlling Obesity-Related Cancers

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Abstract

Obesity related diseases are on the rise worldwide and obesity is an economic burden on the public health system in most developed and developing countries. The control of obesity is a major challenge that is hard to solve using medication and surgical procedures as those often have serious side effects. Our focus in this article is to highlight the extensive research literature on use of plant-derived chemicals or phytochemicals for control of obesity and obesity related diseases with a particular focus on several types of cancer. We discuss the genes, proteins and pathways involved in obesity and its control, and discuss how current research has revealed the beneficial effects of plant-derived chemicals or phytochemicals on these genes and pathways.

Keywords: Obesity; Phytochemicals; Cancer; Adiposity; BMI; Signalling pathways; Genes

Introduction

Obesity is the state of being overweight, a condition caused due to excessive rate of accumulation and storage of fat in the body. Obesity not only affects appearance, but leads to a number of health issues. The disparity in calorie consumption and calorie burning leads to energy imbalance, which is a lead cause of obesity. Body mass index (BMI) is a measure of obesity or adiposity. BMI is defined as the ratio of the weight and square of the height. The root cause of obesity may be due to uncurbed intake of food, sedentary life style without any physical activity and genetic susceptibility [1]. Endocrine disorders, genes, mental disorder and sometimes medications also contribute to obesity [2]. Many obese people eat very little, yet they are overweight due to slow metabolism [3].

Obesity has become an epidemic worldwide in the modern age and millions of people die every year due to obesity. The prevalence of obesity across the globe has become three times since 1975 [4]. Malnutrition is a known cause of death, but deaths due to being overweight is sometimes more alarming than deaths from being underweight. Overweight individuals are often at a higher risk of developing a plethora of medical problems. Comorbidity is a big problem with obese patients. The upward trend of obesity has a tremendous impact on economy, significant comorbidities and hence diminished lifespan in the population [5]. The associated health complications include type 2 diabetes mellitus (T2DM), insulin resistance, hypertension, cardiovascular disease, stroke, arthritis, gout, dyslipidemia, sleep apnea, gastroesophageal reflux, polycystic ovarian disease (PCD) and infertility [6]. Obesity is often associated with psychiatric problems and behavioral alterations. Mood disorders, personality problems, and sometimes schizophrenia may be caused due to obesity [5]. Obesity is the cause of a number of cancers like breast, endometrial and gallbladder cancer in women and prostate and colorectal cancer in men [7]. Obesity enhances mortality from several cancers like colon, kidney, liver, esophagus, gallbladder, multiple myeloma and non-Hodgkin’s lymphoma [8].

Causes of Obesity

The fat cells once consumed as food, remain for an indefinite period in our body. The size of fat cells can be reduced, but we cannot remove fat cells except for metabolic breaking down of the cells. Therefore, it is important to address the approaches for obesity prevention.
and control that can prevent or reduce accumulation of fat cells in the body. In order to do so, it is important to know the risk factors for obesity. These are excessive intake of energy rich food, junk food with more calorie and less nutrients, very little physical activity, sedentary lifestyle, very little or too much of sleep, genetic factors, family history, racial/ethnic differences, depression or mental stress, intake of certain drugs like steroids etc., lack of understanding of nutrition, poor socio-economic and socio-cultural status to name a few. A number of genetic markers are known, which are responsible for increased susceptibility to obesity [9]. Obesity is responsible for the initiation and progression of many types of cancers. Obesity causes low level of inflammation, which may cause damage to the DNA. Chronic inflammation is a risk factor for many cancers [10]. Chronic inflammation of the gallbladder may be caused by gallstones, which are quite common in obese individuals. Therefore, gallbladder cancer may be one of the cancers caused by obesity [11]. One of the causes of esophageal cancer is Barrett disease or gastroesophageal reflux disease, which may be caused due to inflammation. Another inflammatory condition is chronic ulcerative colitis. Hepatitis is also a liver disease, caused by inflammation and both may cause liver cancer [12]. Fat tissues are deposited due to obesity and as a result an excess amount of estrogen is secreted. Overproduction of estrogen enhances the risk of ovarian, breast and endometrial cancers. Overweight individuals generally have high insulin and insulin like growth factor-1 (IGF-1) levels. These conditions increase the risk of colon, endometrial, prostate and kidney cancers [13].

Fat cells in obese people regulate two important proteins, i.e. AMP-activated protein kinase and mammalian target of rapamycin (mTOR). Both are implicated in carcinogenesis. Scaffolding that surrounds breast cells is altered due to obesity [14], also, immune responses are altered, which in turn alter NF-κB and oxidative stress [15]. Therefore, it is clear that excessive body fat increases the risk for a number of cancers. These cancers include breast, endometrial, esophageal, colorectal, gallbladder, gastric, kidney, liver, multiple myeloma, meningioma, ovarian, pancreatic, thyroid etc. BMI, which gives a good measure of obesity, compared to weight alone, is associated with some of the cancers. High BMI is related to breast, colorectal, ovarian and thyroid cancers.

**Genes and Pathways in Obesity Related Cancers**

As mentioned before, obesity has been implicated in several cancers. Metabolic conditions due to obesity quite often result in facilitating tumor growth. We will discuss in this section some of the key metabolic tendencies due to obesity, and how they are related to genes and genetic pathways.

**Genes and proteins related to obesity**

One of the common risk factors for Type 2 diabetes mellitus (T2DM) is obesity. Insulin resistance is the primary cause for T2DM, and insulin resistance results in hyperinsulinemia, the state in which an elevated level of insulin circulates in the blood. This is due to the fact that the receptors on the cells become less sensitive to insulin and as a result excess amount of glucose accumulates in the cells. Elevated levels of insulin may further cause growths in tissues, and may also increase the levels of insulin-like growth factor 1 (IGF-1). The elevated levels of IGF-1 interact with cell surface receptors and promote tumor growth. There are two main genetic pathways involved in the tumor promoting growth due to elevated levels of insulin and IGF-1, the PI3K-AKT-mTOR and the Ras-Raf-MEK-MAPK [16-18]. The literature on use of phytochemicals in cancer prevention and treatment, as well as their effects on the genes and pathways mentioned in this section is vast. We only give some examples here, and refer the reader to the book by Roy and Datta [19] for more details.

Adipose tissue is a loose connective tissue made of adipocytes. The main role of adipose tissue is to store energy in the form of lipids, cushioning and insulation. It is one of the major endocrine organs, producing hormones like estrogen, resistin, leptin, and cytokines like Tumor Necrosis Factor-α (TNF-α). White adipose tissue (WAT) and brown adipose tissue (BAT) are the two types of adipose tissues present in humans and other mammals [20]. They have different colors, functions, different cellular compositions and localization. WAT is the major part of body’s adipose tissue and constitutes most of the total body fat and is also the source of free fatty acids (FFA). Its excessive accumulation might lead to obesity and obesity-related diseases. Most common is “android obesity”, the excess WAT in the upper parts of body. This kind of obesity represents a strong risk factor for several inflammatory pathologies. “Gynoid obesity” is the excess WAT in lower body parts and usually does not have much metabolic complications [20,21]. BAT on the other hand is mainly for adaptation due to cold, generates energy or heat from nonoxidative phosphorylation [22]. Fat cells in BAT are smaller in number, and have richer vascular supplies. They also have more abundant mitochondrial chromogens, responsible for the brown color. Positron emission tomography demonstrates that in adult humans metabolically active BAT are found in axillary, cervical, paravertebral and supraclavicular regions, which are induced in response to cold [23]. BAT is a potential target for both gene expression manipulation and pharmacological intervention for combating human obesity [24,25].

Adipose tissue is the source of soluble molecules (adipokines), secreted by adipocytes. Adipokines like Leptin, Adiponectin, Interleukin-6 (IL-6), TNF-α etc are linked to inflammation and there is a hypothesis that elevated inflammatory adipokines contribute to carcinogenesis. Precise ratio between Adiponectin to Leptin is crucial in cancer. TNF-α is secreted by adipocytes and was the first identified inflammatory cytokine. TNF-α expression in white adipose tissue (WAT) was first shown in rodents, and was significantly increased in obese models [26]. TNF-α may play a role in the development of insulin resistance in several ways, including the inhibition of insulin receptor signaling pathway [27,28]. It is not clearly understood to which extent TNF-α produced in adipose tissue goes into circulation, although it is known that there is a correlation between the TNF-α system (including the soluble receptors) and indices of obesity [29]. It is known that TNF-α is an important regulator for the synthesis of IL-6 [30]. It is also responsible for the biosynthesis of estrogen in adipose tissue in vivo by stimulating aromatase expression [31]. Therefore, there
could be an increase in circulating TNF-α in obese individuals that could contribute to insulin resistance and in turn may lead to breast tumorigenesis by regulating the synthesis of IL-6. IL-6 acts both locally and in other parts through circulation and is secreted by adipocytes. It has been reported that the expression of IL-6 and plasma levels are elevated in adipose tissue of individuals with obesity and insulin resistance [32,33]. It has been proposed that IL-6, along with Leptin, is responsible for the regulation of energy balance by conveying information to the hypothalamus from adipocytes. IL-6 has been reported to be an inhibitor in early-stage breast cancer, however high serum level of IL-6 is associated with poor prognosis in metastatic breast cancer in advanced stages. IL-6 is also known to stimulate aromatase expression in adipose tissue in vivo, thereby stimulating estrogren biosynthesis [34], and possibly directly contributing to breast cancer progression.

As adipose tissue expands in obesity, levels of these adipokines increase in blood and risks of several cancers grow [35,36]. TNF-α levels have been directly correlated with higher body mass index. TNF-α has regulatory effects on several other adipokines and collectively they can promote oncogenesis through inducing cell survival. TNF-α also has anti-apoptotic potential in some tumors through stimulation of NF-κB. Higher level of insulin in circulation promotes higher levels of IL-6 and TNF-β in adipose tissues. Higher levels of IL-6 have been observed in patients with hepatocellular carcinoma and ovarian cancer. IL-6 seems to act through the activation of the JAK/STAT pathway, one of the most important pathways in regulating proliferation and anti-apoptotic actions in tumor cells [37,38]. IL-6 also acts on the PI3K-AKT-mTOR pathway and that increases the expression of cyclin D1, a cell survivor factor [39]. Chronic inflammation has been implicated in the initiation and progression of tumors, through the elevated levels of cytokines like TNF-α and IL-6. Systematic inflammation is one of the major causes of colorectal adenoma [40,41].

Mutations in leptin receptor gene or leptine gene may result in aberrant leptine action and may result in obesity and obesity related diseases [42]. Investigations in human tumor cell lines and mouse models indicate that leptine balance has an effect on pathways that regulate growth, including NF-κB, PI3K-AKT-mTOR and JAK/STAT [43,44]. In recent research, synthetic nanopeptides have been observed to block leptine effects and as a result interfere with these pathways both in vitro and in vivo. The leptin levels are higher among obese individuals compared to their leaner counterparts. This indicates a correlation of leptine levels with the total body fat, and increases the risk of several cancers. For example postmenopausal women with higher body fats have the highest risk for breast cancer [45].

The Janus kinase/signal transducer and activator of transcription (Jak/Stat) pathway is a mediator of cytokine signaling. The Jak/Stat pathway is central to the signalling by various cytokine receptor systems and regulates various fundamental cell functions. IL-6–mediated activation of Stat3 is a principal pathway implicated in promoting tumorigenesis [46]. Aberrant activation of this pathway results in neoplastic transformation and abnormal growth in malignancies e.g., breast cancer. It is unclear how Jak/Stat pathway modulates the pathogenesis of breast cancer. Vascular endothelial growth factor (VEGF), a key angiogenic factor is an important biological marker for breast cancer malignancy and progression [47], which is controlled by pro-inflammatory cytokines like TNF-α. Obesity has been reported to increase tumor growth rate by upregulation of VEGF pathways [48,49]. Obesity is an independent prognostic factor for the development of metastases after breast cancer diagnosis [50]. Leptin, the main adipokine secreted by adipose tissue, is also abnormally expressed by breast cancer cells [51]. The metastatic proteins matrix metalloproteinases (MMPs), particularly MMP-2 and MMP-9 are regulated by Leptin, as evident by the decrease in these two metastatic proteins following Leptin knockdown [52].

Reactive oxygen species (ROS) cause inflammation which is a cause for many types of cancer. It has been observed that elevated levels of insulin increased mitochondrial generation of Reactive oxygen species (ROS) in several cancer cell lines [53]. These elevated levels of ROS cause DNA damage that further exacerbates carcinogenesis. It has also been observed that high-fat diets induce obesity and serve as tumor promoters, resulting in increased number of tumors, as well as larger tumors [54]. This may also cause a higher level of metastasis. The mutation in the Pac gene in mouse provides a suitable model for studying increased levels of colon cancer due to hereditary causes in humans. High fat diets in the mouse model have been observed to induce inflammation through up-regulation of the NF-κB pathway.

Some important pathways implicated in obesity

PI3K-AKT-mTOR is an intracellular pathway that does not depend on any extracellular signalling [55,56]. The main purpose of this pathway is to regulate cell cycle and control cellular proliferation. This pathway is overactive in many cancers by suppressing apoptosis, and increasing cell proliferation, the two major hallmarks of cancer. PI3K are lipid kinases that phosphorylate PI2P into PI3P. The action of PI3P is to activate kinases with the pleckstrin homology domain or PH. An example of such a kinase is PDK1. PTEN inhibits PI3K by controlling the signalling of PI3P. AKT also has a PH domain, and is activated by PDK1 and mTORC2. AKT is capable of phosphorylating many target proteins in duding AKT1, GSK3 and TSC2. Hence, AKT has extensive downstream effects and can control apoptosis and cell proliferation. This pathway is active in ovarian, breast and urothelial cancers.

The signal for the Ras-Raf-MEK-MAPK pathway comes from outside a cell, and eventually reaches the nucleus [57]. Mitogens are the signalling molecules for controlling cell division and proliferation. An example of a mitogen is the EGF (epidermal growth factor). The main molecules involved in this pathway are Ras, RTK, MEK, RAF and ERK. The receptor EGFR binds to the external signalling molecule EGF. This is the activation signal for the pathway. The next step is the phosphorylation of EGFR and the binding of a complex of the docking protein SOS and GRB2 to the phosphorylated EGFR. This activates SOS and the action of SOS is to remove GDP from members of the Ras family of proteins. As a result, Ras binds to GTP, and activates the RAF kinase. RAF
phosphorylates MEK, which in turn phosphorylates MAPK. MAPK
then phosphorylates several transcription factors, which in turn
regulate the transcription of several genes that control the cell cycle
progression.

JAK/STAT is a complex signalling pathway involving many
proteins. It is activated by extracellular ligands and the signal is
finally communicated to the nucleus where transcription factors
regulate gene expression. This pathway is involved in cell cycle
progression, apoptosis and tumor formation and progression,
and as mentioned before, a disruption of this pathway has been
implicated in several cancers. Our brief review of this pathway is
based on the paper by Schindler et al. [58]. The main players in this
pathway are the four JAK proteins, JAK1-3 and TYK2; and the seven
STAT proteins, STAT1-4, STAT5A, STAT5B and STAT6. Cytokines like
IL-6 first binds to cell surface receptor and dimerize the receptors.
This results in the JAK proteins to come in contact with the
receptors, and the JAK proteins phosphorylate each other through
a process called transphosphorylation. The phosphorylation occurs
on the tyrosine residues of the JAKs, and increases the activity of
their kinase domains. In the next step, the JAKs phosphorylate the
tyrosine residues of the receptor and this creates binding sites for
the SH2 domains of the STAT proteins. Once the STATs bind to the
receptor, the JAKs tyrosine-phosphorylate them and this causes
the STATs to disassociate from the receptors. The SH2 domain of
a STAT binds to the phosphorylated tyrosine of the opposite STAT,
forming a dimer. This dimer translocates to the nucleus and acts as
transcription factor for the target genes.

The Keap1-Nrf2 pathway is triggered due to excessive oxidative
stress [59,60]. The important participants in this pathway are Kelch-
like ECH-associated protein (Keap1) and antioxidant response
elements (ARE). AREs are usually sequences in the untranslated
regions of genes and act as regulatory elements for transcription.
Nuclear factor-like 2, or Nrf2 regulates as a transcription factor the
transcription of the proteins that are responsible for controlling
oxidative damage. Nrf2 is kept in the cytoplasm as a complex by
several proteins including Keap1 and Cullin3, and the average
timeframe of Nrf2 is very short when there is no oxidative stress.
Nrf2 is degraded fairly quickly through ubiquitination by Keap1 and
Cullin3 in unstressed conditions. However, if there is oxidative
stress in the cell, the Keap1-Cullin3 complex is disrupted and Nrf2
is not ubiquitinated anymore and released in the cytoplasm. Nrf2 is
then transported to the nucleus and binds to the ARE in the
promoter regions of the target antioxidative genes, resulting in the
increased transcription of these genes.

**Phytochemicals in Obesity Control**

There are a number of ways obesity can be controlled. The
first approach is lifestyle change, which includes proper diet and
exercise. Physical exercise results in diminished body fat owing
to enhanced fat burning, which is nothing but fatty acid oxidation
[61]. Exercise affects fatty acid oxidation and the rate depends on
the extent of physical exercise. For low to moderate exercise, fatty
acid oxidation is elevated, while for higher exercise, the level drops
[62]. With increasing intensity of exercise, carbohydrate oxidation
yields more energy. With enhanced duration of exercise, the energy
rate from fatty acid oxidation increases. If the exercise is continued,
breakdown of muscle glycogen is achieved. Hence, in order to
achieve maximum weight loss, duration of exercise should be more,
but moderate intensity exercise is recommended [63].

Efficient fat burning is the primary goal of treating or preventing
obesity. Bariatric surgery, intragastric balloon and metabolic
surgical procedures are sometimes done for an effective treatment.
When lifestyle modifications like more exercise and less intake of
food fail to control obesity, pharmacotherapy may help [64,65].

**Drugs for controlling obesity**

A number of drugs have been developed for controlling
obesity. The drugs commercially available are Orlistat, Lorcaserin,
Sibutramine, Rimonabant, Metformin, Exenatide and Pramlintide.
Antiobesity drugs can be broadly divided into two categories. The
first category acts centrally by repression of appetite. The second
category acts by peripheral suppression of fat absorption.
Phentermine and Sibutramine are the centrally acting drugs. The
peripherally acting drugs are Benzphetamine, Diethylpropion,
Phendimetrazine, Orlistat, Mitratapide, Dirlotapide etc. [66].
Orlistat, a pancreatic lipase inhibitor breaks down dietary fats to
free fatty acids, preventing absorption of ingested fats. However,
this formulation has certain side effects including gastrointestinal
problems like oily stool, urgency and incontinence of stool. Another
drug is Liraglutide, which is a glucagon-like peptide-1 receptor
agonist. The side effects include gastrointestinal irritation and
may lead to acute pancreatitis. Lorcaserin, a serotonin agonist has
been found to subdue appetite. A combination of Phentermine and
Topiramate facilitates weight loss by enhanced burning of energy
[67]. Phentermine is apparently a safe drug but not much effective
[68]. Some other drugs are Amfepramone, Naltrexone/bupropion
etc. [69].

There are some adverse effects of bariatric surgery, including
acute pancreatitis, deep vein thrombosis, hernia and infections
[70,71]. Often combination therapeutics are used to prevent
weight gain. Some such combinations are Contrave (Bupropion
+ Naltrexone), Empatic (Bupropion + Zonisamide), Qnexa (Topiramate + Phentermine) [72]. Most of the antiobesity drugs
have been withdrawn due to serious side effects. Fenfluramine and
dexfenfluramine, serotonin releasing agents have been withdrawn
from the market due to heart valve damage, headache, diarrhoea,
dizziness, dry mouth, insomnia, erectile dysfunction, anxiety,
irritability, lethargy, and CNS stimulation [73]. Rimonabant, an
anorectic antiobesity drug has several side effects, like increased
risk of serious psychiatric disorders including, depression,
anger, and suicidal attempts [74]. Therefore, it is apparent
that conventional obesity control measures often show various
problems, both physical and psychological. Therefore, there is an
urgency to consider some unconventional treatments.

**Phytochemicals in obesity control**

Considering the side-effects of the antiobesity drugs, a holistic
approach can be of much help in obesity control. We consume a
lot of fruits and vegetables with our daily meals, containing many beneficial chemical compounds. These plant derived molecules with therapeutic activities are coined as phytochemicals and many of them like Withania somnifera, Zingiber officinalis, Dioscorea nipponica, Malucom estica, Nelumbonucifera, Cassia nomame etc. have elicited antiobesity properties [66]. Phytochemicals can be classified into different groups. These are broadly alkaloids, carotenoids, nitrogen containing compounds and organosulfur compounds and phenolics. Phenolics consist of phenolic acids, tannins, stilbenes, coumarins and flavonoids. Flavonoids comprise anthocyanidins, flavan-3-ols, flavonols, flavones, flavanones, isoflavones. Alkaloids may be heterocyclic or non-heterocyclic [19]. Examples of other phytochemicals are glucosinolates (isothiocyanates and indoles), phytates and phytoestrogens (isoflavones and lignans).

Phytochemicals have a number of health promoting activities. They are anticardiogenic, antimutagenic, antioxidant and anti-inflammatory, besides having other beneficial attributes. Fruits, vegetables and spices are good sources of phytochemicals, dietary fibres, and they are low in energy density and impart protection against obesity [75]. The bioactive chemical compounds present in the phytochemicals have minimal energy value. They are also beneficial in weight management and obesity prevention [76, 77]. They suppress appetite, mainly by targeting gut-derived hormones, also called GLP-1 or ghrelin [78,79]. Receptors of the neuronal system, i.e. transient receptor potential cation channel subfamily V member 1 (TRPV1) is another target [80]. A constant feeling of hunger is one of the symptoms of obesity that tempts one to consume more food, and hence eventually may cause obesity. Some plant derived products may help in suppressing this nagging hunger and hence help in achieving weight loss [4]. Some of the possible rudimentary justification for obese condition may be inhibition of absorption of fat, impediment of pre-adiposyte differentiation, lipolysis stimulation and induction of apoptosis of adipocytes [63]. Oxidation of fatty acids by phytochemicals may be another reason for their beneficial effects [81,82].

Phytochemicals with antiobesity potential may be an alternate way to reduce body weight and control adipose tissue mass [63]. Not only adipose tissue growth, differentiation of pre-adipocytes and hunger or appetite may be altered by phytochemicals. All these facilitate loss of weight and hence control obesity [83,84]. Like physical exercise, phytochemical consumption has an effect on fatty acid oxidation and thus they burn fat. There may be an additive effect of phytochemicals and physical exercise on fatty acid oxidation. Apart from being good anticancer agents, isothiocyanates can prevent conversion of sugar and carbohydrates into fat. Hence cruciferous vegetables containing isothiocyanates are promising antiobesity agents [85]. The phytochemicals contained in fruits, vegetables, and other plant parts show anti-inflammatory effect [86]. This anti-inflammatory property may contribute to combat the obesogenic state [77]. One unique feature of phytochemicals is that they aim to target the pathways related to obesity [87].

The metabolic properties of an adipocyte vary during its life cycle. Upon maturation, adipocytes accumulate triglyceride and grow in size. The rate of triglyceride synthesis is more in larger adipocytes. This leads to higher rate of fatty acid flux. Secretion of cytokines is dependent on the size of adipocytes. Larger is the size of adipocytes, profile of cytokine secretion is unfavourable, hence smaller adipocytes are better. The metabolic consequences of obesity are dependent on whether expansion of adipose tissue is achieved primarily by an increase in adipocyte number or adipocyte size [88]. Phytochemicals may act on different stages of the life cycle of adipocytes. Phytochemicals elicit their function via inhibition of cell proliferation and enhancement of programme cell death of fat cells. They also inhibit the absorption of triglyceride [77], hinder the absorption of lipid, reduce the intake of energy and they help to expend more energy. Another beneficial effect is the inhibition of lipogenesis [89]. Phytochemical index is a good measure to assess the role of a particular compound in obesity control. The ratio of the energy obtained from diet and the energy expended per day is called the phytochemical index, which maintains an inverse relation with adiposity and oxidative stress [90].

A cluster of conditions that increases probability of heart disease, stroke, type 2 diabetes, hypertension, high blood sugar, pot belly (excess of body fat around waist), dyslipidaemia (abnormal cholesterol) and hyper-triglyceridemia (high triglyceride levels) is coined as metabolic syndrome (MS). Both obesity and MS have a high mortality rate. MS has increased globally and this is a serious problem, which needs urgent attention. MS and inflammation are intimately connected. Intestine, liver and adipose tissues are the sites where inflammation is initiated in MS. Due to hypertrophy of adipocytes, adipose tissues induce a pro-inflammatory state [91]. Proinflammatory cytokines like tumor necrosis factor alpha (TNF-α), IL-6, IL-8 and the chemokine ligand 2, also called monocyte chemoattractant protein 1 (MCP-1) are secreted by hypertrophic adipocytes. These proinflammatory cytokines lead to serine phosphorylation of insulin receptor substrate-1 by nuclear factor kβ (NF-kβ) and Jun N-terminal kinase (JNK) signalling [92]. Adipocyte hypertrophy leads to hypoxic condition, thereby inducing cell necrosis. This facilitates production of TNF-α, IL-6, plasminogen activator inhibitor-1 (PAI-1) and infiltration of macrophage [93]. TNF-α leads to enhancement of insulin resistance in adipose cells. Therefore, MS needs to be considered in obesity control. Treatment of MS involves lifestyle modifications, which include proper diet and regular physical activity.

Metabolic syndrome (MS) may be treated with a number of polyphenols present in food, including green tea, many types of nuts, red wine, grape seeds, berries and dark chocolate etc. Catechins in green tea is a flavanol and caffeine is an alkaloid, both show metabolic effects. The polyphenols in green tea include epicatechin (EC), epicatechin gallate (ECG), epigallocatechin (EGC), and epigallocatechin gallate (EGCG). Treatment with green tea aids in reduction of body weight, it also controls waist circumference and body fat mass [94]. Caffeine increases energy expenditure and diminishes energy intake, thereby affecting energy balance [95]. Tea shows some antiobesity effects due to the presence of catechins and caffeine, which act on the adrenergic system. The phosphodiester enzyme which facilitates hydrolysis of cyclic
adenosine monophosphate (cAMP) to AMP is inhibited by the purine alkaloid caffeine. Upon activation, cAMP signal induces adrenergic effect that includes reduction of hunger, and enhancement of energy expenditure and lipolysis [96]. The active ingredient in green tea induces overstimulation of the adrenergic system. All these lead to enhanced expenditure of energy, oxidation of fats and lipolysis [97]. It has been reported that the effect of green tea on energy expenditure is higher in the presence of caffeine [91]. Green tea lessens oxidative stress due to cardiac remodelling in patients undergoing dialysis [91,98]. Inflammation in the nervous system is also reduced by green tea [99]. Consumption of green tea results in fullness and satiety [100]. Green tea polyphenol EGCG has been found to inhibit adipogenesis. Some other natural biomolecules that also aid in inhibition of adipogenesis are genisten, resveratrol, capsacin, and procyanidins [101]. EGCG has been found to diminish hunger and increase fullness; however satiety remains unaltered after daily intake of 1796mg of the polyphenol for 3 weeks [102]. Another report says that 100mg of EGCG consumption for 2 months increased satiety and fullness, concomitantly decreasing hunger [103]. Fatty acid oxidation can be greatly increased by EGCG. These catechins also enhance the fat burning index, which is measured by the indirect calorimetry methodology and is indicated by a reduction in the respiratory exchange ratio, designated as RER [63]. The mechanism by which green tea shows its antiobesity action may be through the inhibition of catechol-O-methyltransferase (COMT), an enzyme that degrades norepinephrine. Higher concentration of norepinephrine elevates thermogenesis and fat oxidation, thereby reducing appetite [4].

The delicious Capsicum pepper is very popular and can be eaten raw or cooked. It is a genus of flowering plants in the nightshade family Solanaceae. Some members of this family are used as spices and vegetables. Its medicinal attributes are also well-known. The pungency is due to the presence of Capsaincoids, which have been reported to increase satiety [4]. A study conducted with a capsicum analog nonivamide revealed that its intake diminished the feelings of hunger for several hours [104]. The active ingredient Capsaicin in capsicum controls obesity by burning more calories via enhancement of metabolism [105]. One of the mechanisms by which capsacin promotes satiety is by enhancing gastrointestinal distress and feeling of bloating [78]. A diet rich in red pepper (cayenne) has been found to increase the concentrations of glucagon-like peptide-1 (GLP-1) and reduce the concentration of ghrelin. Glucagon like peptide 1 is produced in the ileum and colon and regulates appetite and food intake, it also acts as an anorexigenic hormone [4]. Capsaicinoids suppress appetite via interference with sympathetic nervous system (SNS). They enhance catecholamine secretion, which results in a reduction in appetite [106,107]. Both capsicum and green tea can fight obesity. Hence a combination of these two may be more effective. Keeping this in mind, trials have been undertaken to assess the efficacy of this combinational treatment in obesity control. Randomized controlled trials have reported that catechins in green tea, together with capsaicin appreciably diminished hunger and enhanced fullness and satiety [102].

Caralluma adscendens var. fimbiata is a medicinal plant that contains glycosides, flavonoids and steroids and shows a number of benefits. It acts as an anti-inflammatory, antioxidant, anti-diabetic, analgesic, anti ulcer, antibacterial and hypoglycemic molecule. This plant finds use for its antiobesity properties [108] as well as its ability to reduce hunger levels [109]. Another group of polyphenols, well-known for their antioxidant properties are Anthocyanins, which are water soluble. These are found in red wine, fruits that are red or purple in color; some vegetables like cabbage, grapes, strawberries, cranberries, cherries, apples, beets and cereals. These are good antioxidants owing to their property to transfer electrons or to donate the hydrogen atoms to free radicals from different hydroxyl groups [110]. Anthocyanins are anti-inflammatory; they are also good anti-carcinogenic, anti-convulsant and anti-diabetic. Growth of adipose tissues is negatively influenced by Anthocyanins. Lipolysis is increased, along with fat mobilisation and fatty acid oxidation [111-116]. Berries rich in anthocyanins break down fats and sugars and thus curb extra fat [117]. In vivo studies also reveal that these compounds help in reducing body weight and accumulation of adipose tissues; therefore they play a vital role in obesity control. Anthocyanins regulate fatty acid oxidation by upregulation of AMP-activated protein kinase (AMPK) and downregulation of carnitine palmitoyltranferase-1 (CPT-1) [117].

Another important polyphenol is resveratrol, which is found in abundance in red grapes, apples, peanuts, blueberries and cranberries. Resveratrol has been found to inhibit adipogenesis and preadipocytes. It does so via down regulation of adipocyte specific gene expression. Resveratrol aims at various molecular targets and decreases the number of adipocytes. This polyphenol upregulates AMPK and down regulates PPAR-α and thereby enhances oxidation of fatty acids [63]. Blueberries control triglycerides, weight gain, liver weight and abdominal fat mass. Besides, they aid in improving adipose and skeletal muscle PARP activity, which is responsible for uptake of glucose and oxidation of fats [118]. Strawberry consumption has been found to decrease total cholesterol and small low-density lipoprotein particles. Apples and blackcurrants have been found to decrease postprandial glucose and insulin via inhibition of intestinal glucose transport [91].

Another important phenolic acid is Synephrine, which is present in bitter orange. It primarily exists in protoalaloid form and is well known for its weight control properties [119]. This compound increases the resting expenditure of energy, lipolysis and fat breakdown during resting phase [63]. Thermogenic effects of this plant molecule are due to activation of β-adrenergic receptor [120]. Another fruit with a lot of health benefits is pomegranate. This fruit also contains polyphenols ellagitannins, anthocyanins etc. It has been found to decrease blood glucose and enhance insulin sensitivity among other benefits. This fruit show its anti-inflammatory effects, through the PPAR pathway. The dried and purple in color, some vegetables like cabbage, grapes, strawberries, cranberries, cherries, apples, beets and cereals.
tree with edible fruit. It is an integral part of Mediterranean diet. Olives contain anti-inflammatory agents and improve blood pressure, glycemic control, endothelial function and oxidative stress. Oil made from olive is therefore good for our health.

An important phytochemical that finds place in our food is turmeric, whose active ingredient is curcumin. It has a wide range of beneficial attributes, e.g., it has anti-inflammatory, hypoglycemic, antioxidant, antiviral and antimicrobial activities. It is also used with anticancer potential. It shows effect on signalling molecules implicated in inflammation and cancer. Activity of different transcription factors, growth factors, inflammatory cytokines, protein kinase and other enzymes are modulated by curcumin. Curcumin reduces low density lipoprotein (LDL) or the bad cholesterol and at the same time controls hypertension [121]. It has impact on weight, glucose and lipid profile, in patients with MS [122]. Another herb cardamom burns fat. Two other spices cinnamon and clove increase insulin function, reduce glucose, total cholesterol and thus help to control obesity [123,124].

There are a number of other plant derived products that aid in obesity control. Among the fruits that help to curb obesity are watermelon, avocado, apple, blueberry, cucumber, orange, quinoa, grapefruit, pear, zizyphus (Indian plum) etc. Vegetables are also equally effective in obesity related diseases. Curry leaves, cabbage, garlic, ginger, green leafy vegetables like spinach, beans, tomato, broccoli, spring vegetables like asparagus etc aid in obesity management. Allicin in garlic reduce cholesterol and unhealthy fats. Besides pulses, millets such as jowar, bajra, ragi, oats etc. are effective in obesity control Normally it is thought that intake of nuts can cause obesity since they are energy rich food. At the same time nuts contain good amount of fibre. Presence of fibres gives a sense of satiety, which plays a role in weight management. Nuts like almond are rich in vitamin E. Peanuts also show favourable lipid and lipoprotein profiles that help in weight control [105]. Antioxidant lycopene rich tomatoes are studied with a number of health benefits. These include less risk of heart disease and cancer. Besides, tomato contains vitamin C, folate, potassium and vitamin K and helps to prevent obesity [125].

Cooking without brushing the pan with oil is practically impossible. Oils and fats give a good texture and flavor to the skin, but a diet containing high amount of unhealthy fats can cause obesity. However, there are some oils that can combat obesity, including coconut oil, mustard oil and olive oil. Mustard oil extracted from the seeds of mustard is rich in fatty acid, oleic acid, erucic acid and linoleic acid and contains low amount of saturated fat. Mustard oil also contains Diacetylglycerol, which is responsible for its anti-obesogenic effect [126]. Coconut oil is used for preparing food and contains more than 65% medium-chain fats, due to which thermogenesis can be achieved, leading to weight loss. Coconut oil increases the high-density lipoprotein (HDL) level, thereby improving the LDL/HDL ratio. Therefore, it has an impact on BMI [127]. Olive oil is good for cooking and it is also used as a good salad dressing. Medium chain fatty acids present in it results in thermogenesis and is known for its antiobesity attributes [128].

Tea seed oil contains monounsaturated fatty acids and can prevent obesity [129]. Antiobesity effects are mediated through their anti-lipase activity, anti-hyperlipidemia etc. These oils inhibit adipogenic transcription factors including PPARy and CEBPα at the protein and mRNA level. Plasma glycerol concentration is increased, which is a marker of lipolysis and intracellular triglyceride and fat accumulation is also diminished by the essential oils [130]. Herbs and spices are often effective in obesity control. They inhibit lipase, down-regulate adipogenesis and thermogenesis. They also affect lipid metabolism and regulate a number of signal transduction pathways implicated in obesity. Antiobesity herbs include Acacia arabica, Aconitum heterophyllum, Aloe vera, Azadirachta indica, Betula utilis, Calatropis gigantean, Cinnamomum zeylanicum, Emblica officinalis, Moringa oleifera and many others [89].

Some other vegetables that we consume have promising potential. Spinach is a leafy vegetable, rich in chlorophyll, antioxidant phyto-constituents such as flavonoids, polyphenols, carotenoids and vitamins. It is very effective in controlling obesity caused due to high fat diets. It aids in reduction of water retention in our body [131]. Beans being low in fat and abundant in fiber increase satiety, but do not add much calorie to a diet [132]. Green coffee bean has been found to regulate body fat in high fat diet induced obesity. These beans have been found to reduce weight, fat mass, size of adipocytes and plasma lipid level. These have been achieved via regulation of adipogenesis and lipogenesis [133]. Two other phytochemicals asparagus and Jerusalem artichokes, rich in dietary fibres are capable of fighting obesity as well as getting rid of toxins and other wastes from the body [134].

Dietary fiber or roughage are found in edible plant foods such as cereals, fruits, vegetables, dried peas, nuts, lentils and grains. These fibers may be in soluble or insoluble forms, of which insoluble forms work better in controlling obesity than the soluble ones [135]. As they remain undigested, they help to relieve constipation and they control hunger by boosting up metabolism. Oats contain good antioxidants and minerals. The fibres in oats bring down cholesterol levels and control obesity. Millets also contain fibres, which absorb cholesterol, enhance bile secretion, emulsify fats and thereby reduce weight [136]. Consumption of dietary fibers is important in preventing MS and obesity. Some fruits like apples, and avocados contain high amount of fibre and other nutrients. Regular consumption of apples, avocados increases satiety, hence appetite is decreased and so is obesity [105]. Apart from being rich in fiber, avocados contain high amount of monounsaturated fatty acid (MUFA), and are one of the highest sources of beta-sitosterol, which inhibits cholesterol [137].

As mentioned before, exercise is one of the key ways to achieve weight loss. Exercise, along with generous consumption of phytochemical may show some add-on effect on fatty acid oxidation. Consumption of Catechins from green tea along with exercise showed an additive effect on fatty acid oxidation rate and lipolysis [138]. Similar findings have been reported in case of p-synephrine and exercise. However, anthocyanins along with exercise did not show any promising outcome, only an accumulative effect
was seen [116]. For many phytochemicals, bioavailability is a big problem, and that could be one of the reasons why additive effects were not seen in case of anthocyanins and exercise. Nature of the phytochemicals and their mode of action on different molecular targets may be some other determining factors for their efficacy. EGCG along with exercise increases fat oxidation rate, fat utilization, plasma glycerol and epinephrine, while respiratory exchange ratio (RER) gets diminished. Resveratrol along with exercise enhances fat oxidation and decreases RER. P-synephrine increases oxidation of fat, when taken in combination with exercise. All these plant molecules have different modes of action. Combination of more than one phytochemical in aid of obesity control has been reported. Resveratrol in conjunction with genistein and quercetin caused a marked decrease in the process of adipogenesis. Resveratrol in cooperation with vitamin D has been found to prevent extra weight gain [139]. Thus, resveratrol along with other phytochemicals or physical exercise may serve as a good candidate in obesity control.

**Phytochemicals in obesity related cancers**

Phenolic phytochemicals have strong antioxidant activities due to the phenolic hydroxyl groups providing hydrogen atoms for capturing reactive oxygen species (ROS). Excessive oxidative stress is mainly due to H2O2 and OH and these agents are responsible for cancer cell proliferation. Phenolic phytochemicals can inhibit the effects of H2O2 by intervening in several pathways including NF-κB and MAPK. Epigallocatechin gallate (EGCG) is a phenolic phytochemical present in green tea and has very effective anti-oxidant effects. The effect of EGCG on human epidermal keratinocytes has been observed by inducing oxidative stress through ultraviolet radiation, as UV radiation increases the concentration of H2O2. This in turn increases the phosphorylation of JNK and MAPK. It was observed that EGCG could inhibit MAPK activation by scavenging of H2O2 [140]. TNF triggers x activation. In another study Manna et al. [141] showed that resveratrol, the phenolic phytochemical found in grapes, prevented NF-κB activation in several cell lines, including Jurkat, HeLa and U-937. Yu et al. [142] showed similar results in HeLa cell lines. Isothiocyanates also have antioxidant properties as well as peroxidase properties [143]. There are many other natural phytochemicals, that act as good antioxidants and help in obesity related cancers. Lycopene, a carotenoid is found in fruits and vegetables having red color, particularly tomatoes. Phloretin, a flavonoid is found in apples. Another flavonoid of importance is kaempferol, found in kale, beans, spinach and broccoli etc. All these three contribute as anticancer, anti-diabetic and anti-obesogenic agents [144]. Quercetin, a flavonoid is found in onions, green tea, apples, berries, St. John’s wort, buckwheat tea etc. Quercetin is an antioxidant that aids in obesity and cancer. Some other antioxidant phytochemicals that contribute in this regard are naringenin, a flavonoid, flavones like luteolin, daidzein, genistein etc. Allicin, an organosulfur compound, phenolic acids like caffeic acid, chlorogenic acid, ferulic acid, tannins are good anti-obesogenic and anti-cancer agents [144].

ROS impacts activation of transcription factors like Activator Protein-1 (AP-1) or NF-κB. These in turn can modulate proinflammatory cytokines such as Tumor Necrosis Factor α (TNF-α), Interleukins like IL-6, IL-8, and IL-1 [145]. Glicosides of apigenin, leuteolin are anti-inflammatory and can work in this regard. Apigenin is capable of suppressing nitric oxide (NO) via inhibition of inducible nitric oxide synthase (iNOS) and COX-2 [146]. Luteolin is capable of inhibiting inflammation of adipocytes. Inflammation in macrophages and phosphorylation of JNK in macrophages can be inhibited by luteolin. Quercetin, found in abundance in apples, isoflavones like genistein, daidzein, glycitein show their anti-inflammatory properties via modulation of NF-κB. These compounds have been found to downregulate TNF-α and IL-6. Anthocyanins are good anti-inflammatory molecules that works through mitogen activated protein kinase (MAPK) pathway. Glicosides of malvidin, delphinidin, cyanidin, petunidin, and peonidin can inhibit IL-1β -activation of NF-κB [147]. Licorice roots contain triterpenes like glycyrrhizin and glycyrrhetic acid, which are also promising anti-inflammatory agents. They act through P13K/Akt/GSK3-β pathway to reduce cytokine production [146].

Polyphenols like curcumin, genistein and epigallocatechin are particularly effective in this respect for mitigating the effects of obesity related tumor onset and progression. Curcumin is a lipophilic polyphenol and a popular Indian spice. The effect of curcumin on the Nrf2 pathway has been studied [143,148]. Das and Vinayak [148] studied the effect of Nrf2 activation on liver cells of mice by inducing Dalton’s lymphoma. When lymphoma was progressing, the Nrf2 pathway was suppressed, but was restored after administering curcumin through intraperitoneal injections. Chen et al. [149] showed that curcumin had dual effects of increasing the expression of Nrf2 as well as decrease in the expression of Fen1, which is known to be expressed in breast cancer as an endonuclease that cleaves the phosphodiester bonds of DNA. Curcumin has been found to inhibit TNF-α, IL-1β and suppress NF-κB [150,151]. Resveratrol has been found to inhibit all three, in addition this biomolecule also inhibits IL-6 [146]. Rosmarinic acid, from Rosemary and sage are good antioxidants that has effects on these pathways [152].

Plant biomolecules induce the expression of several antioxidant enzymes. They also aid as modifier of a number of signal transduction pathways. This includes suppression of stress induced proteins and Keap1 dissociation from Nrf2 as a consequence of stress is facilitated. EGCG, lycopene, resveratrol, curcumin, mulberry leaf help in this regard by modulation of Nrf2/ARE pathway [153]. Echinacea purpurea L, a herbal medicine improves cellular oxidative status and promotes anti-inflammatory activities [154]. As mentioned before, PI3K pathway is one of the most activated pathways in cancer. There is evidence that PI3K/ AKT pathway leads to obesity and type 2 diabetes mellitus [155]. Therefore, it is important to see how and which phytochemicals target this pathway. A wide range of phytochemicals have been found to inhibit PI3K/AKT pathway. These include gingerol, curcumin, dialyl trisulfide, emodin, ginsenoside, plumbagin, apigenin, resveratrol, EGCG, quercetin, fisetin, luteolin, apigenin, indole-3-carbinol, sulforaphane, PEITC, eLLag acid and many more [156]. Another promising therapeutic target in cancer is MAPK. It has been observed that obesity is related to increased level of...
Mitogen-activated Protein Kinase Phosphatase 3 (MKP-3). A direct correlation between diminished level of MKP-3 and adiposity has been observed [157]. A link has been explored between MAPK signalling and endometrial cancer, which is again due to obesity [158]. Among the phytochemicals that target MAPK are soy-leaf, isoflavones, resveratrol, caffeic acid, apigenin, quercetin, kaempferol, genistein, triterpenoids [159]. Besides, INK, phosphor ERK, phosphor AKT are also down regulated.

Two signalling pathways that are highly implicated in solid tumors are Ras-Raf-MEK-ERK and PI3/AKT. Raf kinase is the downstream target for Ras. This facilitates the activation of ERK1/2. Ras/mitogen-activated protein kinase (MAPK) pathway is linked with adipogenicity. Once ERK is activated, it translates to the nucleus and initiates mitotic clonal expansion in preadipocytes, which is important for adipogenesis [160]. Acacetin is a natural flavonoid which is derived from Robinia pseudoacacia. Acacetin has been studied as its antioxidant and anticancer properties. Jung et al. [161] reviewed its anti-cancer potential and noted its ability to modulate the PI3K-Akt-mTor pathway. Chen et al. [162] have shown the modulating effect of black rice anthocyanins on the Ras-Raf-MEK-MAPK pathway. Curcumin, diallyl trisulfide, resveratrol, apigenin, sulfoparane, etc. help to modulate this pathway [163]. MAPK/ERK pathway is being modulated by sibilenin, fisetin, genistein, etc. [164].

Conclusion

Obesity is one of the most endemic public health problems worldwide, and a risk factor for many diseases including cancer. Controlling obesity through medication and surgical procedures is often fraught with severe side effects. Hence controlling obesity through naturally occurring phytochemicals is one of most effective strategies. We have examined the role of obesity as risk factors for many diseases in this review. We have examined the genes, proteins and genetic pathways that are triggered by obesity and subsequently affect the onset and progression of many diseases. We have also provided a broad review of the phytochemicals that are effective in controlling obesity by acting on the related genes and pathways, with a particular focus on their effects on different types of cancers caused by obesity.

Conflict of Interest

The authors have no conflict of interest.

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