Beyond the Cholesterol-Lowering Effect of Soy Protein: A Review of the Effects of Dietary Soy and Its Constituents on Risk Factors for Cardiovascular Disease

D. Dan Ramdath 1,* , Emily M. T. Padhi 1 , Sidra Sarfaraz 1 , Simone Renwick 1 and Alison M. Duncan 2

1 Guelph Research and Development Centre, Agriculture and Agri-Food Canada, Guelph, ON N1G 5C9, Canada; epadhi@gmail.com (E.M.T.P.); sidra9518@gmail.com (S.S.); srenwick@mail.uoguelph.ca (S.R.)
2 Department of Human Health and Nutritional Sciences, University of Guelph, Guelph, ON N1G 2E1, Canada; amduncan@uoguelph.ca
* Correspondence: dan.ramdath@agr.gc.ca; Tel.: +1-226-217-8082; Fax: +1-226-217-8181

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Abstract: The hypocholesterolemic effect of soy is well-documented and this has led to the regulatory approval of a health claim relating soy protein to a reduced risk of cardiovascular disease (CVD). However, soybeans contain additional components, such as isoflavones, lecithins, saponins and fiber that may improve cardiovascular health through independent mechanisms. This review summarizes the evidence on the cardiovascular benefits of non-protein soy components in relation to known CVD risk factors such as hypertension, hyperglycemia, inflammation, and obesity beyond cholesterol lowering. Overall, the available evidence suggests non-protein soy constituents improve markers of cardiovascular health; however, additional carefully designed studies are required to independently elucidate these effects. Further, work is also needed to clarify the role of isoflavone-metabolizing phenotype and gut microbiota composition on biological effect.

Keywords: cardiovascular disease; cholesterol; functional foods; isoflavones; lipids; obesity; Dietary soy; soy protein

1. Introduction

Cardiovascular disease (CVD) describes a collection of disorders affecting the vasculature of the heart, brain and peripheral tissue and is the leading cause of death globally [1]. Atherosclerosis is the underlying cause of coronary heart disease (CHD), the most common form of CVD, and is thought to be initiated through an inflammatory response by the vascular endothelium following injury [2]. The origin of these endothelial lesions is unclear, but implicated factors include: chronic elevations in blood pressure [3]; prolonged hyperglycemia and the resulting formation of advanced glycation end products [4]; elevated LDL cholesterol (LDL-C), particularly molecules that have undergone oxidized modification [2]; and oxidative stress and inflammation [5]. Consequently, major CVD risk factors include hypertension, the presence of type 2 diabetes, dyslipidemia, obesity (body mass index (BMI) >30), and inflammation [1]. Dietary modification lowers CVD risk by attenuating associated risk factors, and in particular, legumes are emphasized as part of a cardioprotective diet because increased consumption is associated with improved weight management and glycemic control, reduced blood pressure, and an improved plasma lipid profile [6].

Soybeans (Glycine max) are widely cultivated for their lipid content, and indeed are the top oilseed produced worldwide [7]. In addition, soybeans are recognized as a valuable source of nutrients as they contain high-quality protein (~40%); polyunsaturated fatty acids (18%); carbohydrates (primarily
sucrose, stachyose, and raffinose); and dietary fibers [8]. Soy foods have been part of the human diet for millennia, but more recently considerable attention has been given to the associated health benefits of soy, particularly reduction of CVD risk via the lowering of LDL-C. Much of the focus on soy has been directed toward the hypocholesterolemic properties of bioactive peptides in soy protein, which exert their effects primarily through mechanisms involving the LDL-C receptor (LDLR), and bile acid regulation [9,10]. These findings are supported by several meta-analyses [11–21] and have culminated in a soy health claim relating 25 g soy protein with a reduced risk of CHD in the United States [22] and Canada [23], but not Europe [24]. However, other constituents in soy have been shown to confer many health benefits, including reduction of CVD risk, and these are worthy of further examination.

Soybeans are a significant source of phytochemicals such as isoflavones, phytosterols and lecithins, as well as soluble fibers, saponins and polysaccharides, which may act collectively or through independent mechanisms to confer unique health benefits [25–27]. For example, soy lecithins and saponins have a role in lipid metabolism; phytosterols and linoleic acid produce hypocholesterolemic effects [25]; and soy fibers have been shown to promote weight loss [28]. Further, the health benefits of soy protein appear to reach beyond its putative LDL-C lowering effect by offering protection against renal dysfunction [29], oxidative stress [30], and by improving markers of endothelial function [31]. Finally, the health benefits of isoflavones, of which soybeans are the single greatest dietary source, have also been extensively examined. They represent a class of phytoestrogens belonging to the flavonoid family, and there are three primary isoflavones (daidzein, genistein, glycitein) which are characterized by their general di-phenolic structure resembling mammalian estrogen [32]. The structural similarity between soy isoflavones and estradiol suggests that isoflavones may elicit estrogenic effects. Although the affinity for the estrogen receptor by soy isoflavones is 100–1000 times less than mammalian estrogen, isoflavone concentrations can appear in plasma >1000-fold greater than that of endogenous estrogen, thus lending support to the suggestion that isoflavones can exert significant physiological effects [33].

The cholesterol-lowering effect of soy protein has generated much interest in the past and has been extensively studied, but other components in soy appear to confer significant cardiovascular health benefits despite receiving less attention over the years. This review serves to examine and expand on the health effects of soy and soy constituents beyond cholesterol reduction. Herein, we provide an up-to-date summary of the epidemiological and clinical evidence associating dietary soy, and other soy-derived components, with reduced CVD risk. This review will examine the cardio-protective effects of dietary soy in the context of major disease outcomes such as hypertension, hyperglycemia, dyslipidemia, inflammation, and obesity.

2. Epidemiological Studies

Current epidemiological findings show an inverse association between consuming whole soy foods/products and CVD risk. For example, Shimazu et al. (2007) studied Japanese dietary patterns and found that increased soybean intake (up to 101 g/day) was associated with lower CVD mortality [34]. However, the recent Takayama study, which examined dietary intakes of soy and natto (fermented soy beans) and CVD mortality among Japanese adults, found that there was a significant decrease in mortality from stroke at the highest quartiles of total soy protein and natto intake [35]. Despite this finding, the authors conclude that except for natto, there were no significant associations between CVD related mortality and intakes of total soy protein, total soy isoflavone, and soy protein or soy isoflavones from soy foods [35]. It appears that different soy foods may vary in their biological efficacy and protective effects. In this regard, it is worthwhile to examine the role of soy isoflavones, given that their estrogenicity may protect against the sharp rise in CVD incidence after menopause, when endogenous estrogen concentrations are depleted [15]. Although plausible mechanistic evidence supports the cardio-protective effects of soy isoflavones, there is discrepancy in
the literature regarding the health effects of isoflavones extracts that are consumed in isolation of other soybean components, and this has created disagreements about their health benefits.

Observational studies among East Asian populations consistently find that chronic disease incidence is lowered with the intake of isoflavone-containing whole soy and traditional soy-based foods [36]. In contrast, studies performed with Western populations frequently utilize various combinations of isolates to evaluate the health effects of soy, and have failed to consistently demonstrate health benefits [36]. Furthermore, there is emerging evidence on the diverging biological effects among individuals capable of metabolizing the soy isoflavone daidzein to the more bioactive metabolite equol, which suggests a complex interplay between gut microbiota composition and host benefit, perhaps accounting for some of the conflicting results in the literature [37]. For example, in exploring the effect of soy on CVD risk factors, it was found that blood pressure, plasma triglycerides (TG) and C-reactive protein (CRP) concentrations were significantly lowered among equol- and O-desmethylangolensin (ODMA)-producing pre-hypertensive, postmenopausal Chinese women, compared to non-producers [38]. In addition, racial differences in the ability to convert daidzein into equol have been reported. For example, 30%–40% of Western populations are estimated to be equol producers while greater proportions have been observed in Japanese, Chinese, and Korean populations, which are purportedly linked to differences in genetics, gut microbiome composition, and perhaps diet [39]. Interestingly, background diet did not influence isoflavone bioavailability in Chinese Asian adults who were acclimatized to a French Western dietary pattern [40].

Other health benefits observed among soybean consumers include lower BMI and mortality due to CHD [41], while lower circulating inflammatory markers are observed among female soy consumers [42]. Soy consumption has also been associated with the lowering of blood pressure. In the US, a pooled analysis of three longitudinal cohort studies (Nurses’ Health Study, Nurses’ Health Study II and Health Professionals Follow-up Study) found that consuming ≥4 servings/week of broccoli, carrots, and tofu/soybeans was associated with a lower risk of hypertension compared to consuming <1 serving/month [43]. However, the Tehran Lipid and Glucose Study, which involved 1546 normotensive adults, did not find an association between incident hypertension and dietary phytochemical intake, which provided an indirect estimate of soybean consumption [44].

Soy intake may also lower CVD risk by mitigating risk factors for type 2 diabetes (T2D), although this protective effect appears to be limited to populations living in East Asia. In the Saku Study, which stratified participants by sex and BMI, Japanese men with a higher BMI (>23.6 kg/m²) who consumed soybean products ≥4 servings/week experienced lower fasting and postprandial blood glucose concentrations, and lower T2D incidence compared to those consuming <1 serving/week [45]. Additionally, higher intake of soy products, daidzein and/or genistein was associated with lower T2D risk among overweight Japanese women [46] and adult Chinese Singaporeans [47]. By contrast, intake of soy foods (tofu/soy milk) among US adults does not appear to lower T2D risk, although an inverse relationship between soy isoflavone intake and T2D risk was identified in one study [48]. Similarly, a multi-ethnic cohort study conducted in Hawaii did not find an association between soy food intake and diabetes risk among Caucasian, Japanese American, and Native Hawaiian adults [49]. More recently, a meta-analysis was unable to establish a relationship between soy intake and risk of stroke or CHD, although the authors reason that this may be attributed to the limited pool of case-control and cohort studies currently available [50]. Taken together, population-based studies suggest soy intake mitigates CVD risk factors, but it is unclear whether Western populations gain similar health benefits observed among populations living in East Asia. The results of observational studies conducted in Western populations often lack rigor and are controversial, quite possibly because the active components of soy-based interventions are not adequately characterized. Additionally, it is possible that there are coexisting dietary and environmental factors in East Asian cultures that confound the outcome of these studies.
3. Interventional Studies Involving Soy

3.1. Blood Pressure Lowering Effect of Soy

Hypertension is defined as systolic blood pressure (SBP) >140 mm-Hg and/or diastolic blood pressure (DBP) >90 mm-Hg [51]. Primary hypertension accounts for 90% of all cases and although the underlying cause is unclear, the major contributing factors include diet, smoking, stress, obesity [51], and possibly genetics [52]. Hypertension increases the risk of vascular injury through pro-inflammatory mechanisms, thereby increasing CVD risk. Further, angiotensin II, a potent vasoconstrictor, is elevated during hypertension and can increase the activity of the free radical-generating lipooxygenase enzyme in smooth muscle cells, which contributes to the formation of oxidized LDL-C (oxLDL), superoxide anion, and hydroxyl radicals [2]. Soy and some of its constituents have been shown to reduce risk for hypertension through effects on vasodilation and inhibition of a key enzyme involved in the regulation of blood pressure.

Isoflavones have been shown to mitigate hypertension by targeting mechanisms involving vasodilation; in particular, interaction with the estrogen-response element of genes related to endothelial nitric oxide (NO) synthase increases endogenous NO production, which improves brachial artery flow [53]. In postmenopausal women, six months of isoflavone supplementation was shown to improve endothelial vasodilation and resulted in a significant reduction in cellular adhesion molecules such as Intercellular Adhesion Molecule 1, Vascular Cell Adhesion Protein 1, and E-selectin [54]. In support of these findings, animal studies have shown that soy isoflavones increase renal blood flow, sodium excretion, and interact with estrogen receptors to inhibit angiotensin converting enzyme activity in the renin-angiotensin-aldosterone system [55]. However, clinical evidence supporting a role for isoflavones in hypertension management remains controversial. A meta-analysis of 14 randomized controlled trials (RCTs) found that daily intake of isoflavone extracts (25–375 mg) over a period of 2–24 weeks significantly decreased SBP (−1.92 mm-Hg, 95% CI: (−3.45 to −0.39 mm-Hg)), but not DBP, in normotensive adults [56]. Another meta-analysis found that daily intake of soy isoflavones (65–153 mg) for 1–12 months significantly lowered blood pressure (SBP: −5.94, 95% CI, (−10.55, −1.34 mm-Hg); DBP: −3.35, 95% CI, (−6.52, −0.19 mm-Hg)) in hypertensive, but not normotensive, adults [57]. This suggests that the hypotensive effect of isoflavones is best achieved in persons with established hypertension. Recently, a meta-analysis of 71 trials investigating the effect of phytoestrogen supplementation on arterial hypertension concluded that reductions in SBP and DBP were not significant [58].

Several studies suggest that the effect of isoflavones on endothelial function may be related to individual capacity to metabolize daidzein into equol [59]. Recently, a double-blind crossover study demonstrated significant improvements in arterial stiffness, blood pressure, and endothelial function with the consumption of purified equol supplements, but only among equol-producing men [60]. Another study showed that soy nuts, which provide a source of both soy protein and isoflavones, attenuated SBP in both healthy women and those with metabolic syndrome (MetS), but DBP was significantly decreased only in participants identified as equol-producers [61]. In equol-producing women with prehypertension or untreated hypertension, neither whole soy nor purified daidzein caused significant changes in 24-h ambulatory blood pressure after six months [62].

Apart from isoflavones, other components of soy have been shown to possess hypotensive properties. Soy pulp, which contains oligopeptides and high amounts of fiber, has exhibited anti-angiotensin-converting enzyme activity in vitro, providing mechanistic evidence for a hypotensive effect [63]. Previously published systematic reviews have indicated difficulty in ascertaining the effect of soy protein on blood pressure, citing a lack of studies that focus solely on the protein component of soy as the active agent [31,64]. Soy protein consumed in combination with isoflavones appears to lower SBP in pre-diabetic, postmenopausal, hypertensive women when compared to milk protein [65,66], and a recent meta-analysis showed that soy protein lowers DBP in patients with T2D and MetS [67]. In another study, nut consumption was associated with lower SBP and DBP in adults without T2D;
however, subgroup analysis revealed that soy nuts alone did not reduce blood pressure [68]. Further, a novel bread fortified with soybean flour did not improve blood pressure in women with T2D [69], and soy lecithin with and without isoflavone-rich soy protein isolate did not ameliorate brachial artery flow-mediated dilation, although improvements in the plasma lipids profile were observed [70]. The amino acid composition of dietary soy is another interesting feature that may help explain its associated hypotensive effects. Soy foods and legumes in general, are rich in arginine which is a precursor to NO in the L-arginine-nitric oxide pathway [71]. It is thought that arginine aids in regulation of blood pressure through increased production and improved NO bioavailability in the vascular endothelium [72]. Two meta-analyses have concluded that supplementation with L-arginine significantly improves blood pressure and endothelial function in adults [72,73]. Additionally, soy foods have increased arginine content relative to lysine, which may impact the hypotensive activity of soy containing foods. Both amino acids compete for the same transporter in the intestinal lumen, so increased lysine relative to arginine could limit uptake of the latter, and thus affect its bio-conversion and consequent downstream hypotensive effects [74]. Table 1 shows the arginine and lysine contents of soy protein isolate relative to other protein sources.

Overall, although soy isoflavones seem to attenuate blood pressure, this effect is more likely to occur in hypertensive or equol-producing individuals. The independent and/or combined hypotensive effects of other soy components such as soy protein and its amino acid composition, fiber, lecithins, and saponins require further studies as the current body of evidence is limited by that number of available RCTs, but currently does not suggest any significant hypotensive effects.

### Table 1. Arginine and lysine content of select foods.

| Food                   | Arginine | Lysine | PDCAAS |
|------------------------|----------|--------|--------|
| Soy protein isolate    | 6670     | 5327   | 1.00   |
| Peanuts, dry-roasted   | 2832     | 850    | 0.52   |
| Almonds, dry-roasted   | 2444     | 563    | 0.23   |
| Black beans, boiled    | 549      | 608    | 0.75   |
| Milk, 2% (cow)         | 94       | 276    | 1.00   |
| Egg, hard-boiled       | 755      | 904    | 1.00   |
| Beef, chuck, braised   | 2054     | 2748   | 0.92   |

1 Adopted from the US Department of Agriculture National Nutrient Database [75], and expressed as mg/100 g edible portion; 2 The Protein Digestibility Corrected Amino Acid Score (PDCAAS) is the WHO preferred method for evaluating protein quality (scored numerically, 0–1) on the basis of amino acid composition relative to human requirements [76]. Scores adopted from previously published work [77–79].

### 3.2. Blood Glucose Lowering Effect of Soy

Hyperglycemia is defined as chronically elevated fasting blood glucose (FBG) >6.1 mmol/L that typically results from abnormal glucose metabolism and impaired insulin sensitivity [80]. Greater CVD incidence among diabetic patients suggests that hyperglycemia is associated with an increased CVD risk, perhaps by exacerbating atherosclerotic lesions through the generation of advanced glycation products [4,81]. Murine models of T2D have shown that supplementation with soy isoflavones lead to favorable effects, such as reduced islet β-cell loss and increased antioxidant enzyme activity [82–84]. In particular, genistein has been shown to possess antioxidant activity and inhibit tyrosine kinase, which collectively may alter insulin secretion. However, quite independently, genistein appears to directly affect β-cell proliferation, glucose-stimulated insulin secretion and confers protection against apoptosis through mechanisms that involve cyclic AMP/Protein Kinase A (cAMP/PKA) signaling as well as epigenetic regulation of gene expression at physiologically relevant doses [85]. Phenolic-rich extracts from soybeans also appear to inhibit α-amylase and α-glucosidase enzymes in vitro, potentially aiding in the regulation of postprandial blood glucose [86]. These studies demonstrate plausible mechanistic evidence supporting the role of soy isoflavones in improving glycemic control, and are supported by human trials that show soy isoflavones can significantly alter
glucose homeostasis. For example, 120 postmenopausal Caucasian women with MetS experienced significant improvements in FBG, insulin, and insulin sensitivity after consuming 54 mg of purified genistein daily for one year [87]. Chinese women given a calcium supplement containing 40 or 80 mg soy isoflavones experienced significant reductions in FBG over the course of one year compared to a group given a calcium supplement alone [88]. In addition, FBG, insulin, and insulin resistance decreased in postmenopausal women at risk for osteoporosis following long-term supplementation with genistein extracts [89]. However, other markers of glycemic control, such as 2-h postprandial glucose and HbA1c did not improve when Chinese women consumed soy protein with or without intact isoflavones [90], or when 50 mg of daidzein or genistein supplements were ingested daily for 24 weeks [91]. A meta-analysis that examined the effect of soy isoflavones supplementation in peri- and postmenopausal non-Asian women found significant treatment effects, including reduced circulating insulin (−1.37 μIU/mL, 95% CI: (−1.92 to −0.81 μIU/mL)) and insulin resistance as measured by the Homeostatic Model Assessment for Insulin Resistance (HOMA-IR) (−0.39, 95% CI: (−0.65 to −0.14)) but not FBG [92]. A more recent meta-analysis concluded that isoflavones, particularly genistein, significantly lowered FBG (−0.22 mmol/L, 95% CI (−0.38 to −0.07 mmol/L) and insulin resistance (HOMA-IR, −0.52, 95% CI: (−0.76 to −0.28) in postmenopausal women, although substantial heterogeneity between studies was noted [93]. Another recent meta-analysis of individuals with T2D and MetS showed significant reductions in FBG and insulin concentrations, compared to placebo, when soy protein was consumed for ≥6 months [94].

Closer examination of the available studies suggests that improvements in glycemic control are achieved more frequently with soy foods rather than isolates. For example, a meta-analysis of 24 randomized controlled trials found no significant improvements in FBG, insulin, or HbA1c with the consumption of soy protein with or without isoflavones; however, a subgroup analysis showed that whole soy, but not purified isoflavones significantly reduced hyperglycemia [95]. Among older women with MetS, significant reductions in FBG, insulin, and HOMA-IR were experienced with soy nuts but not textured soy protein [96]. A similar finding was made in a study that tested the effect of soy nuts or soy protein isolate on blood glucose response in postmenopausal women with MetS [97]. In persons with T2D, blood glucose and insulin responses were lowered in those consuming a nutrition bar made from soybeans compared to an isocaloric cookie control [98], and whole soy powder significantly reduced the glucose, but not insulin, response compared to white rice [99].

The diverging effect of whole soy foods compared to soy protein isolates suggests that other components (e.g., fiber, saponins, polysaccharides, phytosterols) may account for the hypoglycemic effect of soy [95]. Indeed, soluble fibers extracted from soy hulls have in vitro binding capacity and physicochemical properties (solubility, viscosity, water-holding capacity) similar to oat β-glucan, a component of oat fiber with well-documented cholesterol and glucose-lowering properties [100]. In a randomized crossover study, soluble polysaccharides extracted from soy did not alter postprandial blood glucose response in healthy young males; however, an inverse relationship was found between product viscosity (but not fiber concentration), glucose area under the curve (AUC), and glycemic index [101]. Currently, a lack of human studies examining the hypoglycemic effect of soy components other than isoflavones and protein makes it difficult to draw conclusions on the effects of these components. Additional studies are therefore required in order to better clarify the role of these bioactive agents in potentiating the hypoglycemic effect of soy. A summary of findings from randomized control trials based on the impact of soy on hyperglycemia is presented in Table 2.
Table 2. Summary findings from randomized control trials based on the impact of soy on hyperglycemia.

| Study | Sample Size & Population | Duration | Treatment | Dose; Format | Results | Conclusions |
|-------|--------------------------|----------|-----------|--------------|---------|-------------|
| [89]  | 389 osteopenic postmenopausal women | 2 years, parallel RCT | Genistein extract with calcium and Vitamin D3 supplement compared to placebo | Purified supplement (54 mg/day) | Significant reductions in FBG, insulin, HOMA-IR, fibrinogen, F2-isoprostanes, soluble intercellular adhesion molecule-1, and soluble vascular cellular adhesion molecule-1 | Genistein supplements improve markers of glycemic control and other markers of CVD risk |
| [97]  | 42 postmenopausal women with MetS | 8 weeks, randomized crossover trial | Control diet (Dietary Approaches to Stop Hypertension, DASH); DASH diet with red meat replaced with soy nuts; DASH diet with red meat replaced with soy protein | 30 g soy nut = 1 serving red meat; 30 g soy protein = 1 serving red meat. Normal diet for 3 weeks followed by all 3 diets for 8 weeks each, with 4-week washout period in between each diet | Soy nuts reduced HOMA-IR and fasting plasma glucose more than soy protein \((p < 0.01 \text{ for both factors})\) and control \((p < 0.01 \text{ for both factors})\) | Soy nuts have greater role in attenuating blood glucose response markers compared to soy protein in postmenopausal women with MetS |
| [96]  | 75 women with MetS, age 60–70 | 12 weeks, parallel RCT | Soy nuts; textured soy protein (TSP) | 35 g/day soy nut; 35 g/day textured soy protein | Compared to control, serum FBG, HOMA-IR and insulin were lower in soy-nut group \((p < 0.05, p < 0.1, p < 0.05)\) and the mean changes were higher in soy nut group vs. TSP \((p < 0.001)\) | Soy nut leads to greater reductions in markers of glycemic control than soy protein |
| [88]  | 203 postmenopausal women, age 48–62. | 1 year parallel RCT | Calcium tablet with isoflavones | 40 or 80 mg isoflavones/day compared to calcium with 0 mg isoflavones | Both treatment groups significantly lowered FBG. No dose-response effect. No effect on lipids | Isoflavone supplementation favourably influences FBG in postmenopausal women |
| [90]  | 180 postmenopausal women with hyperglycemia | 6 months parallel RCT | Soy protein isolate with or without isoflavone conjugates | 15 g soy protein and 100 mg isoflavones, 15 g milk protein and 100 mg isoflavones, 15 g milk protein | No significant improvements in fasting and 2-h postload glucose, fasting and postload insulin, glycated serum protein, HOMA-IR, and beta-cell function | Soy protein isolate with or without isoflavones does not improve glycemic control and insulin sensitivity |
### Table 2. Cont.

| Study | Sample Size & Population | Duration | Treatment | Dose; Format | Results | Conclusions |
|-------|--------------------------|----------|-----------|--------------|---------|-------------|
| [99]  | 20 healthy males and females | 1 day; within-subject design with 1 week intervals in between interventions | Intervention (1) Bar-type cake made of whole soy powder; (2) cooked paddy-rice; (3) cooked paddy rice with whole soy powder cake | 114 g whole soy powder cake containing 50 g carbohydrates, 144 g cooked paddy-rice containing 50 g carbohydrates; 144 g cooked paddy-rice with 60 g whole soy powder sake | (1) Blood glucose and insulin levels were lower than control (2) Blood was lower, while insulin levels were increased slightly | Postprandial blood glucose and insulin response may be improved with whole soy powder food; beneficial effects of combining soy products with carbohydrate-rich foods are less clear |
| [87]  | 120 postmenopausal women with Metabolic Syndrome (60 received treatment; 60 received placebo) | 1 year, randomized, double-blind, placebo-controlled trial | Genistein | 54 mg/day in 2 tablets | Fasting blood glucose, fasting insulin and HOMA-IR were significantly decreased in the treatment group ($p < 0.001$) | Soy isoflavone, genistein, significantly lowers hyperglycemia and reduces insulin resistance |
| [98]  | 10 type 2 diabetes patients in the 80 kcal meal tolerance test (Study 1); 11 diabetic patients in the 592 kcal meal tolerance test (Study 2) | 1 day, crossover study | Soybean nutrition bar | Study 1: 1 soybean nutrition bar containing 7.0 g carbohydrates, 4.3 g fat, 1.9 g fiber, 2.7 g protein, 8.2 mg isoflavones; Study 2: 4.3 soybean nutrition bars, each containing 50.9 g carbohydrates, 31.3 g fat, 14.4 g fiber, 19.6 g protein, 14 mg isoflavones | Blood glucose was lower in both studies with the soybean nutrition bar intervention ($p < 0.001$); Insulin AUC was lower than the control in study 2 only; no significant changes in blood TGs and non-esterified fatty acids between treatment and control groups in either study | Soybean nutrition bars may have a role in preventing postprandial hyperglycemia in patients with type 2 diabetes compared to isocaloric cookies |
| [91]  | 165 women with impaired glucose regulation, aged 30–70 | 24 week parallel RCT | Soy protein with or without daidzein or genistein extract | 10 g soy protein, or protein with 50 mg daidzein, or 50 mg genistein | No significant changes in FBG, 2-h glucose, HbA1c, fasting, and 2-h insulin, AUC of glucose and insulin. | Purified extracts of daidzein and genistein do not improve glycemic control and insulin sensitivity |

AUC = area under blood glucose response curve; CVD = cardiovascular disease; FBG = fasting blood glucose; HBA1c = glycated hemoglobin; HOMA-IR = homeostatic model assessment of insulin resistance; RCT = randomized controlled trial; MetS = metabolic syndrome.
3.3. Non-Protein Effects of Soy on Blood Lipids

Studies examining the relationship between soy and dyslipidemia, defined as elevated plasma LDL-C and TG concentrations and often accompanied by low HDL-C [102], have highlighted differences in the efficacy between soy isoflavones and protein in reducing LDL-C. In many human trials, a hypolipidemic effect has been demonstrated with soy isoflavones; however, this effect appears to be inconsistent so there is wide disagreement in the literature. Isoflavones are believed to exert their hypolipidemic effects by binding with estrogen receptors when circulating estrogen is low, after which they translocate to the nucleus, bind to DNA sequences near the promoter region of target genes, and induce DNA transcription [103]. Through this mechanism, isoflavones could serve as ligands for lipid-regulating proteins such as the peroxisome proliferator activated receptors (PPARs), the liver X receptor, and the farnesoid X receptor, which would lower hepatic lipid synthesis, bile acid synthesis, and cholesterol reabsorption [104]. As such, there is a mechanism by which soy isoflavones could theoretically modulate circulating cholesterol levels. However, isoflavone supplements taken by postmenopausal women for 12 weeks did not alter the expression of genes associated with the LDLR and scavenger receptor CD36, both of which are important in regulating plasma LDL-C concentrations [105]. Further, no significant changes in body fat and visceral adipose tissue were detected; in fact, a potentially deleterious significant increase in LDL-C was observed [105]. This finding agrees with a meta-analysis which found that an average of 70 mg/day purified isoflavone aglycone extracts (27–132 mg/day), consumed independently of soy protein, did not significantly lower total or LDL-C in normocholesterolemic menopausal women [18]. Another meta-analysis found that soy products, but not isoflavone supplements, significantly improved total cholesterol (TC), LDL-C, HDL-C, and TG [102]. It seems reasonable to conclude that current evidence from randomized control trials in humans do not sufficiently support the claim that isoflavone extracts can independently reduce CVD risk by modulating plasma lipids. It is possible that soy isoflavones may reduce CVD risk by protecting against the oxidation of LDL-C and the development of oxLDL, as opposed to a lipoprotein-lowering effect. However, preliminary studies indicate that conjugated isoflavone metabolites (the abundant form in circulation) are ineffective antioxidants in comparison to aglycone isoflavones [106].

The hypolipidemic effect of soy may be mediated through a synergistic interaction between its proteins and isoflavones. Animal studies have shown that serum lipids are augmented only when soy protein is consumed in combination with isoflavones [107,108]. A meta-analysis that examined the effect of soy protein consumed with varying doses of isoflavones found a lowering effect on TC, LDL-C, and TG, in addition to increased HDL-C among individuals with T2D [109]. There is also evidence suggesting that soy phytochemicals may elicit hypolipidemic effects. For example, hamsters that were fed a high-fat diet along with 200 mg/kg glyceollins (derivative molecules of isoflavones) experienced reduced plasma lipids and hepatic lipid content after 28 days of supplementation [110]. In rats, four-week supplementation with okara, a tofu by-product containing high amounts of protein and insoluble dietary fiber, resulted in lower serum TG and phospholipid concentrations [111]. Furthermore, it has been suggested that soy lipids such as lecithins and phospholipids could also potentiate the hypolipidemic effect of soy by inhibiting intestinal cholesterol absorption and promoting biliary cholesterol excretion [112]. High concentrations of phosphatidylcholine in soy lecithin may account for these modulatory effects on lipid metabolism by solubilizing cholesterol in the intestines, thereby restricting uptake by enterocytes [112]. An early study demonstrated that soybean phospholipid supplementation exerted direct effects on plasma lipids, producing a significant reduction in TC during supplementation, which was followed by a significant increase in cholesterol concentrations when the intervention was stopped [113]. Another study showed that a powder containing soy-derived stanols and lecithins lowered total- and LDL-C in a 10-week RCT of normo- and mildly hypercholesterolemic adults by reducing cholesterol absorption [114].

The combination of soy fibers and phospholipids with soy protein was found to have an additive hypocholesterolemic effect compared to when soy protein was consumed alone [115]. A low-glycemic
index diet supplemented with soy protein and 4 g of soy phytosterols induced hypolipidemic effects in postmenopausal women, improved blood pressure, and lowered the Framingham risk score assessment for CHD in postmenopausal women [116]. Most recently, soy milk powder enriched with phytosterols was shown to significantly lower TC and LDL-C in mildly hypercholesterolemic Chinese adults after six months of daily intake; this effect was independent of the apolipoprotein E genotype [117], suggesting the observed effect was not attributed to inherent differences in cholesterol uptake. However, other factors such as the gut microbiota composition could mediate the cholesterol lowering effect of soy. For example, a recent study demonstrated increased gut microbial diversity in hamsters fed soy protein compared to dairy protein, which was related to an observed lipid lowering effect [118]. It is therefore apparent that the lipid lowering effect of soy is potentiated and mediated by inherent factors as well as constituents that act in synergy with protein.

3.4. Effects of Soy on Inflammation and Obesity

Inflammation is an innate immune response involving both intra- and extracellular pathways that utilize growth factors, cytokines, leukotrienes, and prostaglandins to attack foreign substances and eliminate them. In this regard, the development of atherosclerotic plaque is one such event that positively stimulates the immune response and results in a state of chronic inflammation [119]. Chronic inflammation can further lead to pathologies such as CVD by encouraging the transformation of healthy endothelial tissue into diseased tissue [119]. Available evidence supports an important role for soy isoflavones in mitigating markers of inflammation. For example, the isoflavone genistein protects against endothelial injury by down-regulating the expression of pro-inflammatory genes and inhibiting the production of reactive oxygen species (ROS) [120]. Studies in cell culture models have shown that genistein, but not daidzein, can mitigate cellular damage induced by peroxidase via an increase in glutathione peroxidase activity [121]. Daidzein has been shown to regulate the expression of inflammatory genes by modulating pathways involved in the activation of PPAR-α and -γ, and by inhibiting the c-Jun N-terminal kinase (JNK) pathway [122]. These findings are supported by preliminary human studies testing the independent effects of soy isoflavones. For example, a clinical trial involving prostate cancer patients showed that isoflavone-fortified bread lowered pro-inflammatory cytokines and chemokines [123]. In addition, increased isoflavone intake appeared to confer some protective effects when healthy young adults were challenged with an endotoxin to evoke an inflammatory response [124]. Further, isoflavone extracts taken by obese postmenopausal women improved serum concentrations of leptin, adiponectin, and tumor necrosis factor alpha (TNF-α) after supplementation for six months [125]. However, a six-month RCT involving 265 postmenopausal equol-producing Chinese women showed that significant improvements in plasma CRP concentrations were achieved only when whole soy, but not purified daidzein, was consumed [65].

Studies involving human volunteers have shown that soy protein also confers anti-inflammatory effects. For example, adults with T2D and nephropathy who consumed soy protein had significant reductions in serum CRP [126]. In addition, hemodialytic patients consuming 27 g/day of soy protein for six months had lower ratios of neutrophil:lymphocyte concentration, a marker of systemic inflammation [127], and a diet supplemented with soy nuts improved arterial stiffness in adults at risk for CVD [128]. The reasons for these anti-inflammatory effects are unclear; however, it has been postulated that the amino acid composition of soy protein may partly account for its ability to mitigate systemic inflammation [129]. Notably, glycine (an amino acid found abundantly in soy protein) was shown to promote antioxidant enzyme activity and inhibit inflammatory pathways in a rat model [130]. A particular challenge in interpreting the literature on soy and inflammation is that markers used to assess inflammation vary by study [65]. Moreover, several studies are conducted with postmenopausal women, which limit the generalizability of these results. Together, these limitations create difficulty in ascertaining the net effect of soy and its constituents using pooled analysis.

Obesity, now recognized as a state of disease, is closely linked with inflammation as excess adipose tissue (especially visceral adipocytes) secrete inflammatory cytokines and chemokines that can initiate
and/or promote a pro-inflammatory state [131]. Obesity is also linked to an increase in circulating ROS, which can damage proteins and cellular organelles such as the mitochondria [132,133]. Additionally, low adiponectin concentrations, which are observed during obesity-associated inflammation, promote the development of insulin resistance, MetS, and CHD [131]. Given the epidemiological evidence linking soy consumption to healthier body weights [47], the specific mechanisms underlying this health benefit have been an active area of recent research.

Soy isoflavones are associated with anti-adipogenic effects, although evidence on this relies largely on data obtained from animal studies. For instance, long-term supplementation with isoflavones was shown to reduce body/visceral adipose tissue and serum leptin concentrations in adult female rats [134], and isoflavone extracts reduced body mass and plasma lipid concentrations in diet-induced obese male rats [135]. Another study using Huanjiang mini-pigs found that isoflavones regulated the expression of genes involved in lipid metabolism [136]. Similarly, mice consuming isoflavones extracts (0.15%) experienced reduced body weight, adipose mass, and TG concentration, possibly through the suppression of genes regulating PPAR-γ and SREBP-1c [67]. Isoflavone-induced upregulation of PPAR-α and PPAR-γ coactivator-1α (PGC-1α) was also postulated to promote the breakdown of fatty acids by inducing β-oxidation [137]. Controversially, some isoflavone studies have produced evidence that suggests isoflavone supplementation could have deleterious consequences: increases in adipose tissue were observed among male mice fed a low-fat diet supplemented with purified genistein [138], and TC and leptin concentrations significantly increased when mice were supplemented with 0.45% isoflavone extracts [139]. Further studies have demonstrated other benefits such as reduced body and liver mass, blood glucose, TC, and serum leptin concentrations in mice fed a high-fat diet supplemented with black soybean [140].

Studies examining the effect of soy on weight loss in obese humans are limited. In one of these studies, obese postmenopausal women ingesting 75 mg isoflavone conjugates daily experienced reductions in trunk fat mass after 12 months of supplementation [141]. In contrast, a multi-center dose-response study showed that obese postmenopausal women who ingested soy isoflavone tablets containing 80 or 120 mg isoflavone conjugates daily did not experience significant improvements in body composition [142]. However, in adults (18–95 years), obesity was associated with the status of ODMA non-producer, which suggests additional work is required to clarify the role of isoflavone-metabolizing-phenotype on the bioactivity of soy isoflavones consumed during intervention studies [143].

The effect of other soy components on obesity has also been examined. One study found that biscuits fortified with soy fiber significantly lowered body weight, BMI and total- and LDL-C in healthy adults [28], although a similar effect was not observed when obese participants with non-alcoholic fatty liver disease consumed a low-carbohydrate diet supplemented with soy nuts [144]. Participants enrolled in a weight loss exercise and nutrition program experienced greater, but not statistically significant, weight loss when assigned to a whole soy group compared to the wheat supplemented group despite engaging in equal amounts of exercise [145].

The efficacy of soy protein in improving body composition has been examined in a few human trials and the overall results suggest no intervention effects; moreover, it does not appear to be an effective weight loss aid. For example, postmenopausal obese women consuming a calorie-restricted diet supplemented with soy protein did not lose significantly more weight compared to a diet without soy [146]. Another RCT revealed that soy protein was not as effective as milk protein in lowering visceral and subcutaneous fat mass [147], and postmenopausal women did not experience significant reductions in BMI or waist-to-hip ratio after consuming isoflavone-rich soy protein daily for 12 months [148]. However, the addition of black soy protein led to more significant declines in body weight, body fat mass and leptin in 64 overweight/obese participants [149], and in a 12-month exercise and nutrition intervention study, obese women lost significantly more body weight with a soy yogurt meal replacement compared to control [150].
4. Contribution of Soy Based Foods to Satiety

The effect of consumption of soy and soy products on satiety-related measures has been explored in a few studies. In one such study, soy isoflavone supplementation did not significantly influence energy intake, body weight, nor serum ghrelin concentrations in healthy postmenopausal women after eight weeks [151]. Similarly, isoflavone tablets did not significantly affect plasma concentrations of insulin, leptin, ghrelin, and adiponectin in postmenopausal women after 12 months [142]. In addition, soy flour (27.3%) incorporated into a pretzel-like bread did not significantly alter satiety scores [152], and there were no significant differences in appetite, satiety, or food consumption when identical portions of beef or soy protein were consumed [153]. Similarly, obese men consuming a soy-based, high-protein diet did not experience more weight loss, nor was food intake reduced compared to men consuming a diet with beef [154]. Overweight and obese men who consumed an isoflavone-free soy protein preload 30 min before lunch had reduced appetite and caloric intake after 12 weeks, but whey protein exhibited more substantial effects [155]. However, muffins made with soy flour elicited greater feeling of fullness scores among mildly hypercholesterolemic overweight adults, compared to a wheat muffin supplemented with whey protein [156]. In another study, snacks made with soy protein isolates led to greater reductions in appetite and reduced intake of sugary snacks in the evening among adolescents [157], and a breakfast high in soy protein reduced serum appetite hormones including ghrelin and protein YY [158]. Obese mice also exhibited greater satiety while consuming whey protein as compared to soy protein [159]. Altogether, the limited evidence on the effect of soy and satiety appears to suggest that soy protein, isoflavones, and other soy constituents do not elicit a satiating effect any greater than what is observed with a comparable quantity of animal protein. A summary findings from randomized control trials based on the impact of soy on obesity and satiety is presented in Table 3.
Table 3. Summary findings from randomized control trials based on the impact of soy on obesity and satiety.

| Study | Sample Size & Population | Duration | Treatment | Dose; Format | Results | Conclusions |
|-------|--------------------------|----------|-----------|--------------|---------|-------------|
| [146] | 25 abdominally obese men and women | 12 weeks, parallel RCT | Soy protein meal replacement | 4 replacement meals/day; 44 g soy protein, 60–135 mg isoflavones/day | No significant variations in body composition (including weight, BMI, % fat and % lean mass) or cardiometabolic risk factors were observed in the soy replacement meal compared to control | Soy protein affects weight loss to same extent as other proteins |
| [28]  | 39 overweight and obese adults | Daily breakfast for 12 weeks | Soy fiber supplemented biscuits | 100 g soy fiber/day | LDL-C, TC and BMI decreased after 12 weeks ($p < 0.05$) | Soy fiber supplementation may help in weight management and lowering cholesterol levels |
| [144] | 45 patients with non-alcoholic fatty liver disease | 8 weeks, parallel RCT | 3 diets: low-calorie; low-calorie, low carbohydrate; low-calorie, low soy diet | 30 g soy nuts containing: 7 g fat, 9 g fiber, 11.3 g protein, 10 mg sodium, 102 mg phytoestrogens | Soy group had greatest decline in serum liver enzymes and fibrinogens, and malondialdehyde. There were no changes in BMI or weight between the two groups | Soy may mitigate inflammation when combined with a low-calorie diet, but may not affect weight loss in those with non-alcoholic fatty liver disease. |
| [158] | 11 overweight and obese men | 1 day, parallel RCT | Soy protein breakfast meal replacement, followed by standardized lunch 4 h later | Meal replacement, containing 28.7 soy protein (54.6 g total protein), 19.8 g carbohydrate | Meal replacement group had lower glucose levels, glucose and insulin AUC, ghrelin concentrations following breakfast; Fat oxidation was lower after lunch | Soy protein could have a potential hypoglycemic effect, as well as mediating metabolic risk factors and improving weight management |
| [150] | 380 women, BMI of 30–40 kg/m$^2$, 7% of whom took the soy meal replacement | 12 months, with meal replacement taken for a maximum of 3 months (most did so during first 3 months of study) | Soy-based meal replacement (in addition to baseline lifestyle program administered to both control and treated groups) | Soy-yogurt-honey product: 85% soy-protein isolate, 17% milk protein | Weight loss was greater in meal replacement group ($p = 0.1$); health-related quality of life scores also increased more compared to control | Soy protein consumption may enhance weight loss when combined with a lifestyle intervention program focusing on diet and physical activity |
| [149] | 64 overweight/obese subjects | 12 weeks, parallel RCT | Black soy peptide (BSP) supplementation | 4.5 g/day in tablet form | After 12 weeks, BSP group had statistically significant decrease in body weight, body fat mass, body fat % and plasma leptin (vs. no change in placebo); no change in inflammatory markers or lipid profiles between groups | Black soy peptide seems to have a role in weight and fat mass regulation in overweight subjects, perhaps by altering leptin levels |
Table 3. Cont.

| Study | Sample Size & Population | Duration | Treatment | Dose; Format | Results | Conclusions |
|-------|--------------------------|----------|-----------|--------------|---------|-------------|
| [156] | 116 healthy men and women | 6 weeks, parallel RCT | Soy muffins (vs. control wheat muffins) | 12.5 g soy protein muffin; 2 muffins/day | Higher fullness scores in soy muffin group on a Visual Analog Scale ($p = 0.002$) vs. control group | Replacing wheat flour with soy flour may increase perceived satiety. |
| [145] | 38 obese men and women | 8 weeks, parallel RCT | Whole soy powder bar (vs. wheat control) | 1 bar eaten 1–2 h before dinner daily | Compared to control, soy group had lower BMI, waist circumference and body fat % ($p < 0.05$). No significant differences in weight loss, TC, LDL = C, or insulin were found between the two groups | Whole soy can complement the benefits of a weight-loss program; whether it does so by mediating glycemic response remains unclear based on this study. |
| [147] | 48 obese Japanese adults | 20 weeks, parallel RCT | Soy protein | Soy protein intervention containing 12 g soy protein, 9 g milk protein; consumed at breakfast | Visceral and subcutaneous fat, body weight and BMI decreased significantly after 20 weeks of milk protein ingestion, but not after ingestion of soy protein containing milk protein | Milk protein has a more substantial impact on markers of weight loss compared to a combination of milk and soy protein. |

AUC = area under blood glucose response curve; BMI = body mass index; LDL-C = low density lipoprotein cholesterol; RCT = randomized control trial; TC = total cholesterol.
5. Conclusions

Isoflavones and their metabolites appear to improve blood pressure, glycemic control, obesity, and inflammation. Evidence on the hypotensive effects of other soy components such as protein, fiber, lecithins, and saponins is limited by a dearth of available RCTs and current observations suggest that these components do not elicit significant hypotensive effects. However, these constituents may act in synergy with soy protein to modulate plasma lipids to a greater extent than when soy protein is independently consumed. Few human studies have examined the effect of minor soy constituents on glycemic control, despite convincing mechanistic evidence supporting these effects. The anti-adipogenic effects of isoflavones have been tested mostly in animal models, with promising results; however, soy protein does not appear to improve body composition greater than what is achieved with comparable levels of milk protein. A limited body of literature suggests that soy protein and other constituents may enhance satiety. Together, these findings demonstrate the importance of several soybean bioactive compounds in reducing CVD risk that complement the well-established effects of soy protein. Additional RCTs examining the independent effects of these components, and the role of the microbiome in mitigating these effects, are needed to support the direct health benefits of these novel bioactive compounds.

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Abbreviations

AUC area under the curve
BMI body mass index
cAMP cyclic adenosine monophosphate
CHD coronary heart disease
CRP C-reactive protein
CVD cardiovascular disease
FBG fasting blood glucose
HDL-C high density lipoprotein cholesterol
HOMA-IR homeostatic model assessment of insulin resistance
LDL-C low density lipoprotein cholesterol
LDLR low density lipoprotein cholesterol receptor
oxLDL oxidized LDL cholesterol
MetS metabolic syndrome
NO nitric oxide
ODMA O-desmethylangolensin
PPAR peroxisome proliferator activated receptor
PKA protein kinase A
RCT randomized controlled trial
ROS reactive oxygen species
TC total cholesterol
TNF-α tumor necrosis factor alpha
TG triglycerides
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