The Impact of Coffee on Health

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ABSTRACT
Coffee is one of the most popular and widely consumed beverages worldwide due to its stimulating effects on the central nervous system as well as its taste and aroma. Coffee is a complex mixture of more than 800 volatile compounds whereas caffeine and chlorogenic acids are the most common compounds. During the last years, coffee has progressively moved to a less negative position on health due to its better-known pharmacology. Caffeine, e.g., in a cup of coffee, appears to exert most of its effects through an antagonism of the adenosine receptors. Novel approaches in epidemiological studies and experimental researches suggest that coffee consumption may help to prevent several chronic diseases, including type 2 diabetes mellitus and liver disease. Most prospective cohort studies have not found coffee consumption to be associated with a significantly increased cardiovascular disease risk. There is also evidence that decaffeinated coffee may, in some respect, have similar benefits as regular coffee, indicating that besides caffeine other components contribute to the health protecting effects. For adults consuming moderate amounts of coffee (3–4 cups/d providing 300–400 mg/d of caffeine), there is little evidence of health risks and some evidence of health benefits. This review provides up-to-date information about coffee on health. Topics addressed include the cardiovascular system, liver diseases, and diabetes as well as gastrointestinal disorders.

Introduction
In 2016/17, the production of roar coffee amounted to about 151.62 million 60 kg bags [1], and it is estimated that 2.25 billion cups of coffee are consumed each day worldwide [2]. Coffee is served internationally and most countries have developed its own preferences about how to prepare and present it.

The history of coffee goes at least as far back as the 10th century, with a number of legends surrounding its use. The native (undomesticated) origin of coffee is thought to have been Ethiopia. The earliest substantiated evidence of either coffee drinking or knowledge of the coffee tree is from the 15th century, in the Sufi monasteries of Yemen. By the 16th century, it had reached the rest of the Middle East, South India, Persia, Turkey, and Northern Africa. Coffee then spread to the Balkans, Italy, and the rest of Europe, to Indonesia and then to America [3]. Although coffee was introduced in Europe only a few hundred years ago, consumption of this beverage now occupies a significant place in our national cultures.

Coffee is taken as a brewed beverage that is prepared from the roasted seeds of a bush of the genus Coffea. The coffee beans are contained in berries that, once matured, are processed and dried. The two main species are Coffea Arabica (coffee Arabica) and Coffea canephora (coffee Rustica). They have a large production history and an important role both in the global market and researches [4].

The high consumption of coffee may have a substantial effect on public health. Therefore, it is no wonder that coffee stimulates the interest of researchers and clinicians. In PubMed in May 2017, the term “Coffee” resulted in 12583 hits including 998 reviews and 1666 clinical trials. Nevertheless, the impact of coffee intake on chronic diseases has been a matter of debate in the last two decades, with some conflicting results due the retrospective nature of most of the studies, although coffee has progressively moved to a less negative position due to its better-known phar-
Bioactive Components in Coffee

Coffee includes a complex mixture of compounds. The particular profile of compounds depends on coffee variety, roasting, and processing. Caffeine has been perhaps the most widely known compound and is the most investigated component of coffee. When green coffee beans are roasted at high temperatures, chemical reactions between amino acids and carbohydrates, known as Maillard reactions, create a number of unique compounds. Additionally, coffee is abundant in polyphenols like chlorogenic acids. The main chlorogenic acid in coffee is 5-caffeoylquinic acid, although other caffeoylquinic, feruloylquinic, and dicaffeoylquinic acids are present in significant quantities [5]. Phenolic metabolites of chlorogenic acids have been studied for potential bioefficacy, and controversy still remains as the results are not entirely clear [6, 7]. Only a few studies have investigated the bioavailability of coffee phenolic and chlorogenic acids due to the complex metabolic pathways in humans [8, 9]. Chlorogenic acids may be transformed into phenolic acids (caffeic, ferulic, and isoforulic moieties) and, subsequently, into colonic metabolites (dihydrocaffeic and dihydroferulic acids). With extensive conjugation at the level of the intestine and the liver, many different metabolites (aglycone, sulfate, glucuronide, and methyl) could then be identified from a single cup of coffee. Lactones, diterpenes, including cafestol and kahweol, niacin, and the vitamin B3 precursor trigonelline are also present in coffee (Fig. 1). Cafestol and kahweol found in coffee oil have shown antioxidant activity in cell models and mice models that involved triggering the upregulation of key antioxidant enzymes [10, 11]. On the other hand, the two diterpenes are the main cholesterol-rising compounds in coffee. They are retained in part by paper filters, but are preserved when coffee is directly prepared by boiling the ground beans [12]. Moreover, coffee is rich in vitamin B3, magnesium, and potassium.

Pharmacokinetics and Mode of Action of Caffeine

Caffeine is the most widely consumed behaviorally active substance in the world. It is present in a number of dietary sources, i.e., tea, coffee, cocoa beverages, and chocolate bars as well as soft and energy drinks (Table 1). Caffeine was first extracted from cocoa beans into its purest form, a white powder, in the 1820s by the German Scientist Friedrich Ferdinand Runge. Caffeine is contained in more than sixty plants, which is a remarkable number, thus it has been hypothesized that caffeine was originally a minor nutrient, not essential for the plant, but extremely useful as a pesticide. In fact, caffeine is toxic for several insects and animals, especially herbivores. Through caffeine the plant may defend itself and have a better chance of survival. In this view, caffeine can be considered as a “co-evolutionary protecting agent” [13]. The amount of this natural alkaloid in coffee is influenced by the method of coffee preparation, and amounts between 65 to 120 mg of caffeine have been reported to be contained in a normal cup of coffee, whereas Arabic coffee normally contains less caffeine than the Robusta variety [14]. Soft drinks typically contain about 30 to 60 mg of caffeine per serving. By contrast, energy drinks contain as much as 80 mg of caffeine per serving (Table 1). The caffeine in these drinks either originates from the ingredients used or is an additive derived from the product of decaffeination or from chemical synthesis [15, 16].

Caffeine is completely absorbed by the stomach and small intestine within 45 min of oral ingestion. The hydrophobic properties of caffeine allow its passage through all biological membranes and the peak plasma concentration is reached within 15–20 min after oral ingestion in humans [15]. Caffeine is metabolized in the liver by the cytochrome P450 oxidase system, specifically the CYP1A2 enzyme into three primary metabolites: paraxanthine (84%), theobromine (12%), and theophylline (4%) [17]. Another enzyme involved in caffeine clearance is the NAT2, whose function is to catalyze the transformation of a broad range of xenobiotics [18]. This enzyme has previously been studied with relation to the risk of Parkinson’s disease and in gene-environment interaction studies [19], but with mixed results.

The half-life time of caffeine varies widely among individuals depending on such factors as age, liver function, pregnancy, some

<table>
<thead>
<tr>
<th>ABBREVIATIONS</th>
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<tbody>
<tr>
<td>ALT</td>
<td>alanine aminotransferase</td>
</tr>
<tr>
<td>CD</td>
<td>Crohn’s disease</td>
</tr>
<tr>
<td>CHI3L1</td>
<td>chitinase 3-like protein 1</td>
</tr>
<tr>
<td>CTGF</td>
<td>connective tissue growth factor</td>
</tr>
<tr>
<td>CVD</td>
<td>cardiovascular disease</td>
</tr>
<tr>
<td>CXCL13</td>
<td>small cytokine belonging the CXC family</td>
</tr>
<tr>
<td>CYP1A2</td>
<td>member of the cytochrome P450 oxidase system</td>
</tr>
<tr>
<td>DSS</td>
<td>dextran sulfate sodium</td>
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<tr>
<td>GABA</td>
<td>gamma-amino butyric acid</td>
</tr>
<tr>
<td>GLUT 4</td>
<td>glucose transporter type 4</td>
</tr>
<tr>
<td>GLP 1</td>
<td>glucagon-like peptide 1</td>
</tr>
<tr>
<td>IBD</td>
<td>inflammatory bowel disease</td>
</tr>
<tr>
<td>IL10</td>
<td>interleukin 10</td>
</tr>
<tr>
<td>NAT2</td>
<td>N-acetyltransferase 2</td>
</tr>
<tr>
<td>NF-κB</td>
<td>nuclear factor kappa-light-chain-enhancer of activated B cells</td>
</tr>
<tr>
<td>NHANES</td>
<td>National Health and Nutrition Examination Survey</td>
</tr>
<tr>
<td>TCBQ</td>
<td>tetrachlorobenzoquinone</td>
</tr>
<tr>
<td>TNF-α</td>
<td>tumor necrosis factor-α</td>
</tr>
<tr>
<td>TLR4</td>
<td>toll-like receptor 4</td>
</tr>
<tr>
<td>UC</td>
<td>ulcerative colitis</td>
</tr>
<tr>
<td>WNT10B</td>
<td>member of the WNT gene family</td>
</tr>
</tbody>
</table>
concurrent medications, and the level of enzymes in the liver needed for caffeine metabolism. In healthy adults, caffeine’s half-life time is approximately 3–4 h. In women taking oral contraceptives, this is increased to 5–10 h [20] and in pregnant women the half-life time is roughly 9–11 h [21]. In infants and young children, it may be longer than in adults [15].

The principal mode of action of caffeine at doses in normal human consumption is as an antagonist of adenosine receptors. Adenosine receptors have been cloned, namely, A1, A2A, A2B, and A3. A1 and A3 receptors preferentially couple to Gi proteins and inhibit adenylate cyclase, while A2A and A2B couple to Gs and stimulate production of cyclic AMP (cAMP) [21]. These receptors are widely expressed in the human body and have been implicated in several biological functions, both physiological and pathological. These include cardiac rhythm and circulation, lipolysis, renal blood flow, immune function, sleep regulation, and angiogenesis as well as inflammatory diseases, ischemia-reperfusion, and neurodegenerative disorders [22]. The caffeine molecule is structurally similar to adenosine and is able to potently block adenosine effects on A2A and A1 receptor subtypes already at the low concentration achieved after a single cup of coffee. Twenty times higher concentrations are required to inhibit cyclic nucleotide breakdown via inhibition of phosphodiesterases. To block GABA<sub>A</sub> receptors 40 times higher and to mobilize intracellular calcium stores via activation of ryanodine receptors, 100 times higher concentrations are needed [23]. These high concentrations of caffeine are unlikely to be reached in human by normal use of coffee [15]. For children and young adults, the primary sources of caffeine are soft drinks and teas, while for adults aged 25 and older, it is mostly derived from coffee. Interestingly, people in Asia mainly drink tea instead of coffee.

**Effects of Coffee on the Cardiovascular System**

Considerable controversy exists regarding the association between coffee consumption and CVD risk. The relationship between coffee consumption and the risk of coronary heart disease was first studied in the 1960s, given that the prevalence of coffee drinking and CVD were both high in Western countries [24]. Since 2000, the association between coffee consumption and other CVD outcomes such as stroke, heart failure, and total CVD mortality has also been more frequently studied and summarized in meta-analyses [25–27]. These meta-analyses did not support an association between coffee consumption and a higher CVD risk, but the shape of the association remains uncertain. Interestingly, a meta-analysis published in 2014 [28] concluded that moderate coffee consumption (3–5 cups/day) was associated with a lower CVD risk, and heavy coffee consumption (≥6 cups/day) was neither associated with a higher nor a lower risk of CVD. A further meta-analysis [29] has also showed that heavy coffee consumption was not associated with risk of CVD mortality. In contrast, the cohort study by Liu et al. [30] found that 4 cups per day of coffee consumption was associated with increased mortality, but the association was only significant for participants under 55 years of age. The results from this study contradict those from other meta-analyses and the majority of studies in the literature. Possible reasons for the discrepancy may be a relatively small size, lack of updated dietary assessment, and subgroup analysis in Liu’s study. In addition, coffee brewing methods were not assessed in the included studies.

The question remains whether mechanisms lower the risk of CVD. Pharmacological studies have confirmed that A1 receptor activation has a number of effects in the cardiovascular system, including a reduction in heart rate and atrial contractility, and the attenuation of the stimulatory actions of catecholamines on the heart, and the A2A receptors are involved in vasodilation in the aorta and coronary artery [31]. The blockade of these receptors by caffeine can contribute to the protective effect of coffee in CVD. Apart from caffeine, further studies suggested the involvement of other components confirmed by results from studies with unfiltered and paper-filtered coffee [32] or caffeinated and decaffeinated coffee [33]. Chlorogenic acids and their metabolites, for example, attenuate oxidative stress (reactive oxygen species), which leads to the benefit of blood pressure reduction through improved endothelial function and nitric oxide bioavailability in the arterial vasculature [34]. It is concluded that the available evidence, although limited, allows for stating that there is no clinical basis for associating moderate coffee intake with an increased risk of cardiovascular diseases, including stroke [35]. Not only caffeine but also other components of coffee may contribute to the cardiovascular effects.

**Coffee and Type 2 Diabetes**

Coffee has recently received scientific attention as a current epidemiologic, and *in vivo* studies have revealed its health benefits against metabolic disorders, especially type 2 diabetes [36]. An
inverse correlation has been confirmed in most but not all studies. Therefore, an updated systematic review and a dose-response meta-analysis of all available data on the correlation of both caffeinated and decaffeinated coffee consumption with the risk of type 2 diabetes have been published in 2014 by Ding et al. [37]. The systematic review and meta-analysis based on 1,109,272 study participants and 45,335 cases of type 2 diabetes demonstrate a robust inverse correlation between coffee consumption and the risk of diabetes. Compared with no coffee consumption, 6 cups/day of coffee was associated with a 33% lower risk of type 2 diabetes. The correlation was consistent for men and women [37]. By contrast, a multiethnic cohort study [38] suggested that the protective effect of coffee intake was stronger for women (34% lower diabetes risk) than for men (14% lower diabetes risk). The discrepancy may be due to the coffee intake assessment through self-reported dietary questionnaires. Misclassification, therefore, cannot be excluded. A further aim of Ding’s systematic review [37] was to compare the effects of caffeinated and decaffeinated coffee consumption and the risk of type 2 diabetes. Decaffeinated coffee consumption was associated with the same level of protection as seen for caffeinated coffee and therefore confirmed previous findings of the European Prospective Investigation into Cancer and Nutrition (EPIC)-Germany study [39], which reported a 23% lower incidence for caffeinated and a 30% lower risk for decaffeinated intake of ≥4 cups/day. It remains unclear, however, by which mechanism(s) the diabetes-preventive effect is brought about. There is evidence for both increased insulin secretory responsiveness and increased insulin sensitivity [40]. Coffee appears to act primarily, if not exclusively, through postprandial, as opposed to fasting, glucose homeostasis [41].

Studies observed a protective effect of decaffeinated coffee pointing to a relevant role of constituents other than caffeine, such as polyphenols, which are a major source of antioxidants in the Westernized diet [42, 43]. As seen in knockout mice (Lepr/db), chlorogenic acid inhibited gluconeogenesis by affecting expression and activity of enzyme glucose-6-phosphatase. Moreover, it improved skeletal muscle glucose uptake by increasing expression and translocation of GLUT 4. A 2.5-fold increase in glucose transport was described as an additive action with insulin [44]. Polyphenols in coffee stimulate GLP 1, which is a major intestinal hormone that activates glucose-induced insulin secretion from β-cells [45]. Prolonged activation of the GLP-1 signal has been shown to attenuate diabetes in animals and human subjects. This hypothesis has been confirmed by studies suggesting that polyphenols in coffee bean extract may bring an additive effect in decreasing body weight gain and increasing insulin sensitivity [46]. These beneficial effects are possibly due to the downregulation of genes associated with WNT10B- and galanin-mediated adipogenesis and the TLR4-mediated proinflammatory pathway as well as stimulation of GLUT4 translocation to the plasma membrane in white adipose tissue of mice [47]. So far, coffee has consistently been inversely associated with type 2 diabetes, and polyphenols, among other bioactive compounds, are the best candidates to be responsible for the beneficial actions.

### Coffee and Liver Diseases

There is increasing evidence in favor of protective effects of coffee consumption in the development and progression of liver disease due to various causes. The clinical evidence of benefit of coffee

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**Table 1** Caffeine content chart (according to data from [16]).

<table>
<thead>
<tr>
<th>Item</th>
<th>mg of caffeine</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Typical</strong></td>
<td><strong>Range</strong></td>
</tr>
<tr>
<td>Coffee (240 mL)</td>
<td></td>
</tr>
<tr>
<td>Brewed, drip method</td>
<td>85</td>
</tr>
<tr>
<td>Instant</td>
<td>75</td>
</tr>
<tr>
<td>Decaffeinated</td>
<td>3</td>
</tr>
<tr>
<td>Espresso (30 mL)</td>
<td>40</td>
</tr>
<tr>
<td>Teas (240 mL)</td>
<td></td>
</tr>
<tr>
<td>Brewed, major U.S. brands</td>
<td>40</td>
</tr>
<tr>
<td>Brewed, imported brands</td>
<td>60</td>
</tr>
<tr>
<td>Instant</td>
<td>28</td>
</tr>
<tr>
<td>Iced</td>
<td>25</td>
</tr>
<tr>
<td>Soft drinks (e.g. Cola 360 mL serving)</td>
<td>40</td>
</tr>
<tr>
<td>Energy drinks (250 mL serving)</td>
<td>80</td>
</tr>
<tr>
<td>Cocoa beverage (240 mL)</td>
<td>6</td>
</tr>
<tr>
<td>Chocolate milk beverage (250 mL serving)</td>
<td>5</td>
</tr>
<tr>
<td>Solid Milk chocolate (30 mL serving)</td>
<td>6</td>
</tr>
<tr>
<td>Solid Dark chocolate, semi-sweet (30 mL serving)</td>
<td>20</td>
</tr>
<tr>
<td>Baker’s chocolate (30 mL serving)</td>
<td>26</td>
</tr>
<tr>
<td>Chocolate flavored syrup (30 mL serving)</td>
<td>4</td>
</tr>
</tbody>
</table>

*Due to brewing method, plant variety, brand, formulation etc.
consumption in hepatitis B and C, as well as nonalcoholic fatty liver disease and alcoholic liver disease, has been reviewed by Wadhawan and Anand in 2016 [48] and for hepatic fibrosis and cirrhosis by Liu et al. in 2015 [49]. The two meta-analyses clearly indicated that coffee intake of more than 2 cups per day in patients with preexisting liver disease is associated with a lower incidence of fibrosis and cirrhosis, lower hepatocellular and carcinoma rates, as well as decreased mortality. In favor of this protection, two recent population-based studies, the NHANES I and III [50–52], have reported that higher coffee consumption (> 2 cups per day) was associated with a lower risk of elevated ALT levels by 44% and a lower risk of chronic liver disease compared to non-coffee drinkers. Additionally, a recent large cohort study [53] of 330 patients with alcoholic and non-alcoholic cirrhosis showed a strong inverse relationship between coffee drinking (> 4 cups per day) and elevated serum enzyme levels, especially in those who drank large quantities of alcohol. Moreover, coffee consumption decreased liver stiffness, which may indicate less fibrosis and inflammation in patients with nonalcoholic fatty liver disease, and hepatitis C and B virus infection [54]. Although coffee consumption has been associated with a reduced frequency of liver disease, it is unclear whether the effect is from caffeine or other components. The beneficial effects of caffeine against liver fibrosis have been demonstrated by several studies using standard rodent models of experimental liver fibrosis. In almost every study, ingestion of coffee/caffeine blocked toxin-induced liver fibrosis/cirrhosis [55]. In particular, in experimental models of fibrosis, caffeine was shown to inhibit hepatic stellate cell activation by blocking A2A receptors, and emerging evidence indicates that caffeine might also favorably impact angiogenesis and hepatic hemodynamics. Successively, Gressner et al. [56] have shown that caffeine inhibits TGF-β-induced CTGF expression in hepatocytes. TGF-β levels are reduced by coffee and caffeine administration to rats subjected to chemical-induced liver fibrosis [52, 57, 58]. On the other hand, Vitagliano et al. [59] reported that consumption of decaffeinated espresso coffee was able to reduce not only liver steatosis but also inflammation and fibrosis in rats. It is therefore suggested that caffeine in coffee is not essential and that specific coffee components contribute to the hepatoprotective effect [37, 60, 61]. Chlorogenic acid possesses a hepatoprotective nature [62]. In a recent study [63] conducted on TCBQ-induced liver damage in mice, the acid pretreatment seems to be effective in suppressing TCBQ-induced oxidative stress, therefore possessing a hepatoprotective nature. Additionally, chlorogenic acid reduced liver fibrosis and the expression of collagen I and III. These rats displayed reduced concentrations of VEGF, TGF-β, and α-smooth muscle actin [64]. The diterpenes cafestol and kahweol may offer protective effects against aflatoxin B1-induced liver damage in rats and in hepatocyte cultures [58, 65]. Cafestol and kahweol may also induce the synthesis of glutathione, which has a role in detoxification and the prevention of liver damage. Taken together, a growing body of literature has consistently shown an inverse relationship between coffee consumption and liver diseases. However, there are not enough data to make firm conclusions about the relative importance of caffeine or other components within coffee in the development and progression of liver disease.

Coffee and inflammatory Bowel Disease

Studies to date suggest that there is no association between coffee consumption and the risk of dyspepsia [66], gastroesophageal reflux disease [67], peptic ulcers [68], gastritis, and stomach cancer [69, 70]. People with IBD are coffee users as well, but the question remains as to whether coffee consumption is safe for people living with a chronic digestive disease. IBD consists of two major forms, CD and UC. In a study performed by Ng et al. [71], coffee had a protective effect against UC development. Another report studying 41,836 postmenopausal women for 15 years showed that high coffee consumption is inversely correlated to the severity of inflammatory diseases [72]. It is well known that the herbal mixture of myrrh, dry extract of chamomile flowers, and coffee charcoal has anti-inflammatory and antidiarrheal properties. In a randomized, double-blind, double-dummy study [73], 96 patients with inactive UC were randomized to receive either the herbal preparation or mesalazine over a 12-month period. There was no significant difference in the relapse rate between the two groups. No significant differences were also shown in relapse-free time, endoscopy, and fecal biomarkers. In vitro, the herbal mixture influenced gene expression of activated human macrophages within the cytokine/chemokine signalling pathway. Particularly, chemokine gene expression was suppressed. Subsequently, the production of CXCL13 (which controls the organization of B cells within follicles of lymphoid tissues) and, to a minor extent, cytokine TNF-α were inhibited, whereas IL10 release from activated macrophages was enhanced by coffee charcoal extracts [74]. In vivo, mice treated with caffeine (2.5 mM, equivalent to the concentration of caffeine in 2–3 cups of coffee) displayed a delayed response towards DSS-induced colitis, characterized by lower body weight loss and clinical and histological scores. Bacterial translocation into other organs and proinflammatory cytokine production were also reduced in the caffeine-treated mice with DSS-induced colitis. Caffeine treatment also resulted in the loss of CHI3L1-associated signalling pathway activation [75]. The disease-associated CHI3L1 expression was also observed in colonic tissue samples obtained from CD and UC patients, but was undetected in normal control individuals [76].

It is, however, considered that the effects of coffee are not exclusively due to caffeine but seem to be linked with other specific constituents [77]. Chlorogenic acid displayed a significant anti-inflammatory activity in a well-established mouse model of experimental colitis, as evidenced by a reduction in the macroscopic damage score, myeloperoxidase activity, and inhibition of the NF-κB dependent pathway [78]. Apart from these, the inhibition of cyclooxygenase-2, inducible nitric oxide synthase (iNOS) with the lack of a cytotoxic effect, attenuation of IL-1β and IL-6 along with TNF-α in a dose-dependent manner, and inhibition of NF-κB by chlorogenic acid in a DSS-induced colitis have been reported in a recent study [79].

Although clinical practice guidelines [80] recommend that people with IBD avoid caffeine, there are more clinical and experimental evidences indicating a possible prospective effect of coffee and its components to IBS symptoms or other inflammatory diseases of the gastrointestinal tract.
Summary

To date, there are still many misconceptions about coffee and health that can lead to confusion about whether coffee consumption can be enjoyed as part of a healthy, balanced diet. Therefore, the potential effect of coffee on the risk of many diseases has been studied intensively during the last years, sometimes with contradictory results. The acknowledgment that coffee and caffeine are not equivalent has increased the interest in whether other components of coffee might contribute to the protective action in the human body and, should that be the case, in which sense. In this case, the majority of research has mainly focused on polyphenols. Nevertheless, there is not enough information to answer this question.

It is important to note that individual differences exist in responses to coffee. Some people are more sensitive to the effects than others. Part of such variability is due to tolerance, but there are indications that it might have a genetic basis as well [81]. Another interesting fact is that males and females differ in their responses to caffeine and that these differences may be mediated by changes in circulating steroid hormones [82]. In most people, moderate coffee consumption of up to 4 cups of coffee per day (around 400 mg caffeine) can be enjoyed as part of a healthy, balanced diet and an active lifestyle. Lower levels are recommended for pregnant women who are advised to limit caffeine intake to 200 mg from all sources, as well as in children where the intake should be reduced because of a lower body weight [83]. Finally, there is a need for well-designed, randomized, controlled trials further investigating the effect of different doses of coffee or coffee bioactive components on healthy individuals as well as on patient populations to clarify important points discussed controversially in the literature.

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Conflict of Interest

The author reports no financial interests or potential conflicts of interest.

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