**Bioactive components of tea**

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**Abstract**

Tea (Black tea and Green tea) are one of the most widely consumed beverages in the world. However, with the increasing interest in the health properties of tea and a significant rise in scientific investigation, this review covers some of the recent findings on the health benefits of both green and black tea. The mechanisms of action of various black and green tea components have been presented. Green tea contains a unique set of catechins that responsible for its biological activity potentially relevant to the prevention of diseases. Although there has been much focus on the biological property of the major tea catechins, black tea offers major health benefits either due to the presence of the catechins in epimerized form or some other active components of both varieties of tea. Characteristics unrelated to the antioxidant properties of green and black tea might also be responsible for their therapeutic potential in preventing diseases. Synergistic effect of the tea constituents is increasingly recognized as being potentially important to the medicinal benefits of black and green tea. The studies indicate that tea has the potential of being a part of diet for healthy living.

**Introduction**

Tea and infusion of the leaves of the *Camellia sinensis*, is a widely consumed aromatic beverage. Consumers vary in their preferences on the type of tea they consume, which in turn is dependent on the degree of fermentation, taste and color [1]. Tea is considered as a rejuvenator and often acts as therapeutic adjuvant for several ailments for people from all walks of life [2]. Various types of tea are popular in different regions of the world. Black tea (Bt) is consumed as beverage principally in India, Pakistan, Sri Lanka, Russia, Europe, North America, North Africa etc. and most of the processing of Bt involves intense crushing and fermentation process [3]. Green tea (Gt) is a lightly processed tea that is not fermented at all [4] and is widely consumed in China and Japan. Tea has been considered as home medicine since time immemorial. It is one of the plant products with highest total flavonoid content and these compounds are responsible for the distinctive taste and color of tea and also the health benefits associated with consumption of tea [5-20]. Polyphenols in tea are secondary metabolites of plants and are generally involved in defense against ultraviolet radiation or aggression by pathogens. Polyphenols may be classified into different groups as a function of the number of phenol rings that they contain and on the basis of structural elements that bind another [21]. The main classes of polyphenols found in tea include phenolic acids, flavonoids, and lignans. Gt and Bt differ in their chemistry, only because of their processing methods but not in leaves itself. The bioactive components in both type of tea has been summarized in table 1. Thus, which type of tea is superior in terms of health benefits, is a question of much interest. Therefore, this review was aimed at evaluating the health benefits of both types of tea with respect to their mechanism of pharmacological action.

**The Key players – Flavanols**

Reactive oxygen species (ROS) and free radicals can cause severe damage to the normal cells of the body. These damages can be to the DNA, proteins, and other biological macromolecules, thereby causing pathological changes in the cellular environment leading to a wide variety of chronic diseases. There are numerous studies that reveal that these diseases are mediated by oxidative stress and imbalance between prooxidant and antioxidant factors. Antioxidants play a pivotal role in preventing or slowing the progression of these conditions. In the last decade, there has been much interest in the potential health benefits of tea polyphenols as antioxidant. Epidemiological studies and associated meta-analyses strongly suggest that long term consumption of diets rich in tea polyphenols offer protection against development of cancers, cardiovascular diseases, diabetes, osteoporosis
and neurodegenerative & ocular diseases [22,23]. The phenolic groups in bioactive components of tea can donate an electron to form relatively stable phenoxyl radicals, thereby disrupting the cascade of oxidation reactions in cellular components. The most common bioactive compound found in tea are flavonoids like flavan-3-ols (flavanols or flavans), which are present in relatively large amounts in tea compared to their levels in other foods. The flavan-3-ols create the chemical signature pattern that is distinctive in each type of tea. The flavan-3-ol sub-classes are ranked by degree of polymerization [5,6,8-10,24]. The catechins are monomers (catechin, epicatechin, epicatechin gallate, epigallocatechin, and epigallocatechin gallate), the Tfs are dimers (theaflavin, thearubigin, Theaflavin 3-O-gallate, Theaflavin 3,3′-digallate, and epigallocatechin gallate, Theaflavan 3,3′-digallate, Theaflavan 3-O-gallate), and Tr are oligomers [25]. Gt is rich in monomeric catechins but lack flavanols in dimeric and oligomeric form.

### Flavonoids

Flavonoids found in Gt and Bt include flavonols like quercetin, kaempferol, myricetin [26]. Flavonoids are indispensable components in variety of nutraceuticals because of their anti-inflammatory, anti-oxidative properties coupled with their capacity to modulate various cell signaling. These flavonoids possess numerous health benefits and further research on their therapeutic potential needs to be explored. Unlike flavanoids, flavonoids are usually present in tea as glycosides.

### Antioxidant property

Antioxidant property of Gt lies in the monomeric catechins either with or without galloyl moiety. Although Bt contains much lower concentrations of monomeric catechins than Gt, it is generally believed that dimeric or oligomeric catechins contribute greatly to antioxidant action of Bt [27]. Consumption of catechins by humans either in its monomeric form or after its epimerization results in modest transient increase in the total antioxidant capacity of plasma, measured through ferric-reducing antioxidant potential (FRAP), oxygen radical absorbance capacity (ORAC), or Trolox-equivalent antioxidant capacity (TEAC) assays [28-30]. The studies comparing the antioxidant activities of dimeric and monomeric catechin had revealed that all catechins and Tf inhibited Cu²⁺-mediated LDL oxidation in the order: TF₃ > ECG > EGCG > TF₂B > TF₂A > TF₁ ≥ EC > EGC [31-34]. Examination of the scavenging effects of tea catechins and their glycosides on free radicals had revealed that the presence of at least an ortho-di-hydroxyl group in the B ring and a galloyl moiety at the 3 position was important in maintaining the effectiveness of the radical scavenging ability [35,36]. Yoshino, et al. [37] had also showed that Gt and Bt infusions have almost similar antioxidant activities in rat liver homogenate. The most important catechin oxidation products in Bt are TF and its mono and digallates [38]. Tfs possesses a characteristic benzoazepinone moiety, which is produced by condensation between a catechol-type B-ring of EC and a pyrogallol-type B-ring of EGC [39]. Tfs, are formed via the co-oxidation of pairs of epimerized catechins, one with a vic-trihydroxyphenyl moiety, and the other with an ortho-dihydroxyphenyl structure. Apart from epimerized catechins are four main Tf derivatives that reserve two Arings, two C-rings from their precursors and possess a characteristic element of the fused seven-member benzoazepinone ring [40]. The benzoazepinone moiety of Tfs play an important role in affording antioxidant protection for the preferred oxidation site for electron donation because of the existence of resonance forms [41]. TF radicals have a higher reduction potential than the tea catechin EGCG. Tfs have significantly higher reaction rates with superoxide radicals than EGCG [42,43]. It has been suggested that the existence of resonance form in the benzoazepinone moiety might be responsible for electron donation [40,41] and it might play an important role in affording antioxidant protection for the preferred oxidation site in the oxidant models of 2,2-diphenyl-1- picrylhydrazyl (DPPH) and hydrogen peroxide. Another important aspect of polyphenols is that the gallated catechins have lower gap energies between the HOMO and LUMO orbitals than those without galloyl moiety. Thus the carbon atoms of galloyl moiety has higher susceptibility toward nucleophilic attack. The antioxidant activity of Tfs were more effective than glutathione (GSH), L(+)-ascorbic (AsA), dN- tocopherol, butylated hydroxytoluene (BHT), and butyl hydroxyanisole (BHA) in in-vitro peroxidation of rat liver homogenate induced by tert-butyl hydroperoxide (BHP) [42]. Both Bt and Gt have monomeric gallated catechins of the flavanol class and EGCG is the precursor of TF [43] which has the most positive effect in scavenging free radicals [44]. The effectiveness of TF was increased by esterification with gallic acid [45]. The higher the number of phenyl hydroxyl groups in Tf derivatives the more

### Table 1: Major bioactive components of tea.

<table>
<thead>
<tr>
<th>Sl No</th>
<th>Source (Tea)</th>
<th>Form</th>
<th>Compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Green tea and Black tea</td>
<td>Flavonol (Monomer)</td>
<td>Catechin, Epicatechin, Epicatechin gallate, Epigallocatechin and Epigallocatechin gallate</td>
</tr>
<tr>
<td>2</td>
<td>Black tea</td>
<td>Flavanol (Dimer)</td>
<td>Theaflavin, Theaflavin 3-O-gallate, Theaflavin 3,3′-digallate</td>
</tr>
<tr>
<td>3</td>
<td>Black tea</td>
<td>Flavanol (Oligomer)</td>
<td>Theaflavins</td>
</tr>
<tr>
<td>4</td>
<td>Green tea and Black tea</td>
<td>Other flavonoid</td>
<td>Quercetin, Kaempferol, Myricetin</td>
</tr>
<tr>
<td>5</td>
<td>Green tea and Black tea</td>
<td>Amino acid</td>
<td>L-theanine, Glutamine Arginine</td>
</tr>
<tr>
<td>6</td>
<td>Green tea and Black tea</td>
<td>Methyl xanthine</td>
<td>Caffeine</td>
</tr>
<tr>
<td>7</td>
<td>Green tea and Black tea</td>
<td>Phytoestrogens</td>
<td>Lignans</td>
</tr>
<tr>
<td>8</td>
<td>Green tea and Black tea</td>
<td>Polysaccharides</td>
<td>Galactose, Arabinose, Rhamnose, Xylose, Galacturonic acid, Mannose, Ribose and Glucuronic acid</td>
</tr>
<tr>
<td>9</td>
<td>Green tea and Black tea</td>
<td>Trace minerals</td>
<td>Copper, Manganese, Iron and Zinc</td>
</tr>
</tbody>
</table>

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it can interact with ROS and this and benzoletropolone moiety might have enhanced the scavenging of the O$_2^·$, H$_2$O$_2$ and OH in in vitro models [43]. The relation of the amount of metal ions in living cells to the oxidation of lipid, the ability to chelate metal ions with Tf are also important while evaluating its health benefits [46]. Catechins in monomeric or dimeric form can prevent the peroxidation of lipid effectively by cutting off the chain reaction in oxidation of lipid [45].

Trs are known to be heterogeneous polymers of flavan-3-ols and flavan-3-ol gallates and their bonds are presumably present at C-4, C-6, C-8, C-2′, C-5′, and C-6′ in the flavan-3-ol unit [47]. The MALDI-TOF results from a more recent study reveal that molecular weight of Trs are not over 2100 Da and these oligomers of catechins in which the 3-OH group is more and less esterified by gallic acid [48]. Tfs and Trs like their major precursor catechins are capable to eliminate free radicals, which is mainly due to the contribution of the residue of the active phenolic hydroxyl groups and benzoletropolone groups at the Tr molecule [42].

Catechin in dimerized form, prevents DNA damage by suppressing oxidative stress and inhibiting cytochrome P450 1A1 and other oxidant enzymes under in vitro conditions [49]. Tf ameliorates cerebral injury through anti-inflammatory effects and modulation of signal transducer and activator of transcription (STAT)-1 [50]. Tf targets miRNA-128-3p and leading to the activation of Nrf2 pathway thereby reducing the oxidative stress [51].

Increased expression of inducible nitric oxide synthase (iNOS) and subsequent production of large amounts of nitric oxide results in its reaction with superoxide to form peroxynitrite, and other NO-derived oxidants capable of damaging DNA, proteins and contributes to vascular failure and end-organ damage during endotoxemia and to diseases such as asthma, short- and long-term lung disease, septic shock and other diseases [52], ulcerative colitis, and Crohn’s disease [53]. Infusions prepared from Gt, Bt and individual tea polyphenols can suppress iNOS gene expression and iNOS activity in cultured macrophages [54,55]. Pharmacological suppression of iNOS-dependent NO production might be helpful in the treatment and prevention of chronic diseases [56].

Tea polyphenols can also inhibit the formation of ROS by inhibiting the enzyme, xanthine oxidase which catalyzes the oxidation of both hypoxanthine and xanthine to uric acid, while reducing O$_2$ to O$_2^·$ and H$_2$O$_2$ [57]. Apart from Monomeric and dimeric catechins Quercetin found in both Bt and Gt have strong ability to sequester free radicals and bind transition metal ions [58]. Beyond antioxidant activity: as modulators of cell signaling. Despite significant advances in our understanding of the biology of tea polyphenols, they are still mistakenly regarded simply as antioxidants. The evidences suggest that their beneficial effects involve decreases in oxidative/inflammatory stress signaling, increases in protective signaling and neuro hormetic effects leading to the expression of genes that encode antioxidant enzymes, phase-2 enzymes, neurotrophic factors, and cytoprotective proteins [59,60]. Specific examples of such pathways include the sirtuin- FoxO pathway, the NF-κB pathway, and the Nrf-2/ARE pathway [61].

The mechanism(s) of the inhibitory effects of Gt or Bt on carcinogenesis are related to inhibition of NAPDHcytochrome P450 reductase; inhibition of N-O-acetyltransferase; induction of CYPIA2 and UDP-glucuronosyl transferase (leading to increased metabolism of IQ and rapid elimination of detoxification products in the urine); and electrophile scavenging/degradation [62]. The antioxidant properties of tea also might be important during the post initiation phase of carcinogenesis [63,64].

Bt and Gt can help fight cavities and prevent gum disease [65]. Tea polyphenols suppressed the growth of cavity-causing bacteria in plaque and reduced acid production levels [66]. Tea polyphenols also inhibited an enzyme produced by the bacteria, glucosyltransferase, thus preventing the formation of the matrix material by which dental plaque adheres to tooth surfaces [67]. In addition, the size and stickiness of dental plaque were reduced because certain bacteria, when exposed to black tea, lost their ability to form aggregates with other bacteria [65,68].

Monomeric catechins and flavonoids, found in Gt can maintain stable mitochondrial membrane potential (ΔΨm) of target cells and decrease the lens epithelial cell death rate under high glucose [69]. Tea polyphenol containing ophthalmic gel could maintain stable ΔΨm, reduce the generation of ROS, and prevent lenticular epithelial cells from apoptosis in experimental model and possesses great potential in clinical practice. IgE and FceRI are essential in the development of allergic diseases. Studies revealed that tea polyphenol could suppress the expression on the Fc epsilon receptor I (FceRI) by highly associating with plasma membrane microdomains and lipid. Studies confirmed that EGCG attenuated the mucus production and the MAPK expression in an asthma model-mouse [70].

Ocular hypersensitivity including contact dermatitis and corneal allograft rejection are major type IV allergies where new formulations are required. It has been found that Tf is involved in inhibition of the alterations of cytokines and maintenance of antioxidant status in animals with type IV allergies [71]. Further studies in this direction is required for commercial preparation of drugs. Flavonoid component has a ability in inhibiting Lyn/PLCγ/IP3R-Ca$^{2+}$, Lyn/ERK1/2, and Lyn/NF-κB signaling and there by dampening the inflammatory response during allergic conjunctivitis [72].

Many beneficial physiological and pharmacological effects have been ascribed to tea consumption is at least in part, related to the inhibition of gelatinases involved in both tumor
invasion and neo-angiogenesis (the growth of new blood vessels into a tumor) [73]. Some aggressive bacterial strains produce high amounts of gelatinases that contribute to their pathogenesis green tea polyphenols have the ability to inhibit bacterial gelatinases [74]. Some important flavonoid found in tea improves retinal ganglion cell survival and function in glaucomatous neurodegeneration however more studies in this area is required [75].

A prodrug of tea polyphenol alleviated the hypoxia-inducible factor-1α (HIF-1α)/vascular endothelial growth factor (VEGF)/VEGF receptor 2 (VEGFR2) pathway that contributes to the pathogenesis of choroidal neovascularization [76,77]. However, studies with Bt polyphenols are lacking.

Reports suggest, tea compounds mimic the effect of insulin signaling to FOXO1a and PEPCK, key downstream effectors of cellular insulin/longevity signaling [78]. Bt and Gt polyphenols also suppress certain genes in the liver that are responsible for glucose metabolism [79]. Bioactive components found in tea including kaempferol, myricetin, quercetin, ECG, Tf3, Tf2B, Tf2A have been reported as anti-HIV-1 compounds. Their high biological activity and low toxicity tea polyphenolics represent a valuable natural source of molecules that might be favorable to HIV patients due to enhanced lymphocyte proliferation, which could restore disturbances in T-cell homeostasis [80]. Molecular docking and molecular dynamic simulation studies have revealed that Tf3, is one of the most potent gp41 inhibitor [81]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82]. A common conclusion drawn from scientific evidence is conversion of monomeric catechins during fermentation [82].

Reports suggest that tea contains the highest quantity of theanine present in tea is dependent upon the cultivation process, growing conditions and type of tea [83]. Recent studies have shown that topical administration of chitinase inhibitors, caffeine and dexamethasone in combination produced a remarkable reduction of inflammatory signs. Caffeine has more anti-inflammatory property than the chitinase inhibitors and significantly reduces acidic mammalian chitinase activity in tears [78].

**Amino acids:** In Gt and Bt, L-theanine is one of the most important amino acid and it accounts for almost 50% of the total free amino acids found in tea. It was first discovered in an aqueous extract of Gt, later it was reported in Bt [84]. Reports suggest that theanine is synthesized in the tea roots and then proceed to the developing shoot tips under the catalytic activity of a specific enzyme, L-glutamate ethylamine ligase and using the amino acid alanine as the precursor of ethyl-amide in presence of light [85]. The chemical structure of L-theanine is gamma-ethyl amide of glutamic acid. The amount of theanine present in tea is dependent upon the cultivation process, growing conditions and type of tea [86]. Reports suggest that black tea contains the highest quantity of theanine. Darjeeling black tea contains the highest quantity of theanine (250 mg/100 gms). Although caffeine can have certain negative effects on anxiety disorders, studies reveal that L-theanine neutralizes negative effects of caffeine without reducing its mind-energizing, fat-burning features [87,88].

Amino acids other than theanine found in Gt and Bt include conditionally essential amino acids glutamine and arginine. Under stress conditions, these amino acids become essential and it has to be taken as food or supplements. Gt and Bt also contains amino acids like asparagine, and serine.

**Lignans:** Gt and Bt contain lignans, which are compounds that form the building blocks of plant cell walls. Lignans are phytoestrogens that help regulate the body's estrogen production. Lignans are converted by intestinal bacteria into the enterolignans, enterodiol and enterolactone. Enterolactone, the primary lignan metabolite circulates in

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**Figure 1:** Health benefits of bioactive components of tea.
the blood & produces weak estrogenic activity and possess several biological activities. Studies have shown enterolactone helps to reduce risk of breast, prostate and colon cancers, and cardiovascular diseases. Moreover, lignans can support healthy weight and glucose metabolism, reducing the risk of insulin resistance, metabolic syndrome and diabetes. Although observations that drinking tea, appears to be protective against osteoporosis in older women [62,89]. This necessitates further investigation.

**Tea polysaccharides:** Tea polysaccharides (soluble fibers) were found to be mostly glycoconjugates in which a protein carries one or more carbohydrate chains covalently attached to a polypeptide backbone, usually via N- or O- linkages [90]. Reports suggest polysaccharides present in Gt and Bt blunt the spike in blood sugar level after meal and retard the absorption of glucose and may thus be beneficial to people with diabetes [91-93]. Interestingly, the polysaccharides in Bt also seemed to possess the strongest ability to mop up free radicals, which are believed to play a role in the onset of numerous cancers. The Bt was found to contain lower molecular weight polysaccharides (3.8 - 32.7 kDa) than Gt 9.2 - 251.5 kDa. Studies show that the Bt polysaccharides showed a dose-dependent effect on α-glucosidase inhibitory activity [94]. α-glucosidase inhibitors are often considered as oral anti-diabetic drugs that prevents the breakdown of carbohydrates. Studies suggest that Gt polysaccharides hardly inhibited α-glucosidase activity under similar conditions [94]. A mathematical analysis of data from 50 Countries reveal that the prevalence of Type 2 diabetes is low in countries where people consume Bt regularly [95]. The scavenging rates of Gt polysaccharides and Bt polysaccharides on DPPH radicals were 47.9%, and 61.7%, respectively [95]. Deoxyribose assay, the photo-reduction of NBT assay and the lipid peroxidation inhibition studies indicated Tea Polysaccharide of molecular weight 26.8 kD, 11.8 kD, 4.2 kD exhibited the highest antioxidant activities [96]. While detailed studies on cause- bioavailability - effect relationship between Gt and Bt drinking and various diseases are yet to be confirmed, the scientific findings are consistent with a number of biological, physiological, epidemiological and clinical studies suggesting that Gt and Bt have a positive effect on human health.

**Trace elements:** Tea (Bt and Gt) is rich in essential trace elements, which might play an important role in ionic homeostasis in the physiological system. Among the minerals and essential trace elements, Ca, Na, K, Mg, and Mn are present in tea leaves at g/kg level, while Cr, Fe, Co, Ni, Cu, Zn are present at mg/kg level. Reports suggest that the presence of trace elements in green tea is lower than that of black tea. Besides essential macro- and microelements, *Camellia sinensis* strongly accumulate aluminum. Aluminum in most of its forms has no harmful effect on living organisms. However, under certain conditions aluminum might form toxic species [97]. However, these studies appear to have major limitation and may need more detailed investigations.

**Bioavailability:** The worldwide popularity of tea and the absence of toxicity as a natural dietary agent, has made tea an excellent candidate for dietary prevention of chronic diseases [98]. Results from various studies show that the bioavailability of many important tea polyphenolics is low. The poor bioavailability of EGCG, theaflavins, and thearubigins may be explained by Lipinski’s Rule of 5, which is based on the ability of a molecule to pass through transient pores formed in the plasma membrane by the movement of the phospholipid acyl tails and also a molecule’s ability to form hydrogen bonds. Many of the mechanistic studies of the tea polyphenolics have been conducted on cell lines. But it is unlikely that the concentration of polyphenol used in these experiments can be obtained in target tissues other than the skin and GI tract under *in vivo* conditions. So, correlating mechanistic data observed *in vitro* with that found *in vivo* should be done with careful consideration of the poor bioavailability of the tea polyphenols.

Glucuronidation, sulfation, and methylation represent the major metabolic pathways for tea polyphenolics [99]. There are species and tissue-specific differences in tea polyphenolics glucuronidation, with humans and mice being more similar than humans and rats [99,100]. Methylated catechins have been observed in the rat bile of the rat following oral EGCG administration. In humans, EGC is detected mainly as the glucuronidation form or sulfated form with only a small amount present as the free form [100,101]. Methylation of EGC also occurs in humans leading to the formation of glucoronicide or sulfite conjugate. The sulfated form of E is more abundant than the glucuronidation form [101] while EGCG is present mainly in the free form in the plasma [102]. In addition to these conjugation reactions, the tea catechins undergo metabolism in the gut to form the ring fission products gamma valerolactone [103] which may be further broken down by gut flora to phenylacetic and phenylpropionic acids.

Recent studies reveal that polyphenols interact with model membranes in a structure-dependent manner. Liposomes showed morphological changes with higher percentage in the presence of Tf > Tf > EGCG > GC corresponding to the number of phenolic gallate groups and hydroxyl groups in polyphenols influencing how the polyphenols interact with the membrane, following initial H-bond driven interaction. These polyphenols mainly affected hydrophilic region of lipid bilayers [104].

Decaffeinated tea supplements provide large doses of polyphenols without the unwanted side effects of caffeine. Studies reveal polyphenols administered in the form of solid tea showed enhanced bioavailability compared with that of Gt or Bt. Administration of Gt and Bt solids led to significant increase in antioxidant capacity [105]. Flavanol metabolite formation may have contributed to the antioxidant effect and this issue deserves further investigation. However, efforts to translate the results of *in vitro* and animal studies to human
interventions have met with limited success [106]. This may be due to the fact that during in vitro studies, the cells are exposed to supra-physiologic concentrations of tea catechins and there is limited information on the bioavailability of tea flavanols post consumption of tea brew [107,108]. Use of nanotechnology, improves the bioavailability of tea bioactive components and consequently enhance the bioactivity [109].

**Conclusion**

It is also likely that tea components by their ability to control intracellular signaling cascades that target multiple signal transduction pathways, are involved in the inhibition of various chronic diseases. Among the numerous mechanisms proposed, it is also important to further investigate, which are the primary and subsequent events. The relative importance of these pathways needs to be determined under in vivo conditions. The flavonols and flavonoids present in Gt and Bt interact synergistically in their disease modifying actions. However, the absorption rate of black tea polyphenols is higher than that of green tea polyphenols. Novel strategies to enhance the absorption of tea polyphenols and subsequently improve their bioavailability, should be probed. However, it remains the healthiest beverage amongst all.

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