Chapter

Remedial Effects of Tea and Its Phytoconstituents on Central Nervous System

Manisha Singh, Vandana Tyagi and Shriya Agarwal

Abstract

Tea in all its forms is one of the commonly consumed beverages globally, after water. Apart from just being a beverage, it also has extensive therapeutic values. The phytoconstituents of tea either in their pure form or as an extract are essential part of traditional as well as modern day medicines. Tea has shown its medicinal benefits in treating, improving and preventing many of the ailments ranging from being potential antimicrobial, antioxidant agent to being central nervous system (CNS) stimulants. This chapter focuses specifically on physiological impacts that each of its constituents have over our nervous system like role of L-theanine to enhance dopamine and serotonin levels, theobromine, and theophylline for stimulating CNS, caffeine to inhibit adenosine receptors, hence, causing increase in brain activity etc. along with many more neuroprotective properties of tea constituents.

Keywords: central nervous system (CNS), epigallocatechin-3-gallate (EGCG), epigallocatechin (EGC), epicatechin-3-gallate (ECG), and epicatechin (EC), theaflavin (TF-1), Laminin receptor (67LR)

1. Introduction

Tea is known as a part of many traditional medicinal practices (Ayurvedic, Chinese etc.) and as a health supplement of daily usage from ancient era. Tea (Camellia sinensis) belongs to Theaceae family and is known as a perennial shrub/tree which reaches up to the height of 30 feet, however it is pruned cropped at a lesser height of around 2–5 feet for cultivation. It is of various types such as black, white, green, oolong varieties. Rooibos or “Red” and Pu-erh tea are produced from tea plant leaves, which are oval and dark green in color, with notched boundaries, and its flowers are usually white, fragrant bunched, together or separately. The tea plant, C. sinensis, initially was an indigenous species that belonged to China but later spread to other parts of the world like—Indian subcontinent, Japan, Russia and then to Europe in the late seventeenth century. The various forms of tea (Green, oolong, and black tea) originated from the same plant (C. sinensis) but got differentiated, depending on their color display, organoleptic taste, distinctive flavor and their phytochemical content which was eventually a result of different fermentation processes adopted for their production [1].

There are two main varieties of the tea plant, named as Camellia sinensis and Camellia sinensis var. assamica. The Chinese variant, Camellia sinensis, has smaller
leaves and is more tolerant to cold weather. It is observed as a perennial plant going up to the height of 3 m in case of *C. var sinensis*, whereas it was up to 10–15 m tall with less branching in *C. var assamica* [2, 3]. But since in tea cultivation practices the plants are usually pruned and are kept at lower height (1–2 m) hence, promoting them to spread their branches horizontally. In the ancient scriptures of China, tea processing and consumption from these two varieties are reported to be practiced from last 4000 years. The second variety, *Camellia sinensis var. assamica*, is a native to the Assam region in India and thrives well in tropical and low elevation areas in the Indian subcontinent. This variety of tea plantation is commonly cultivated in the tropical and subtropical regions of India and apart from its use as a beverage it has been reported for many utilities like—it has high medicinal value, used for extraction of oil (Tea Tree oil). As per the recent global studies conducting in 2016, it was found that Turkey was listed as one of the highest tea devouring country with consumption of approximately 6.96 pounds per year per capita. On the contrary, China was observed with little less annual consumption of around 1.25 pounds per person but, it has shown highest tea production globally, followed by India and Kenya at second and third positions respectively.

Also, on global platform it was estimated that, almost 3.8 billion gallons of tea, in which black tea has 80%, green tea 16% and remaining 4% was oolong, white and dark tea share was consumed in United States in the same year (2016) [4]. These data exhibit the ever-growing popularity of tea consumption among the masses irrespective of their region of cultivation.

### 2. Potential health benefits of tea constituents

The commonly found and highest content of chemical constituents found in leaves of tea are polyphenols (catechins and flavonoids), inorganic elements (e.g., fluorine, aluminum, and manganese), alkaloids (caffeine, theobromine, theophylline, etc.), amino acids, volatile oils, lipids, polysaccharide, and vitamins. However, the polyphenolic content which is present in the highest concentration, is primarily responsible for its most of the therapeutic benefits. Consequently, flavonoid contents impart its antimicrobial, antioxidant, anti-allergic and anti-inflammatory effects. The phenolic content variants are further elaborated and sub-classified as catechin, gallocatechin, epigallocatechin, epicatechin gallate, epicatechin, and epigallocatechin-gallate (EGCG), the latter being the most active component [2, 5]. Further, the molecular structure of green tea polyphenols exhibits active hydroxyl hydrogen which effectively scavenge free radicals hence, slowing down the detrimental changes in most of the physiological processes existing in human body. Reportedly, tea polyphenols strongly exhibits the movement of glutathione peroxidase and superoxide dismutase causing higher scavenging rate. The phytoconstituents of tea reflects multiple therapeutic benefits on our various diverse physiological systems through various biochemical and pharmacological processes like—antioxidant activities, inhibition of cell proliferation, induction of apoptosis, cell cycle arrest and modulation of carcinogen metabolism [6, 7]. Similarly, in CNS, another constituent in green tea, L-theanine increases the dopamine and serotonin levels resulting in mood elevation and stress reduction. Also, caffeine content in same sources aids in increasing the focus, vigilance, concentration and reasoning ability [8]. Theobromine and theophylline are known as potential CNS stimulants. Numerous studies have shown that most of the tea polyphenols have reactive oxygen and nitrogen species (ROS) scavenging activity along with an ability to chelate down redox-active transition metal ions. Currently, apart from all the listed...
health benefits exhibited by the tea and its polyphenols, the focus is towards exploring its chemo preventive, hypolipidemic and anti-obesity effects in all sorts of possible in vitro and in vivo model systems [9].

3. Types of tea variants

Tea leaves are either classified on the basis of their consumption and texture it has or on the processing method adopted for their leaves. Hence, the classification, studied commonly for tea is based on its varied fermentation degree process and is comprised of basically three types: non-fermented (green), semi-fermented (oolong) and entirely fermented (black) [10]. The tea processing starts firstly, from picking up the appropriate and selected tea leaves from shrub or tea tree which undergoes fractional withering. Then roasting the same leaves to inactivate oxidative enzymes, followed by rolling up, drying and sorting the same leaves. The color of the final tea product is usually green tasting slightly constringent. So many countries like China, the taste of green tea is improvised by supplementing aromatic fruits (orange) or flowers (jasmine). Further, the tea processing steps in case of black tea is more complex, as after withering process the tea leaves are subjected for two steps fermentation processes, in the last step of fermentation they have been rolled up and then fermented. Lastly, they are roasted till they become dark-brown or brown black in color imparting a roasting aroma so as to block the activity of enzymes (polyphenol oxidase and glycosidase) along with further, fermentation of the same [5]. Another variant, oolong tea which is partially fermented type usually has shorter fermentation time in comparison to the black one.

3.1 Green tea

Green tea is a non-fermented tea which is largely consumed by the population of china and japan. After cultivation, tea leaves are first withered for the inactivation of enzyme (polyphenol) which is liable for oxidation of tea catechins into their oligomeric forms (thearubigins and theaflavins). To avoid the oxidation and polymerization of tea leaves, they are steamed up and dried at high temperatures [6, 9, 11]. The Chinese traditional dietary system do have another packed form of green tea called “black powder”, named after type of leaves processing method. Where these leaves individually are stirred and wrapped into a round pellet looking like explosives. It prevents it from any kind of physical damage and maintains its fragrance and flavor. Polyphenols present in green tea are flavonols (quercetin, kaempferol, and rutin), caffeine, phenolic acids, theanine, flavor, and leucoanthocyanins, which show 40% of dry weight of leaves [12, 13]. The highly water-soluble parts of tea comprises of biochemical components like (−) epigallocatechin-3-gallate (EGCG), epigallocatechin (EGC), epicatechin-3-gallate (ECG), and epicatechin (EC) (As listed in Table 1) [9, 14]. Further, it’s also been reported that 1 kg of green tea has around 191 g of the above listed catechins, 36 g of caffeine, and 5.2 g of flavonols [15]. In green tea there are 10–15% of polyphenols present whereas it’s lesser in black tea, i.e. around 5%. Dry weight of green tea constitutes about 42% polyphenols which is composed of 26.7% of catechin-gallate components such as ECG (2.25%), EGC (10.32%), Catechins (0.53%), EGCG (11.16%) and Epicatechin (2.45%) [16]. It’s been estimated that in one cup of green tea the expected concentration of EGCG is between 2.1–2.4 mg/mL and after testing the effects of both Green tea and EGCG (equivalent of 4–8 cups per day) on human subjects there was no appreciable side effects observed [17, 18]. Epidemiological
| S. No | Polyphenols present | Structure of the polyphenols | Therapeutic benefit |
|-------|---------------------|------------------------------|-------------------|
| 1.    | (+)-Catechin        | ![Structure of (+)-Catechin](image1) | Increases cellular lipid antioxidant activity, act as brain permeable iron chelator |
| 2.    | Epicatechin         | ![Structure of Epicatechin](image2) | It inhibits lipid peroxidation in cell membrane and generation of hydrogen peroxide ions in keratinomyocytes. Also, can induce ATF3 (tumor suppressor proteins) through EGR-1 activation. |
| 3.    | Epigallocatechin    | ![Structure of Epigallocatechin](image3) | Act as prophylactic agents against *Bordetella pertussis* infection. |
| 4.    | (-)-Catechin gallate (GC) | ![Structure of (-)-Catechin gallate (GC)](image4) | Antioxidant |
studies too, have suggested protective and suppressive effects against many types of human cancer (including that of skin, lung, liver, esophagus, and stomach) after tea consumption [19–21].

3.2 Black tea

This variety of tea is very famous in North America, Europe, and India. Black tea is extracted from the new, soft, firstly appeared leaves of *Camellia sinensis* and is one of the most broadly devoured non-mixed drinks. The flavor, quality and taste of these drinks tends to change with variation in their topographical and climatic conditions [22, 23]. This variety of tea offers, its simple quality parameters, specifically, theaflavins, thearubigins, and caffeine. Theaflavins adds to the abstinence (liveliness) and splendor, while thearubigins adds to the shading and body (mouth

| S. No | Polyphenols present | Structure of the polyphenols | Therapeutic benefit |
|-------|---------------------|-----------------------------|--------------------|
| 5.    | (+)-Epicatechin gallate (ECG) | ![Image](image1.png) | Radical scavengers and Protective effect on lipid peroxidation in phospholipid Bilayers, Antioxidants |
| 6.    | (−)-Epigallocatechin gallate (EGCG) | ![Image](image2.png) | Has strong antioxidant and anti-inflammatory activities, induces cell apoptosis by hindering cellular cycles in pancreatic cancer, and decreases autoimmune reactions. |

Table 1. Representing the types and structure of phytocompounds present in the green tea with their therapeutic benefits.
feel); and caffeine is responsible for stimulatory impact of dark tea. In Black tea, the compound is permitted to act in a way that the leaves are completely aged to give the trademark fragrance and shade of dark tea [24, 25]. Arranged by squashing tea leaves and permitting enzyme mediated oxidation, which leads to the formation of oligomeric flavanols by tea catechins, including theaflavins, thearubigins, and different oligomers [26–28]. Further, the associated compounds like Theaflavins includes, combination of theaflavin (TF-1), theaflavin-3-gallate (TF-2a), theaflavin-3’-gallate9TF-2b), and theaflavin-3, 3’-digallate (TF-3), having lower tea catechin content (3–10% [w/w]), with theaflavins and thearubigins showing around 2–6% (w/w) and 10–20% (w/w) dry weight. Theaflavins, are orange or orange-red colored benzotropolone structure formed due to co-oxidation and oxidative dimerization of catechins (Table 2) [29, 30].

| S. No | Polyphenols present | Structure of the polyphenols | Therapeutic benefit |
|-------|---------------------|-----------------------------|--------------------|
| 1.    | Thearubigins        | ![Image](image1.png)         | Shows antioxidative properties, prevent activity of enzymes participating in enzymatic lipid peroxidation |
| 2.    | Theaflavins         | ![Image](image2.png)         | Shows inhibitory effect against phytopathogenic bacteria, induce apoptosis, strong antioxidants, prevent free radical generation, show metal chelating abilities |

Table 2. Representing the types and structure of phytocompounds present in the black tea with their therapeutic benefits.

3.3 Oolong tea

Oolong tea is a conventional Chinese tea with a different, unique production method and one of the most popular beverages in china with its Chinese name meaning as “Black dragon tea”. It is a semi fermented tea with restricted time of oxidation as compared to black tea and contains phytocompounds of both black and
| S. No | Polyphenols present | Structure of the polyphenols | Therapeutic benefit |
|-------|---------------------|------------------------------|--------------------|
| 1.    | (+)-Catechin         | ![Image](image1.png)         | Increases cellular lipid antioxidant activity, act as brain permeable iron chelator |
| 2.    | Epicatechin          | ![Image](image2.png)         | It inhibits lipid peroxidation of cell membrane and induces tumor suppressor proteins |
| 3.    | (-)-Epigallocatechin (EGC) | ![Image](image3.png)      | It acts as prophylactic agents against *Bordetella pertussis* infection. |
| 4.    | (-)-Catechin gallate (GC) | ![Image](image4.png)      | Antioxidant |
| 5.    | (+)-Epicatechin gallate (ECG) | ![Image](image5.png)      | Radical scavengers and Protective effect on lipid peroxidation in phospholipid Bilayers, Antioxidants |
| 6.    | (-)-Epigallocatechin gallate (EGCG) | ![Image](image6.png)      | Has strong antioxidant and anti-inflammatory activities, induces cell apoptosis by hindering cellular cycles in pancreatic cancer, and decreases autoimmune reactions. |

Table 3.
*Representing the types and structure of phytocompounds present in the oolong tea with their therapeutic benefits.*

green tea. It has approximately half of the EGCG from green tea, while double quantity of polymerized polyphenols and theaflavins of black tea. The procyanidins produced in oolong tea are formed due to its unique fermentation process. The
leaves are first withered, sun dried and then allowed for oxidation before rolling and twisting. *Camellia sinensis* is used for the production of Oolong tea and tastes very different from green and black tea. The tea processing method differs which makes them significantly different from each other, even if they are produced from the same plant [31].

As all tea leaves are green when they are plucked. Green tea undergoes, heating process in order to inhibit the oxidation of tea leaves. They are rolled up to break the cell structure. While, oolong tea is plucked and kept in optimized condition and allowed for oxidation. Due to difference in its processing method oolong tea tastes different from its sub varieties. It shows a sweet and fruity flavor with striking honey odors to woody and dense with roasted aromas, or even green and fresh with flowery aromas. They are processed by different methods as some are wrapped-curved into small beads and others are rolled into curly leaves. In china, oolong tea is added with flavors like jasmine flowers (Tables 3 and 4) [32].

### 4. Therapeutic benefits of tea in CNS health

As discussed earlier, the health-promoting properties of the tea plants are often credited to their active ingredients including polyphenols. Tea flavanols are a group of natural polyphenols (epicatechins) found in most of the varieties of tea. Their therapeutic benefits although are immense, but they do have contributed exclusively in neural health of living beings. Likewise, the polyphenols of green tea are reported extensively in preventing neuronal degradation by inhibiting neurotoxin formation in cells [33, 34]. Also, in one of the recent study done, with transitional metal (iron and copper) chelating property or EGCG, suggested its possible effective role in treating certain forms of neurodegenerative diseases. Similarly, the antioxidative property of EGCG exhibits protection against advanced glycation end

| Polyphenols                           | Chemical structure | Oolong tea phytophosphol concentration (mg/g) | Green tea phytophosphol concentration (mg/g) |
|---------------------------------------|--------------------|----------------------------------------------|---------------------------------------------|
| Caffeine                              | C₈H₁₀N₄O₂          | 64                                           | 53                                          |
| Flavanol with galloyl moiety          |                    |                                              |                                              |
| Catechin                              | C₁₅H₁₄O₆           | 30                                           | 43                                          |
| Epicatechin                           | C₁₅H₁₄O₆           | 6                                            | 25                                          |
| Gallicatechin                         | C₁₅H₁₄O₇           | 10                                           | 5                                           |
| Epigallocatechin                      | C₁₅H₁₄O₇           | 2                                            | 8                                           |
| Flavanol without galloyl moiety       |                    |                                              |                                              |
| Epigallocatechin gallate              | C₂₅H₁₈O₁₁          | 14                                           | 29                                          |
| Gallicatechin gallate                 | C₂₅H₁₆O₁₁          | 16                                           | 19                                          |
| Epicatechin gallate                   | C₂₅H₁₆O₁₁          | 3                                            | 6                                           |
| Catechin gallate                      | C₂₅H₁₆O₁₀          | 7                                            | 5                                           |
| Oolong tea polymerized polyphenols (OTPP) |               | 114                                          | –                                           |

Table 4. Comparing the polyphenolic contents of oolong and green tea.
products (AGEs) induced neuronal cells injury along with inhibit AGEs—AGE receptor (RAGE) interaction intervened pathways, suggesting a possible therapeutic role of tea catechins for neurodegenerative diseases. Hence, both black and green tea varieties are reported to contribute immensely for the protection against neurodegenerative diseases [34–36]. Also, oxidative variations of cellular components such as nucleic acids, lipids, and proteins are prevented by bidirectional antioxidative property [37]. The oxidation of these components in aqueous phase is responsible for initiation of membrane lipid peroxidation [35].

Moreover, these water soluble tea polyphenols, particularly catechins have effective potential to scavenge free radicals and reduce the versatility of free radicals in the lipid structures too. Polyphenols enters the phospholipids bilayer, coating it with film and, balancing out the impact, by adjusting the lipid pressing ability [38]. They also contain higher amount of chemically dynamic metal particles (iron and copper) creating in-situ oxygen radicals by Fenton’s response [39, 40].

Due to the existence of hydroxyl ions on polyphenol ring metal chelation effects can be observed. Metal Chelating effects by Green and Black Tea additionally, restricts lipid per oxidation and secures the essential lipid structures present in cerebrum leading to reduce oxidative stress [10, 27, 41]. Furthermore, it’s been observed in research studies that the phytocompounds of tea (Green/Black) also prevents, the division of mitochondrial layer against iron induced lipid per oxidation and enhanced the survival rate in many in vivo models [42, 43]. Hence, it can be concluded from the recent research updates, that the high metal chelating quality of its constituents may provide a unique essential neuroprotection against many neurological disorders [44].

One of the essential pathological cause in Alzheimer’s disease (AD) is irregular contact of free chelatable iron which is responsible for the deposition of neocortical amyloid peptide and deposition of metals, phosphorylation of tau and formation of tangles due to production of tau protein from microtubules [45, 46]. Also, the activation of amyloid cascades, which produces amyloid by β-amyloid precursor protein (APP), accumulates in the presence of divalent metal ions into amyloid fibrils leading to a major cause of AD [47, 48].

Recent studies have reported that the delay in onset or slowdown of the neurodegenerative process along with minimal neural deterioration was observed in the population consuming tea infusions on regular basis [49]. There scientific correlation suggests that the reduction in amyloid beta (Aβ) fibril production in the presence of EC and EGCG is suspected to regulate the amyloid protein precursor (APP) enzyme activity [50]. Additionally, it been also suggested that the regular consumption of tea (green and black) may lead to the acetyl cholinesterase (AChE) activity inhibition, further causing halt in acetylcholine production [51, 52]. Besides this, it was found that there was inhibition of butyrylcholinesterase (BuChE) enzyme deposits in the brain of AD subjects after consuming green tea or black tea for certain time [53]. These research findings advises that active phytocompounds present in tea can be used to obstruct the development of AD [54].

5. Mechanism of tea polyphenols

5.1 Mechanism of action of EGCG to improve cardiovascular function and anticancer activity

Tachibana et al. [55] studied the effect of tea polyphenols, and suggested that EGCG directly binds to the Laminin receptor (67LR), located on the peptide LR161-170. This receptor shows a high expression only in cancerous cells. This
suggests that EGCG specifically binds to the cancer cells and binding of EGCG with 67LR receptor activate the enzyme protein kinase B which further activate ENOS pathway leading to vasodilation that contributes to the improvement of cardiovascular function of cell [16, 55]. It also elevates the activity of CGMP that activate the PKC/Acidic sphingomyelinases that induces the apoptosis in cancerous cells.

5.2 Antagonistic actions of theanine on glutamate receptors

Nozawa et al. [56] discovered the death of 50% of neurons at higher concentration of glutamate but when pre-treated with theanine, the possibility of death was significantly decreased. Many more recent updates suggested that increased glutamate level in cell may lead to massive influx of Ca⁺ ions and increases the formation of ROS which leads to the death of neuronal cells. In order to avoid the toxicity of glutamate, the glutamate receptors binds with theanine. Theanine has same structure as glutamate so in presence of theanine it shows a competitive inhibition and inhibit the binding of glutamate to its receptor. Furthermore, Kakuda et al. [57] studies the inhibiting effect of glutamate receptors by theanine that suggests the neuroprotective role of theanine. It shows the specific binding of theanine to NMDA receptor to inhibit the glutamate binding affinity. Theanine has an antagonistic effect to glutamate receptors. Glutamine, derived from glutamate, is synthesized by glutamine synthetase. Theanine can inhibit the transport of glutamine and regulate the glutamate-glutamine cycle in the neurons and, thus, shows the neuroprotective effect of tea (Figure 1) [58].

![Figure 1](image.png)

*Schematic representation. Inhibition effect of theanine on glutamate receptor.*

5.3 Therapeutic limitations of tea compounds

Although being therapeutically crucial compound tea phytocompounds do have certain harmful side effects, if over consumed or overdosed like—higher Caffeine
content, Aluminum presence and the effects of tea polyphenols on iron bioavailability [59, 60]. In the study done by Lin et al. [31], it was been reported that the caffeine content in tea is available in following order: black tea > oolong tea > green tea > fresh tea leaf. Similarly, Cabrera et al. also studied the caffeine content and its after effects in 45 samples of tea and determined that black tea has the high concentration of caffeine (41.5–67.4 mg/g), whereas oolong and green tea samples have less amount of caffeine content of 32.5 and 29.2 mg/g, respectively [61]. The harmful effects of caffeine content in tea are listed as—vomiting, sleep disorder, nervousness, tachycardia, and epigastric pain etc. [62]. Hence, tea intake is strictly restricted in patients suffering from cardiovascular problems. Breastfeeding and pregnant women should avoid over-consumption of green tea because it do causes tachycardia in them giving rise to higher health risks to fetus [63, 64]. The presence of aluminum in black and green teas also suggested increased accumulation of the same inside the body affecting the neural well-being and causing neurological disorders [65].

6. Conclusion

It can be concluded in the review that tea polyphenols with other constituents have a very high therapeutic potential including the potency to decrease the threat of diseases such as cancer, cardiovascular, diabetes and neurodegenerative diseases. It has proven to be a strong antioxidant agent that shows a therapeutic effect of tea. To evaluate the efficacy of tea many experiments are being conducted which shows a promising data from many trials and other ongoing trials are conducted to study the therapeutic effect of tea. Because less information is available about bioavailability of tea polyphenols after intake of tea, studies of bioavailability polyphenols of tea is needed on animals and humans to evaluate its protective role.

Author details

Manisha Singh*, Vandana Tyagi and Shriya Agarwal
Department of Biotechnology, Jaypee Institute of Information Technology, Noida, UP, India

*Address all correspondence to: manishasingh1295@gmail.com

IntechOpen

© 2019 The Author(s). Licensee IntechOpen. This chapter is distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.
References

[1] Da Silva Pinto M. Tea: A new perspective on health benefits. Food Research International. 2013;53(2): 558-567

[2] Wierzejska R. Tea and health—A review of the current state of knowledge. Przegląd Epidemiologiczny. 2014;68(3):501-506

[3] Fern K. Camellia sinensis assamica 2014. Available from: http://tropical.theferns.info/viewtropical.php?id= Camellia+sinensis+assamica

[4] Grigg D. The worlds of tea and coffee: Patterns of consumption. GeoJournal. 2002 Aug 1;57(4):283-294

[5] George VC, Vijesh VV, Amararathna DIM, Lakshmi CA, Anbarasu K, Naveen Kumar DR, et al. Mechanism of action of flavonoids in prevention of inflammation-associated skin cancer. Current Medicinal Chemistry. 2016 Oct 1;23(32):3697-3716

[6] Riegecker S, Wiczynski D, Kaplan MJ, Ahmed S. Potential benefits of green tea polyphenol EGCG in the prevention and treatment of vascular inflammation in rheumatoid arthritis. Life Sciences. 2013;93(8):307-312

[7] Das S, Tanwar J, Hameed S, Fatima Z, Manesar G. Antimicrobial potential of epigallocatechin-3-gallate (EGCG): A green tea polyphenol. Journal of Biochemical Pharmacology Research. 2014;2(3):167-174

[8] Gramza-Michalowska A. Caffeine in tea Camellia sinensis—Content, absorption, benefits and risks of consumption. The Journal of Nutrition, Health & Aging. 2014;18(2):143-149

[9] Narotzki B, Levy Y, Aizenbud D, Reznick AZ. Green tea and its major polyphenol EGCG increase the activity of oral peroxidases. In: Respiratory Regulation—The Molecular Approach. Dordrecht: Springer; 2013. pp. 99-104

[10] Chan EW, Soh EY, Tie PP, Law YP. Antioxidant and antibacterial properties of green, black, and herbal teas of Camellia sinensis. Pharmacognosy Research. 2011;3(2):266

[11] Senanayake SN. Green tea extract: Chemistry, antioxidant properties and food applications—A review. Journal of Functional Foods. 2013;5(4):1529-1541

[12] Le Fanu S. Green Tea. UK: Penguin; 2016

[13] Afzal M, Safer AM, Menon M. Green tea polyphenols and their potential role in health and disease. Inflammopharmacology. 2015;23(4): 151-161

[14] Reygaert WC. The antimicrobial possibilities of green tea. Frontiers in Microbiology. 2014;5:434

[15] Colon M, Nerin C. Molecular interactions between caffeine and catechins in green tea. Journal of Agricultural and Food Chemistry. 2014; 62(28):6777-6783

[16] Kim HS, Quon MJ, Kim JA. New insights into the mechanisms of polyphenols beyond antioxidant properties; lessons from the green tea polyphenol, epigallocatechin 3-gallate. Redox Biology. 2014;2:187-195

[17] Dietz C, Dekker M. Effect of green tea phytochemicals on mood and cognition. Current Pharmaceutical Design. 2017;23(19):2876-2905

[18] Iwasaki M, Mizusawa J, Kasuga Y, Yokoyama S, Onuma H, Nishimura H, et al. Green tea consumption and breast cancer risk in Japanese women: A case-control study. Nutrition and Cancer. 2014;66(1):57-67

[19] Wang H, Provan GJ, Helliwell K. Tea flavonoids: Their functions,
utilisation and analysis. Trends in Food Science & Technology. 2000;11(4–5): 152-160

[20] Pang J, Zhang Z, Zheng TZ, Bassig BA, Mao C, Liu X, et al. Green tea consumption and risk of cardiovascular and ischemic related diseases: A meta-analysis. International Journal of Cardiology. 2016;202:967-974

[21] Ide K, Yamada H, Takuma N, Park M, Wakamiya N, Nakase J, et al. Green tea consumption affects cognitive dysfunction in the elderly: A pilot study. Nutrients. 2014;6(10):4032-4042

[22] Both S, Chemat F, Strube J. Extraction of polyphenols from black tea—Conventional and ultrasound assisted extraction. Ultrasonics Sonochemistry. 2014;21(3):1030-1034

[23] Butt MS, Imran A, Sharif MK, Ahmad RS, Xiao H, Imran M, et al. Black tea polyphenols: A mechanistic treatise. Critical Reviews in Food Science and Nutrition. 2014;54(8):1002-1011

[24] Kumar RS, Murugesan S, Kottur G, Gyamfi D. Black tea: The plants, processing/manufacturing and production. In: Tea in Health and Disease Prevention. Amsterdam, USA; 2013. pp. 41-57

[25] Ho CT, Zheng X, Li S. Tea aroma formation. Food Science and Human Wellness. 2015;4(1):9-27

[26] Van Duynhoven J, Vaughan EE, Van Dorsten F, Gomez-Roldan V, De Vos R, Vervoort J, et al. Interactions of black tea polyphenols with human gut microbiota: Implications for gut and cardiovascular health. The American Journal of Clinical Nutrition. 2013;98(6):1631S-1641S

[27] Grassi D, Desideri G, Di Giosia P, De Feo M, Fellini E, Cheli P, et al. Tea, flavonoids, and cardiovascular health: Endothelial protection. The American Journal of Clinical Nutrition. 2013;98(6):1660S-1666S

[28] Bansal S, Choudhary S, Sharma M, Kumar SS, Lohan S, Bhardwaj V, et al. Tea: A native source of antimicrobial agents. Food Research International. 2013;53(2):568-584

[29] Wang C, Li Y. Research progress on property and application of theaflavins. African Journal of Biotechnology. 2006;5(3):213-218

[30] Tounekti T, Joubert E, Hernández I, Munné-Bosch S. Improving the polyphenol content of tea. Critical Reviews in Plant Sciences. 2013;32(3):192-215

[31] Lin YS, Tsai YJ, Tsay JS, Lin JK. Factors affecting the levels of tea polyphenols and caffeine in tea leaves. Journal of Agricultural and Food Chemistry. 2003;51(7):1864-1873

[32] Zhu QY, Hackman RM, Ensuns J, Holt RR, Keen CL. Antioxidative activities of oolong tea. Journal of Agricultural and Food Chemistry. 2002;50(23):6929-6934

[33] Pan MH, Lin CC, Lin JK, Chen WJ. Tea polyphenol (—)-epigallocatechin-3-gallate suppresses heregulin-beta1-induced fatty acid synthase expression in human breast cancer cells by inhibiting phosphatidylinositol 3-kinase/Akt and mitogen-activated protein kinase cascade signaling. Journal of Agricultural and Food Chemistry. 2007;55(13):5030-5037

[34] Mandel S, Weinreb O, Reznichenko L, Kalfon L, Amit T. Green tea catechins as brain-permeable, non toxic iron chelators to "iron out iron" from the brain. Journal of Neural Transmission Supplementum. 2006;71:249-257

[35] Bastianetto S, Yao ZX, Papadopoulos V, Quirion R. Neuroprotective effects of green and black teas and their catechin gallate
esters against beta-amyloid-induced toxicity. The European Journal of Neuroscience. 2006;23(1):55-64

[36] Oboh G, Rocha JB. Antioxidant and neuroprotective properties of sour tea (Hibiscus sabdariffa, calyx) and green tea (Camellia sinensis) on some pro-oxidant-induced lipid peroxidation in brain in vitro. Food Biophysics. 2008 Dec 1;3(4):382

[37] Reznichenko L, Amit T, Youdim MB, Mandel S. Green tea polyphenol (−)-epigallocatechin-3-gallate induces neurorescue of long-term serum-deprived PC12 cells and promotes neurite outgrowth. Journal of Neurochemistry. 2005;93(5):1157-1167

[38] Seeram NP, Henning SM, Niu Y, Lee R, Scheuller HS, Heber D. Catechin and caffeine content of green tea dietary supplements and correlation with antioxidant capacity. Journal of Agricultural and Food Chemistry. 2006;54(5):1599-1603

[39] Lorenz M. Cellular targets for the beneficial actions of tea polyphenols. The American Journal of Clinical Nutrition. 2013;98(6):1642S-1650S

[40] Swomley AM, Förster S, Keeney JT, Triplett J, Zhang Z, Sultana R, et al. Abeta, oxidative stress in Alzheimer disease: Evidence based on proteomics studies. Biochimica et Biophysica Acta (BBA)-Molecular Basis of Disease. 2014;1842(8):1248-1257

[41] Rodrigues MJ, Neves V, Martins A, Rauter AP, Neng NR, Nogueira JM, et al. In vitro antioxidant and anti-inflammatory properties of Limonium algarvense flowers’ infusions and decoctions: A comparison with green tea (Camellia sinensis). Food Chemistry. 2016;200:322-329

[42] Higuchi A, Yonemitsu K, Koreeda A, Tsunenari S. Inhibitory activity of epigallocatechin gallate (EGCG) in paraquat-induced microsomal lipid peroxidation—A mechanism of protective effects of EGCg against paraquat toxicity. Toxicology. 2003;183(1–3):143-149

[43] Sutherland BA, Rahman RM, Appleton I. Mechanisms of action of green tea catechins, with a focus on ischemia-induced neurodegeneration. The Journal of Nutritional Biochemistry. 2006;17(5):291-306

[44] Lardner AL. Neurobiological effects of the green tea constituent theanine and its potential role in the treatment of psychiatric and neurodegenerative disorders. Nutritional Neuroscience. 2014;17(4):145-155

[45] Weinreb O, Amit T, Mandel S, Youdim MB. Neuroprotective molecular mechanisms of (−)-epigallocatechin-3-gallate: A reflective outcome of its antioxidant, iron chelating and neuritogenic properties. Genes & Nutrition. 2009;4(4):283

[46] Yang CS, Hong J. Prevention of chronic diseases by tea: Possible mechanisms and human relevance. Annual Review of Nutrition. 2013;33:161-181

[47] Weinreb O, Mandel S, Amit T, Youdim MB. Neurological mechanisms of green tea polyphenols in Alzheimer's and Parkinson’s diseases. The Journal of Nutritional Biochemistry. 2004;15(9):506-516

[48] Bush AI. The metallobiology of Alzheimer's disease. Trends in Neurosciences. 2003;26:207-214

[49] Rezai-Zadeh K, Arendash GW, Hou H, Fernandez F, Jensen M, Runfeldt M, et al. Green tea epigallocatechin-3-gallate (EGCG) reduces β-amyloid mediated cognitive impairment and modulates tau pathology in Alzheimer transgenic mice. Brain Research. 2008;1214:177-187
[50] Checkoway H, Powers K, Smith-Weller T, Franklin GM, Longstreth WT Jr, Swanson PD. Parkinson's disease risks associated with cigarette smoking, alcohol consumption, and caffeine intake. American Journal of Epidemiology. 2002;155(8):732-738

[51] Wei CC, Yu CW, Yen PL, Lin HY, Chang ST, Hsu FL, et al. Antioxidant activity, delayed aging, and reduced amyloid-β toxicity of methanol extracts of tea seed pomace from Camellia tenuifolia. Journal of Agricultural and Food Chemistry. 2014;62(44):10701-10707

[52] Chakrabarti S, Sinha M, Thakurta IG, Banerjee P, Chattopadhyay M. Oxidative stress and amyloid beta toxicity in Alzheimer’s disease: Intervention in a complex relationship by antioxidants. Current Medicinal Chemistry. 2013;20(37):4648-4664

[53] Harada M, Kan Y, Naoki H, Fukui Y, Kageyama N, Nakai M, et al. Identification of the major antioxidative metabolites in biological fluids of the rat with ingested (+)-catechin and (−)-epicatechin. Bioscience, Biotechnology, and Biochemistry. 1999;63(6):973-977

[54] Skrzydlewska E, Ostrowska J, Farbiszewski R, Michalak K. Protective effect of green tea against lipid peroxidation in the rat liver, blood serum and the brain. Phytomedicine. 2002;9(3):232-238

[55] Fujimura Y, Sumida M, Sugihara K, Tsukamoto S, Yamada K, Tachibana H. Green tea polyphenol EGCG sensing motif on the 67-kDa laminin receptor. PLoS One. 2012;7(5):e37942

[56] Kakuda T, Nozawa A, Sugimoto A, Niino H. Inhibition by theanine of binding of [3H] AMPA, [3H] kainate, and [3H] MDL 105,519 to glutamate receptors. Bioscience, Biotechnology, and Biochemistry. 2002;66(12):2683-2686

[57] Kakuda T. Neuroprotective effects of theanine and its preventive effects on cognitive dysfunction. Pharmacological Research. 2011;64(2):162-168

[58] Kakuda T, Hinoi E, Abe A, Nozawa A, Ogura M, Yoneda Y. Theanine, an ingredient of green tea, inhibits [3H] glutamine transport in neurons and astroglia in rat brain. Journal of Neuroscience Research. 2008;86(8):1846-1856

[59] Jain A, Manghani C, Kohli S, Nigam D, Rani V. Tea and human health: The dark shadows. Toxicology Letters. 2013 Jun 20;220(1):82-87

[60] Ruchi V. An overview on tea. International Journal of Pharmaceutical Research. 2013;3(3):36

[61] Hayat K, Iqbal H, Malik U, Bilal U, Mushtaq S. Tea and its consumption: Benefits and risks. Critical Reviews in Food Science and Nutrition. 2015;55(7):939-954

[62] Turnbull D, Rodricks JV, Mariano GF, Chowdhury F. Pharmacology. Caffeine and cardiovascular health. Regulatory Toxicology and Pharmacology. 2017;89:165-185

[63] Lu JH, He JR, Shen SY, Wei XL, Chen NN, Yuan MY, et al. Does tea consumption during early pregnancy have an adverse effect on birth outcomes? Birth. 2017;44(3):281-289

[64] Bedrood Z, Rameshrad M, Hosseinzadeh H. Toxicological effects of Camellia sinensis (green tea): A review. Phytotherapy Research. 2018

[65] Hamdaoui MH, Chabchoub S, Hédhili A. Iron bioavailability and weight gains to iron-deficient rats fed a commonly consumed ‘Tunisian meal ‘bean seeds ragout’ with or without beef and with green or black tea decoction. Journal of Trace Elements in Medicine and Biology. 2003;17(3):159